Post-polyploidisation morphotype diversification associates with gene copy number variation

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Genetic models for polyploid crop adaptation provide important information relevant for future breeding prospects. A well-suited model is *Brassica napus*, a recent allopolyploid closely related to *Arabidopsis thaliana*. Flowering time is a major adaptation trait determining life cycle synchronization with the environment. Here we unravel natural genetic variation in *B. napus* flowering time regulators and investigate associations with evolutionary diversification into different life cycle morphotypes. Deep sequencing of 35 flowering regulators was performed in 280 diverse *B. napus* genotypes. High sequencing depth enabled high-quality calling of single-nucleotide polymorphisms (SNPs), insertion-deletions (InDels) and copy number variants (CNVs). By combining these data with genotyping data from the Brassica 60 K Illumina® Infinium SNP array, we performed a genome-wide marker distribution analysis across the 4 ecogeographical morphotypes. Twelve haplotypes, including *Bna.FLC.A10*, *Bna.VIN3.A02* and the *Bna.FT* promoter on C02_random, were diagnostic for the diversification of winter and spring types. The subspecies split between oilseed/kale (*B. napus ssp. napus*) and swedes/rutabagas (*B. napus ssp. napobrassica*) was defined by 13 haplotypes, including genomic rearrangements encompassing copies of *Bna.FLC*, *Bna.PHYA* and *Bna.GA3ox1*. De novo variation in copies of important flowering-time genes in *B. napus* arose during allopolyploidisation, enabling sub-functionalisation that allowed different morphotypes to appropriately fine-tune their lifecycle.

Polyplod crops like wheat, potato, oats and rapeseed have been enormously successful as field crops because of their huge adaptation potential. Indeed, the fact that all flowering plants derive from ancient or recent polyploidisation events¹,² points to an enormous evolutionary advantage associated with polyploidy. On the other hand, most polyploid events do not lead to a successful establishment of a new species³. Understanding how polyploids achieve adaptive potential has important implication for breeding in the context of environmental change.

On the other hand, the complexity of polyploid genomes has considerably restricted large-scale genetic studies of polyploid species⁴–⁶, so broad conclusions are often drawn based on diploid model plants like *Arabidopsis thaliana*. The polyploid crop most closely related to *A. thaliana* is rapeseed (*Brassica napus*), making it an excellent system to transfer information from the model to the crop. Despite its very recent origin and strong allopolyploidisation bottleneck⁶, rapeseed can be grown from boreal to subtropical and semi-arid areas, a result of strong differentiation into distinctly different morphotypes⁷.

The morphotype with highest seed yields is the biannual winter oilseed type⁸. The prerequisites for this lifecycle are winter hardiness for winter survival, along with vernalisation requirement to avoid pre-winter flowering⁹. In subtropical areas, cultivation of semi-winter types that can be vernalised in warmer temperatures is possible⁵. Boreal or semi-arid regions have periods of low plant survival rates, either due to strong winter freezing or extreme heat stress. In these regions, annual spring types are prominent. These are neither winter-hardy nor vernalisation-dependent, and the short growing season strongly limits yield potential. *B. napus* can also be grown as beet-like forms, known as swedes or rutabagas, which form a different subspecies (ssp. napobrassica)⁷. Swedes are generally of winter type, however have limited winter-hardiness and require extended vernalisation to flower (Fig. 1). No wild-types of *B. napus* are known, hence the species is assumed to have arisen in cultivation⁷, with at least one origin believed to be as recent as a few hundred years ago⁹. The different cultivated forms are bred in separate breeding pools, with introgression between morphotypes only in cases of extreme introgression benefit.
However, this necessitates tedious backcrossing programs to restore the required ecogeographic adaptation characters\textsuperscript{10}. Knowledge of the factors determining lifecycle traits like vernalisation requirement and flowering time is crucial for successful exchange of genetic material between \textit{B. napus} gene pools\textsuperscript{10}.

Although the mechanisms of vernalisation have been studied in depth in Arabidopsis, specific winter or spring alleles were not yet defined for \textit{B. napus}. The predominant assumption is that the underlying genetic mechanisms are identical or very similar across crucifer species. The allopolyploid \textit{B. napus} carries two almost intact subgenomes from the ancestors \textit{Brassica rapa} (A subgenome donor) and \textit{Brassica oleracea} (C subgenome donor). Both ancestral subgenomes arose from a common, hexaploid ancestor, raising the theoretical copy number of Arabidopsis gene homologs to six. Due to post-polyploidisation genome reduction, the average gene copy number is 4.4\textsuperscript{11}, whereby considerable variation has been observed among different gene families, with copy number ranging from 1 to 12\textsuperscript{12}. Homology-driven chromosome rearrangements during allopolyploidisation are a key driver of such variation\textsuperscript{6,12}. Copy number variations (CNVs) have been found to impart large phenotypic influence in several plant species like Arabidopsis\textsuperscript{13}, wheat\textsuperscript{14}, potato\textsuperscript{15} and maize\textsuperscript{16}, but also in domestic animals\textsuperscript{17} and humans\textsuperscript{18}.

In Arabidopsis, \textit{FLOWERING LOCUS C} (\textit{FLC}) is the major repressor for the activity of the central flowering transcription factor \textit{FLOWERING LOCUS T} (\textit{FT})\textsuperscript{19}. This gene cannot be expressed before \textit{FLC} protein levels drop\textsuperscript{19}, however when this occurs \textit{FT} can be activated by the photoperiod pathway via the transcription factor \textit{CONSTANS} (\textit{CO})\textsuperscript{20}. Downregulation of \textit{FLC} takes place at the transcriptional level. The \textit{FLC} chromatin is modified and rearranged in order to stabilize a new inactive form\textsuperscript{21,22}. Different mechanisms are involved in the structural regulation of \textit{FLC} gene activity, including both autonomous regulators and the vernalisation pathway\textsuperscript{22,23}. Three different mechanisms may exist for the breakdown of vernalisation requirement: (i) alteration of \textit{FLC} regulating factors like \textit{FRIGIDA} (\textit{FRI}); (ii) alteration of \textit{FLC} gene sequence or activity; (iii) alteration of \textit{FLC} binding sites or \textit{FT} promoter sequences. Arabidopsis annuals and biannuals have been found to differ either in \textit{FRI} or in \textit{FLC}\textsuperscript{24}, indicating that the Arabidopsis winter-spring split is governed by the first two levels of regulation. As a consequence, research on \textit{B. napus} vernalisation has been heavily focused on investigating \textit{FLC} homologs\textsuperscript{25-27}. Indeed, a number of QTL studies in different mapping populations have suggested \textit{FLC} loci as candidates for flowering time in \textit{B. napus}, including populations without vernalisation requirement\textsuperscript{28-32}. Moreover, it has been reported that a transposon insertion in the first intron of \textit{Bna.FLC.A10} is associated with the vernalisation requirement of winter-type rapeseed\textsuperscript{27}.

The aim of the present work was therefore the definition of morphotype-specific alleles or haplotypes that might further our understanding of vernalisation control in a complex allopolyploid, and simultaneously allow breeders to successfully select for desirable lifecycle traits. By comparing results of vernalisation experiments with data from genome-wide marker distribution analysis, targeted deep-sequencing of essential flowering time regulators and the \textit{FT} promoter, and coverage analysis to estimate CNV, we provide novel insights that reveal the complexity of post-polyploidisation morphological diversification in an important crop species.

Material and methods

\textbf{Plant material and phenotyping.} A panel of 280 genetically diverse \textit{B. napus} inbred lines (selfed for 5 or more generations) was grown in Giessen, Germany (50° 35' N, 8° 40' E) in 2012. The plant material was part of the ERANET-ASSYST \textit{B. napus} diversity set that has been described previously\textsuperscript{25,34}. Winter-type rapeseed and swede
accessions were grown in autumn-sown trials, whereas spring-type and semi-winter accessions were grown in spring-sown trials. Plots were sown in a completely randomized block design with a harvest plot size of $3 \times 1.25$ m in a single replicate (containing around 200 plants).

In a separate experiment, a selection of 33 genotypes from the same set was grown in the greenhouse under semi-controlled conditions ($20^\circ$C). These genotypes were selected to represent spring, winter and swede material with different CNV patterns for $Bna.FLC$. Twenty seeds were sown in vermiculite, before being transplanted after one week into plates in soil, with 5 replicates per treatment. Four weeks after planting, these plants were either transferred to a climate chamber for vernalisation at $4^\circ$C and short-day conditions for 6 weeks (mild

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**Figure 2.** Distribution of SNPs/InDels (above) and CNV events (below) over all target gene copies. The chromosomal locations of the copies are given below the common Arabidopsis gene name (white background), with colors representing the respective type of sequence variation observed (see color code below each diagram). Upper panel: Silent SNPs are not indicated if synonymous or non-synonymous SNPs are present in the same copy, and synonymous SNPs are not indicated if non-synonymous SNPs are present in the same copy. Lower panel: Gene copies showing two different colors are deleted in some lines and duplicated in some others.
Figure 3. 2D plots of PCA for the total population. The explained variance is given in brackets. Colors indicate the cluster. Cluster 1 is shown in red, cluster 2 in green and cluster 3 in black. Letters indicate the morphotype: w for winter, s for spring, e for semi-winter and d for swedes.

Figure 4. Genome-wide distribution of χ² p-values tested against equal distribution in winter and spring material. The chromosomes are coloured differently. The solid lines indicate the marker cut-off threshold of 0.1%.
beginning of flowering (BBCH 61) was tracked daily for every single plant.

Marker name | Chromosome | Position | winter allele | spring allele | —log(p) | located in gene | clear winter | clear spring | mixed | deletions
--- | --- | --- | --- | --- | --- | --- | --- | --- | --- | ---
Bn-A01-p4641747 | chrA01 | 4261277 | 132 | 13 | 4 | 101 | 39.3 | 136 | 102 | 12 | 3
Bn-A01-p4641802 | chrA01 | 4261332 | 132 | 13 | 4 | 101 | 39.3 | 129 | 104 | 12 | 8
Bn-A01-p4803773 | chrA01 | 4413051 | 128 | 9 | 6 | 105 | 39.8 | 148 | 101 | 2 | 2
Bn-A02-p3207085 | chrA02 | 695308 | 131 | 7 | 4 | 106 | 42.8 | 129 | 113 | 9 | 1
Bn-A02-p3208827 | chrA02 | 700399 | 129 | 7 | 5 | 106 | 41.7 | 87 | 65 | na | 7
Bn-A02-p3255898 | chrA02 | 786193 | 130 | 8 | 5 | 101 | 39.9 | 128 | 104 | 12 | 8
Bn-A02-p3297592 | chrA02 | 787627 | 128 | 6 | 6 | 104 | 40.6 | 136 | na | 5
Bn-A02-p3299206 | chrA02 | 789246 | 130 | 6 | 5 | 102 | 41.2 | 136 | na | 3
Bn-A02-p3300731 | chrA02 | 790766 | 128 | 6 | 5 | 105 | 39.0 | 129 | 113 | 9 | 1
Bn-A02-p3302725 | chrA02 | 792753 | 126 | 6 | 7 | 105 | 39.8 | 129 | 113 | 9 | 1
Bn-A02-p3361391 | chrA02 | 849106 | 126 | 6 | 8 | 104 | 39.1 | 126 | 102 | 12 | 8
Bn-A02-p35097701 | chrA02 | 3096806 | 136 | 12 | 1 | 99 | 41.9 | 126 | 102 | 12 | 8
Bn-A02-p3517045 | chrA02 | 3104382 | 137 | 11 | 2 | 101 | 42.9 | 129 | 113 | 9 | 1
Bn-A03-p6576575 | chrA03 | 5874703 | 126 | 3 | 9 | 111 | 42.6 | 126 | 102 | 12 | 8
Bn-A03-p6636780 | chrA03 | 5928259 | 131 | 5 | 6 | 109 | 44.0 | 129 | 113 | 9 | 1
Bn-A03-p9836757 | chrA03 | 9057095 | 128 | 8 | 7 | 103 | 39.1 | 136 | 110 | na | 7
Bn-A07-p15352802 | chrA07 | 12760795 | 126 | 7 | 9 | 106 | 39.0 | 136 | 111 | na | 5
Bn-A09-p30805314 | chrA09 | 2855763 | 125 | 4 | 9 | 107 | 40.2 | 133 | 115 | na | 5
Bn-A09-p30805387 | chrA09 | 2855709 | 125 | 4 | 9 | 110 | 41.4 | 133 | 115 | na | 5
Bn-A09-p30875173 | chrA09 | 28628531 | 131 | 11 | 5 | 103 | 39.9 | 129 | 113 | 9 | 1
Bn-A09-p30909393 | chrA09 | 28655613 | 131 | 9 | 4 | 105 | 41.7 | 129 | 101 | 19 | 4
Bn-A09-p30918224 | chrA09 | 28662308 | 131 | 13 | 4 | 101 | 38.9 | 126 | 102 | 12 | 8
Bn-A10-p7354742 | chrA10 | 9020292 | 128 | 7 | 9 | 106 | 39.8 | 135 | 115 | na | 3
Bn-A10-p735555 | chrA10 | 9020402 | 127 | 6 | 9 | 106 | 39.8 | 135 | 115 | 0 | 3
Bn-scaffold_17109_2-p79906 | chrA10 | 14916811 | 121 | 1 | 17 | 111 | 38.9 | 122 | 128 | na | 3
Bn-scaffold_16002_1-p1767743 | chrA02 | 12604057 | 127 | 7 | 9 | 105 | 39.0 | 134 | 114 | na | 5
Bn-scaffold_18206_3-p62755 | chrA02 | 18925562 | 131 | 12 | 4 | 100 | 38.9 | 143 | 104 | na | 6
Bn-scaffold_16912_1-p190291 | chrA02 | 12697195 | 129 | 9 | 8 | 104 | 39.0 | 137 | 112 | 2 | 2
Bn-scaffold_20836_1-p198391 | chrA02 | 12804839 | 129 | 8 | 9 | 105 | 39.4 | 137 | 112 | 2 | 2
Bn-scaffold_20836_1-p197940 | chrA02 | 12805469 | 130 | 8 | 9 | 105 | 39.4 | 137 | 112 | 2 | 2
Bn-scaffold_20836_1-p196601 | chrA02 | 12807036 | 129 | 8 | 9 | 104 | 39.0 | 137 | 112 | 2 | 2

Table 1. Marker distributions of SNP markers associated with the winter-spring morphotype split in B. napus, along with the most closely associated markers from deep sequencing for the winter-spring split. The table shows the marker name, chromosomal position and the number of lines carrying either a winter or a spring allele in the respective winter-type and spring-type populations. The table also gives the —log(p-value) used to determine the split markers, along with the gene ID where the marker is located. If empty, the marker is non-genic. The markers with the highest —log(p-value) in each split region are shown in bold letters. The last four columns of the table show how many clear winter or spring haplotypes were counted, along with the number of mixed haplotypes and deletions. For regions only containing one marker, mixed haplotypes do not apply (na).

Regions from deep sequencing

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vernalisation) or 12 weeks (strong vernalisation), or kept in the greenhouse (no vernalisation). Begin of flowering (BBCH 61) was tracked daily for every single plant.
DNA isolation. Leaf material for genomic DNA extraction was harvested in spring 2012 from the field trial in Giessen, Germany. Pooled leaf samples were taken from at least 5 different plants per genotype, immediately shock-frozen in liquid nitrogen and kept at \(-20^\circ C\) until extraction. Leaf material was ground in liquid nitrogen with a mortar and pestle. DNA was extracted using a common CTAB protocol modified from Doyle and Doyle (1990) as described earlier. DNA concentration was determined using a Qubit fluorometer and the Qubit dsDNA BR assay kit (Life Technologies, Darmstadt, Germany) according to the manufacturer's protocol. DNA quantity and purity was further checked on 0.5% agarose gel (3 V/cm, 0.5xTBE, 120 min).

Selection of target genes. As described previously, a set of 29 *A. thaliana* flowering time genes was selected to cover the entire genetic network controlling flowering time, including circadian clock regulators (*CYCLING DOF FACTOR 1* (CDF1), *EARLY FLOWERING 3* (ELF3), *GIGANTEA* (GI) and *ZEITLUPE* (ZTL)), the input pathways for vernalisation (*EARLY FLOWERING 7* (ELF7), *EARLY FLOWERING IN SHORT DAYS* (EFS), *FLOWERING LOCUS C* (FLC), *FRIGIDA* (FRI), *SHORT VEGTATIVE PHASE* (SVP), *SUPPRESSOR OF FRIGIDA 4* (SUF4), *TERMINAL FLOWER 2* (TFL2), *VERNALISATION 2* (VRN2), *VERNALISATION INSENSITIVE 3* (VIN3)), photoperiod sensitivity (*CONSTANS* (CO), *CRYPTOCHROME 2* (CRY2), *PHYTOCHROME A* (PHYA), *PHYTOCHROME B* (PHYB)) and gibberellin (*GIBBERELLIN-3-OXIDASE 1* (GA3ox1)), along with downstream signal transducers (*AGAMOUS-LIKE 24* (AGL24), *APETALA 1* (AP1), *CAULIFLOWER* (CAL), *FLOWERING LOCUS D* (FD), *FLOWERING LOCUS T* (FT), *FRUITFUL* (FUL), *LEAFY* (LFY), *SQUAMOSA PROMOTOR PROTEIN LIKE 3* (SPL3), *SUPPRESSOR OF CONSTANS 1* (SOC1), *TEMPRANILLO 1* (TEM1), *TERMINAL FLOWER 1* (TFL1)). On top, we also included *CIRCADIAN CLOCK ASSISTED 1* (CCA1), *FLAGELLIN-SENSITIVE 2* (FLS2), *GLYCIN-RICH PROTEIN 7* (GRP7), *GLYCIN-RICH PROTEIN 8* (GRP8), *GORDITA* (GORD) and *SENSITIVITY TO RED LIGHT REDUCED 1* (SRR1).

A full list of gene names and putative functions is provided in Supplementary Table 1.

Bait development. In order to perform target enrichment, complementary sequences of 120 nt length were first developed for each target region. A group of 120mer oligonucleotide sequences covering a certain target region is hereinafter referred to as a bait group for that target region, while collectively all bait groups are referred to as the bait group pool. In the present study the bait group pool for the sequence capture, developed mainly...
### Table 2. Marker distributions of SNP markers associated with the swede vs. non-swede morphotype split in *B. napus*, along with the most closely associated markers from deep sequencing for the swede vs. non-swede split.

The table shows the marker name, chromosomal position and the number of lines carrying either a non-swede or a swede allele in the respective non-swede and swede populations. The table also gives the $-\log(p$-value) which used to determine the split markers, along with the gene ID or the name of the gene where the marker is located. If empty, the marker is non-genic. The markers with the highest $-\log(p$-value) in each split region are shown in bold letters. The last four columns of the table show how many clear non-swede or swede haplotypes were counted, along with the number of mixed haplotypes and deletions. For regions only containing one marker, mixed haplotypes do not apply (na).

| Marker name | Chromosome | Position | non-swede allele | swede allele | located in gene | clear non-swede | clear swede | mixed | deletions |
|-------------|------------|----------|------------------|--------------|----------------|----------------|-------------|-------|-----------|
| chrA03_4639027 | chrA03 | 4639027 | 260 0 0 11 | 57.7 | Bna.VIN3.A03 | 260 11 na 0 | 263 8 na 0 |
| chrA04_12696607 | chrA04 | 12696607 | 0 8 260 3 | 55.8 | Bna.CCR1.A06 | 257 6 8 0 | 263 8 na 0 |
| chrA06_5607262 | chrA06 | 5607262 | 2 11 258 0 | 56.0 | Bna.CCR1.A06 | 260 11 2 2 | 257 6 8 0 |
| chrA06_5607744 | chrA06 | 5607744 | 0 9 260 2 | 56.3 | Bna.CCR1.A06 | 257 6 8 0 | 263 8 na 0 |
| chrA06_5608016 | chrA06 | 5608016 | 1 11 259 0 | 56.9 | Bna.CCR1.A06 | 260 11 2 2 | 257 6 8 0 |
| chrA06_5608089 | chrA06 | 5608089 | 1 11 259 0 | 56.9 | Bna.CCR1.A06 | 260 11 2 2 | 257 6 8 0 |
| chrA06_5614815 | chrA06 | 5614815 | 0 8 260 3 | 55.8 | Bna.CCR1.A06 | 260 11 2 2 | 257 6 8 0 |
| chrA08_14983629 | chrA08 | 14983629 | 260 3 0 8 | 55.8 | Bna.TEM1.A08 | 263 8 na 0 | 263 8 na 0 |
| chrA9_11993194 | chrA9 | 11993194 | 0 8 260 3 | 55.8 | BnaA09g19070D | 263 8 na 0 | 263 8 na 0 |
| chrA9_11993662 | chrA9 | 11993662 | 0 8 260 3 | 55.8 | BnaA09g19070D | 263 8 na 0 | 263 8 na 0 |
| chrA9_11995810 | chrA9 | 11995810 | 0 8 260 3 | 55.8 | BnaA09g19070D | 263 8 na 0 | 263 8 na 0 |
| chrA10_p12192238 | chrA10 | 19312044 | 0 8 260 3 | 55.8 | BnaA09g19070D | 263 8 na 0 | 263 8 na 0 |
| chrA9_32435440 | chrA9 | 32435440 | 2 11 258 0 | 56.0 | Bna.PHYA.A09 | 263 8 na 0 | 263 8 na 0 |
| chrA9_32435455 | chrA9 | 32435455 | 2 11 258 0 | 56.0 | Bna.PHYA.A09 | 263 8 na 0 | 263 8 na 0 |
| chrA9_32437048 | chrA9 | 32437048 | 0 10 260 1 | 56.9 | Bna.CCR1.A06 | 263 8 na 0 | 263 8 na 0 |
| chrBna.ELF7.A10 | chrBna.ELF7.A10 | 17067264 | 0 8 260 3 | 55.8 | BnaA09g19070D | 263 8 na 0 | 263 8 na 0 |
| chrC01_14477235 | chrC01 | 14477235 | 0 9 260 2 | 56.3 | Bna.FD.C01 | 262 8 1 0 | 263 8 na 0 |
| chrC01_14477273 | chrC01 | 14477273 | 260 2 0 9 | 56.3 | Bna.FD.C01 | 262 8 1 0 | 263 8 na 0 |
| chrC01_1447516 | chrC01 | 1447516 | 0 9 260 2 | 56.3 | Bna.FD.C01 | 262 8 1 0 | 263 8 na 0 |
| chrC01_1447693 | chrC01 | 1447693 | 260 2 0 9 | 56.3 | Bna.FD.C01 | 262 8 1 0 | 263 8 na 0 |
| chrC01_1447972 | chrC01 | 1447972 | 0 9 260 2 | 56.3 | Bna.FD.C01 | 262 8 1 0 | 263 8 na 0 |

**Regions from deep sequencing**

- chrA01_random_477115
- chrA02_6053137
- chrA03_6243410
- chrA04_12695445
- chrA05_5425314
- chrA07_23775522
- chrA10_13359226
- chrA10_14998679
- chrAnn_random_610372
- chrAnn_random_20504534
- chrC03_8403949
- chrC03_random_5400150
using gene sequences from *B. rapa* or *B. oleracea*\(^1\), was modified in order to improve specificity. Enriched regions captured in our previous study\(^12\) were classified into target regions and non-target regions. The bait pool was then blasted against target and non-target regions with an E-value cut-off of \(10^{-10}\). Baits which had excessive non-target hits were manually removed. This was the case for bait groups on *FT*, *FUL* and *PHYA*. For some bait groups (*AP1*, *CO*, *SOC1*), too many baits (>30%) were deleted. In these cases, bait groups were created using a pre-publication draft (version 4.0) of the *B. napus* ‘Darmor-Bzh’ reference genome sequence assembly, which was kindly made available prior to public release by INRA, France, Unité de Recherche en Génomique Végétale\(^6\), using the Agilent Genomic Workbench program SureDesign (Agilent Inc., Santa Clara, CA, USA). These replaced the corresponding bait groups developed previously using *B. rapa* or *B. oleracea*. Bait groups were created using the ‘Bait Tiling’ tool. The parameters were set as follows: Sequencing Technology: ‘Illumina’, Sequencing Protocol: ‘Paired-End long Read (75 bp+),’ ‘Use Optimized Parameters (Bait length 120, Tiling Frequency 1x),’ ‘Avoid Overlap: ’20,’ ‘User defined genome,’ ‘Avoid Standard Repeat Masked Regions’. Baits for genes on the minus-strand were developed in sense, while baits on the plus-strand were developed in antisense.

In total, 63 bait groups were created for *B. rapa* copies of the target genes, 71 bait groups for *B. oleracea* copies and 24 bait groups for *B. napus* copies.

**Sequence capture and sequencing.** Custom bait production was carried out by Agilent Technologies (Agilent Inc., Santa Clara, CA, USA) using the output oligonucleotide sequences from eArrayXD. Sequence capture was performed using the SureSelectXT 1 kb–499 kb Custom Kit (Agilent Inc., Santa Clara, CA, USA) according to the manufacturer’s instructions. The resulting TruSeq DNA library (Illumina Inc., San Diego, CA, USA) was sequenced on an Illumina HiSeq 2500 sequencer at the Max Planck Institute for Breeding Research (Cologne, Germany) in 100 bp single-read mode.

**Sequence data analysis.** Quality control of the raw sequencing data was performed using FASTQC. Reads were mapped onto version 4.1 of the *B. napus* ‘Darmor-Bzh’ reference genome sequence assembly\(^6\). Mapping was performed using the SOAPaligner algorithm\(^35\), with default settings and the option \(r=0\) to extract uniquely aligned reads. Removal of duplicates, sorting and indexing was carried out with samtools version 0.1.19\(^36\). Alignments were visualised using the IGV browser version 2.3.12\(^37\). Enriched regions and coverage

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**Figure 6.** Distribution of clear non-swede haplotypes (above) and clear swede haplotypes (below) in the total population for all 13 identified split regions. Mixed haplotypes were not counted. The distribution on morphotypes is colour-coded.
Haplotype differences were calculated using the bedtools software with multiBamCov\(^9\). Calling of single nucleotide polymorphisms (SNPs) was performed with the algorithm mpileup in the samtools toolkit. SNP and InDel annotation was performed using CooVar\(^9\). Target regions were defined using the gene annotation list from the B. napus ‘Darmor-bzh’ v4.1 reference genome\(^6\) and BLAST position results of the bait pool (E-value cut-off \(10^{-100}\)) on the mapping reference, and used to calculate the fraction of target covered. For InDel calling, a separate mapping using Bowtie2\(^40\) was performed, as described previously\(^41\). Removal of duplicates, sorting and indexing was carried out with samtools version 0.1.19. An initial InDel calling was performed using samtools mpileup, and realignment of reads around InDels was performed using GATK RealignerTargetCreator, version 3.1.142. A final InDel calling was then performed as described above. InDels were filtered for a minimum mapping quality of 30 and a read depth of 10 or more using vcftools\(^43\).

Read coverage for each captured region was normalised as follows: normalised coverage = \((\text{number of reads per region} * \text{total length of genome}) / (\text{total number of aligned reads per genotype} * \text{average read length})\). Copy number variation (CNV) in a given target region was assumed if the ratio of normalised coverage(genotype)/normalised coverage(all genotypes) was smaller than 0.5 or higher than 1.5, respectively.

Sequencing data for 3 genotypes from a former experiment (Silona, Campino, Magres Pajberg)\(^12\) were analysed separately with the same pipeline to allow inclusion in the marker distribution analysis.

SNP genotyping and pre-processing. The 283 accessions were genotyped using the Brassica 60 K Illumina® Infinium SNP array by TraitGenetics GmbH (Gatersleben, Germany). We used the SNP positions as published in\(^44\). Heterozygous calls were treated as missing values. Moreover, we used the deep sequencing data to include all confidently called SNPs in biallelic state which lay in the analysed regions. Confidently called InDels were included by coding reference alleles as AA, insertions as CC, deletions as TT and heterozygous calls as missing values. The SNP matrix from the SNP array and the SNP and InDel data from deep sequencing were combined to one single marker file and sorted by position. The subsequent marker set contained 43733 markers. After pre-processing the marker set for non-missing marker values >0.9, minor allele frequency >0.01 and individuals (genotypes) with non-missing individual markers >0.8, we retained 33944 unique SNP markers and a population of 271 individuals for marker distribution analysis. Data pre-processing was performed with R (version 3.1.0) using the package GenABEL\(^45\).

Population structure. Population structure analysis and visualization were performed in R (version 3.1.0) using the package SelectionTools (http://fb09-pg-s207.agrar.uni-giessen.de/~frisch-m/), which applies principal component analysis based on genetic distances calculated according to the euclidean modified Rodger’s distance method. The most likely number of population subclusters was determined to be 3 by plotting the within-cluster
We concluded that the winter material was genetically least diverse, while spring material was more diverse, second one 93 spring type accessions and a semi-winter type accession, and the third and most diverse cluster (Fig. 3). The population falls into three main clusters: the first cluster contained 137 winter type accessions, the component explained 24.1% of the variation, while further components explained 5.4, 2.5 and 2.1%, respectively a Principal Component Analysis (PCA) showed a strong population substructure, as the first principal com-

| Gene ID | Chromosome | start | stop   | deletions | duplications | mean coverage | Gene name |
|--------|------------|-------|--------|-----------|--------------|---------------|------------|
| BnaA08g15780D | chrA08  | 13097823 | 13098361 | 11 | 9 | 0 | 1641.5 | no annotation |
| BnaA09g48410D | chrA09  | 32434233 | 32438771 | 31 | 0 | 10 | 1136.2 | Bna.PHYA.chrA09 |
| BnaA09g57140D | chrA09_random | 4043861 | 4045464 | 1 | 0 | 4 | 8 | 1675.6 | Bna.GA3ox.chrA09_random |
| BnaA10g22080D | chrA10  | 14998617 | 15003197 | 2 | 0 | 2 | 9 | 1321.5 | Bna.FLC.chrA10 |
| BnaC08g38580D | chrC08  | 34776298 | 34779240 | 9 | 8 | 1 | 1226.1 | Bna.CCR1.chrC08 |
| BnaC08g38810D | chrC08  | 34907098 | 34908735 | 4 | 9 | 0 | 0 | 1581.1 | Bna.GA3ox.chrC08 |
| BnaC09g42660D | chrC08  | 36746642 | 36751390 | 6 | 10 | 15 | 0 | 1537.7 | Bna.PHYA.chrC08 |
| BnaC09g42670D | chrC08  | 36752307 | 36753108 | 7 | 9 | 14 | 0 | 1660.2 | germpl like protein |
| BnaC09g46500D | chrC09  | 46335350 | 46350992 | 2 | 9 | 13 | 0 | 1096.9 | Bna.FLC.chrC09 |
| BnaC09g46540D | chrC09  | 46366645 | 46371880 | 3 | 9 | 11 | 0 | 1031.5 | Bna.FLC.chrC09 |

Table 3. Distribution of deletion and duplication events in the non-swede and swede populations. The table shows the gene ID, chromosomal position and the number of lines in the respective non-swede and swede populations which carry either a deletion or duplication. The table also gives the mean coverage of the respective gene and the common gene name.

sum of squares against the possible number of clusters, ranging from 1 to 15. K-means clustering was then performed in R using SelectionTools.

Marker distribution analysis. For every marker, we counted the allele frequency of the alternative allele in each morphotype pool. The ratio between the frequency of the allele in the winter pool (winter + swedes) and the spring pool (semi-winter + spring) was used to assign the allele as a winter or spring allele. If the ratio was <1, the alternative allele was denoted spring (s), if it was >1, the alternative allele was denoted winter (w). We then first tested if the marker would be suitable to explain a morphotype split, by comparing the observed distribution of w alleles in the winter pool (without swedes) and s alleles in the spring pool (without semi-winter) with the expected distribution (139/114), using a χ² test. Only markers which did not show significant deviation from this distribution (p-value > 0.1) were considered in the next step. In the next step, we tested the distribution against random distribution between the pools, by comparing the observed distribution of w alleles in winter/s alleles in spring against the expected random distribution of 69.5/57/69.5/57. We then considered the top 0.1% of −log(p-value) as split markers. The same was done for the swede and non-swede material.

Results

Deep sequencing and variant calling. We defined regions as genetic regions which were covered with a mean coverage in the population of at least 10. In total, we analyzed 1184 regions, of which 637 regions were annotated as genes. Of these, 184 corresponded to the intended target genes. Two target genes copies for VERNALISATION INSENSITIVE 3 (VIN3) (Bna.VIN3.A01 and Bna.VIN3.C01) had insufficient coverage for this analysis and were not considered. Among the non-genic regions, we found 33 regions giving a BLAST hit to the FLOWERING LOCUS T (FT) promoter. A further 12 regions were identified as pseudogenes of the target genes. Those regions which were assigned to one of those classes (target genes, target pseudogenes and FT promoter) were summarized as target regions. A gene group is defined as all copies of a specific gene.

We called and annotated 13053 SNPs, of which 4806 were located in the target regions. InDel calling revealed a total of 1894 InDels, with 506 in the target regions. Only 25 InDels were frameshifts, amino acid insertions or splices variants. All gene groups showed potentially functional variation, i.e. at least one copy of the gene group carried either a non-synonymous SNP, stop codon mutation, amino acid insertion, splice variant or frameshift InDel. Altogether, only 7 copies were completely conserved, while 16 copies carried only silent or synonymous variation. Interestingly, no functional variation was observed in two copies of Bna.FLOWERING LOCUS C (FLC) (on chromosomes A02/C02) and two copies of Bna.FT (also on chromosomes A02/C02). Respectively. On the other hand, other copies of Bna.FLC (A03, A10) and Bna.FT (C06) carried a surprisingly large range of variation. Among the genes with frameshift variants were copies of Bna.FRIGIDA (FRI), Bna.PHYTOCHROME A (PHYA), Bna.EARLY FLOWERING IN SHORT DAYS (ELS), Bna.EARLY FLOWERING 7 (ELF7), Bna.PHYTOCHROM B (PHYB), Bna.VERNALISATION 2 (VIN2) and Bna.LEAFY (LFY) (Fig. 2). We also calculated copy number variation (CNV) based on read depth. No gene group was found without CNV, and only two lines were found which did not carry any CNV among the target copies. The distribution of SNPs, InDels and CNVs is shown in Fig. 2.

Population structure. Among the analysed population of 271 accessions, we had 139 winter type accessions, 7 semi-winter type accessions, 114 spring type accessions and 11 swedes. Analyzing this population with a Principal Component Analysis (PCA) showed a strong population substructure, as the first principal component explained 24.1% of the variation, while further components explained 5.4, 2.5 and 2.1%, respectively (Fig. 3). The population falls into three main clusters: the first cluster contained 137 winter type accessions, the second one 93 spring type accessions and a semi-winter type accession, and the third and most diverse cluster contained 11 swedes, 6 semi-winter type accessions, 21 spring type accessions and 2 winter type accessions. We concluded that the winter material was genetically least diverse, while spring material was more diverse,
| Chromosome | Marker with highest p-value | Candidate | Distance from closest split marker [kbp] |
|------------|-----------------------------|-----------|----------------------------------------|
| chrA01    | Bn-A01-p4803773             | dna topoisomerase 2 | 502.7 |
|           |                             | pseudo-response regulator 2 | 57.0 |
|           |                             | agamous-like protein 1 | 386.3 |
|           |                             | probable lysine-specific demethylase jmjd14-like | 877.5 |
|           |                             | small rna 2-o-methyltransferase | 990.8 |
|           |                             | flowering locus c | 557.1 |
|           |                             | transcription factor hy5 | 342.1 |
|           |                             | embryonic flower 1 | 281.6 |
|           |                             | nuclear transcription factor y subunit a-1 | 127.5 |
|           |                             | flowering time control protein fy | 8.1 |
|           |                             | zinc finger protein knuckles | 44.4 |
|           |                             | dna (cytosine-5)-methyltransferase dmr2 | 121.7 |
| chrA02    | Bn-A02-p3207085             | histone deacetylase | 403.8 |
|           |                             | auxin response factor 4 | 81.4 |
|           |                             | e3 sumo-protein ligase szr1 | 53.5 |
|           |                             | ap2-like ethylene-responsive transcription factor toe2 | 1.9 |
|           |                             | pseudo-response regulator 3 | 8.2 |
|           |                             | sensitivity to red light reduced protein | 215.9 |
|           |                             | multicity suppressor of ira1 | 580.9 |
|           |                             | vernalization insensitive 3 | 757.8 |
|           |                             | flowering locus c | 311.8 |
| chrA03    | Bn-A03-p6636780             | histone h2a | 539.8 |
|           |                             | chromatin structure-remodeling complex protein syd | 445.8 |
|           |                             | protein early flowering 4-like | 164.3 |
|           |                             | btb poz domain-containing protein | 48.3 |
| chrA07    | Bn-A07-p15352802            | two-component response regulator arr5 | 352.9 |
|           |                             | sin3 histone deacetylase complex | 654.6 |
|           |                             | protein argonaute 7-like | 894.1 |
|           |                             | floral homeotic protein aptata 1 | 964.5 |
|           |                             | protein early flowering 3-like | 17.9 |
|           |                             | e3 ubiquitin-protein ligase orthrus 2 | 67.9 |
|           |                             | dna methyltransferase | 727.3 |
| chrA09    | Bn-A09-p30909393            | histone acetyltransferase type b catalytic subunit-like | 187.2 |
|           |                             | transcription factor hy5 | 261.7 |
|           |                             | flowering-promoting factor 1-like | 99.2 |
|           |                             | flowering locus c | 81.8 |
| chrC03    | Bn-scaffold_16002_1-p1767743 | btb poz domain-containing protein | 67.5 |
|           |                             | two-component response regulator arr16 | 12.8 |
|           |                             | phy rapidly regulated 1 | 743.0 |
| chrC06    | Bn-scaffold_18206_3-p62755  | histone z | 939.5 |
|           |                             | squamosa-promoter binding protein | 213.7 |
|           |                             | shatterproof1 | 749.7 |
| chrC09    | Bn-scaffold_20836_1-p197940 | set domain isoform 1 | 845.4 |

Table 4. Selected candidate genes for all 12 regions associated with the split between winter-type and spring-type B. napus accessions. The table lists the chromosome and the marker with the most significant deviation from the expected random distribution, together with candidate genes selected based on gene ontology and literature. The last column specifies the distance of the gene to the closest marker within the split-associated region. The candidate gene closest to the split region is shown in italics.
followed by semi-winter and swede material. Overlap between winter and spring pools is minimal, while all other types show more overlap, although swedes are more distant from the winter and spring core clusters.

**Marker distribution analysis.** To analyse marker distribution on a genome-wide scale, we used SNP data from the Brassica 60 K Illumina® Infinium SNP array and combined it with data from deep sequencing (SNPs and InDels). In order to find the most indicative marker patterns for the differential flowering behaviour of winter and spring material, we analyzed the differential marker pattern between the different morphotypes using the χ² test. We first defined "winter" and "spring" alleles by allele frequencies in the different morphotypes and assessed their distribution in both pools. First, we excluded all markers with non-suitable allele frequencies. We regard all markers as non-suitable if their minor allele frequency was too low to explain a population split. This was tested in a foregoing χ² test (see Methods). The remaining markers were tested against random distribution in the

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**Table 5. Selected candidate genes for all defined regions associated with the split between swede and non-swede B. napus morphotypes.** The table lists the chromosome and the marker with the most significant deviation from the expected random distribution, together with candidate genes selected based on gene ontology and literature. The last column specifies the distance of the gene to the closest marker within the split-associated region. A value of "0.0" indicates that the marker lies within the gene. The candidate gene closest to the split region is shown in italics.

| Chromosome | Marker with highest p-value | Candidate | Distance from closest split marker [kbp] |
|------------|-----------------------------|-----------|----------------------------------------|
| chrA03     | chrA03_4639027              | vernalization insensitive 3 | 0.0 |
|            |                             | histone acetyltransferase type b catalytic subunit | 163.4 |
| chrA04     | chrA04_12696607             | protein ovule abortion 4 | 45.0 |
|            |                             | protein early flowering 3-like | 0.1 |
|            |                             | terminal flowering 1 protein 1 | 413.7 |
| chrA06     | chrA06_5608089              | gibberellin 3-oxidase | 181.8 |
|            |                             | cinnamoyl- reductase | 0.0 |
|            |                             | transcriptional factor b3 family protein | 248.7 |
|            |                             | histone acetyltransferase hac12 | 263.1 |
|            |                             | protein elf4-like 4 | 473.9 |
| chrA08     | chrA08_14983629             | ap2-erebp rave subfamily protein rav2 | 0.0 |
|            |                             | cycling dol factor 2 | 170.7 |
|            |                             | cullin 3 | 187.3 |
| chrA09     | chrA09_11993662             | cullin 4 | 982.5 |
|            |                             | agamous-like mads-box protein agl3 | 348.6 |
|            |                             | protein ovule abortion 4 | 367.7 |
| chrA09     | Bn-A09-p21922383            | mads-box protein gordita | 875.9 |
|            |                             | protein suppressor of fri 4 | 759.6 |
|            |                             | nuclear transcription factor y subunit a-7 | 151.7 |
| chrA09     | chrA09_32437048             | phytochrome a | 0.0 |
|            |                             | phytochrome interacting factor 3 | 11.9 |
|            |                             | histone-lysine n-methyltransferase atx2 | 810.9 |
| chrA10     | chrA10_17106744             | probable lysine-specific demethylase elf6-like | 457.6 |
|            |                             | protein lby cca1-like 1 | 51.2 |
|            |                             | pseudo-response regulator 7 | 41.0 |
|            |                             | protein early flowering 7 | 0.0 |
| chrC01     | chrC01_1447516              | bzip transcription factor | 0.0 |
| chrC08     | Bn-scaffold_16770_1-p1357882 | dek domain-containing chromatin associated protein | 942.4 |
| chrC08     | chrC08_36752954             | phytochrome a | 1.5 |
|            |                             | phytochrome interacting factor 3 | 10.5 |
|            |                             | medea | 551.6 |
| chrC08     | Bn-scaffold_16389_1-p12505   | histone-lysine n-methyltransferase atx2 | 434.0 |
|            |                             | dna helicase | 405.8 |
|            |                             | tata-box-binding protein 2 | 107.5 |
| chrC09     | chrC09_43739821             | chromo domain-containing protein lhp1-like | 985.9 |
|            |                             | della protein | 866.0 |
|            |                             | coa | 0.0 |
|            |                             | coa | 5.9 |
|            |                             | sepallata2 | 47.5 |

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Figure 8. Distribution of flowering plants in the vernalisation trial. The height of the bars represents the number of replications which could be phenotyped. Yellow indicates flowering plants, grey indicates non-flowering plants. The different treatments are framed in different colors, with green indicating no vernalisation, blue mild vernalisation (6 weeks) and red strong vernalisation (12 weeks).
respective morphotypes. The same was done for swedes against non-swede accessions. Choosing a cut-off which considers the top 0.1% of markers (-log[p-value] = 38.9 for winter/spring and 55.8 for swedes), we detected 12 regions on chromosomes A01, A02, A03, A07, A09, A10, C03, C06 and C09 for the winter-spring split (Fig. 4), and 13 regions on chromosomes A03, A04, A06, A09, C01, C08 and C09 for the swede split.

Analysis of split regions. We subsequently counted how many of the 12 winter-spring split regions have a clear winter or spring pattern in each genotype, i.e. the number of cases where every split marker in the haplotype corresponded to the winter or spring state (Table 1). The distribution of lines carrying clear winter and spring haplotypes is shown in Fig. 5. Mixed haplotypes were excluded here, as they account for less than 5% of the haplotypes. From this distribution, we concluded that characterizing these regions for their haplotype pattern is sufficient to distinguish winter from spring morphotypes, but not to distinguish semi-winter or swede morphotypes. The same analysis on 13 split regions identified for the swede vs. non-swede split revealed a more explicit distribution (Table 2 and Fig. 6). Genotyping these loci is therefore sufficient to distinguish swede morphotypes from non-swedes.

In order to exclude candidates for the respective morphotype split, we specifically looked at the variant distributions from deep sequencing in our marker set. Because these derived from sequence data, a poorly fitting distribution excludes the sequence from being a major cause for this morphotype, as sequencing covers the total variation of a gene. This is not the case for data from the SNP array, as even genic SNPs are not always completely predictive for their neighbor SNP. For the winter-spring split, only 6 sequenced regions with a distribution comparable to the detected split markers could be found, among them Bna.VIN3.A02, Bna.FLC.A10 and the Bna.FT promoter on the non-assembled scaffolds of C02_random (Table 1). With the exception of Bna.FLC.A10 (R10P mutation), all those SNPs are either synonymous or located in an intron. For the non-swede vs. swede split, we found 17 sequenced regions carrying variants with an acceptable distribution, for example three copies of Bna.FLC, two copies of Bna.CO and a further copy of Bna.FLOWERING LOCUS D (FD) (Table 2).

Figure 9. Barplots of flowering time recorded in days after sowing (DAS) for FLC-affected and non-affected plants among the spring genotypes in the vernalisation trial. Whiskers show standard errors. The asterisks denote the level of significance for Student’s t-test (*p-value < 0.05, ***p-value > 0.001). NV: No vernalisation, MV: mild vernalisation (6 weeks), SV: strong vernalisation (12 weeks).

In order to exclude candidates for the respective morphotype split, we specifically looked at the variant distributions from deep sequencing in our marker set. Because these derived from sequence data, a poorly fitting distribution excludes the sequence from being a major cause for this morphotype, as sequencing covers the total variation of a gene. This is not the case for data from the SNP array, as even genic SNPs are not always completely predictive for their neighbor SNP. For the winter-spring split, only 6 sequenced regions with a distribution comparable to the detected split markers could be found, among them Bna.VIN3.A02, Bna.FLC.A10 and the Bna.FT promoter on the non-assembled scaffolds of C02_random (Table 1). With the exception of Bna.FLC.A10 (R10P mutation), all those SNPs are either synonymous or located in an intron. For the non-swede vs. swede split, we found 17 sequenced regions carrying variants with an acceptable distribution, for example three copies of Bna.FLC, two copies of Bna.CO and a further copy of Bna.FLOWERING LOCUS D (FD) (Table 2).

Upstream of the gene Bna.FT on C02_random we found two regions, spanning 4622 and 4904 bp, respectively, which retrieved BLAST hits to the A02 or C02 copies of the Bna.FT promoter listed by NCBI. We therefore identified these sequences as the promoter of Bna.FT on C02_random. Both sequences contain a CArG box core motif, whereby the first sequence also contains 3 additional FLC binding sites known from A. thaliana and the second sequence contains 2 such FLC binding sites. No SNP is located in those motifs. We found that most (143 of 145) winter types are unchanged in both sequences or carry only minor changes, whereas most (71 of 116) of the spring population carried one of two distinctive haplotype patterns involving a SNP at position C02_random:980227 (Fig. 7). These patterns were shared by only two putative winter-type accessions, one of which is an exotic accession that may not need vernalisation, whereas the other is an accession which the vernalisation experiments revealed to have vernalisation-independent flowering (see below).

We furthermore compared the numbers of deletion and duplication events in the different morphotype pools. For the winter-spring split, we found no specific pattern for the total population. In contrast, we found several patterns of deletions and duplications which were almost exclusive to swedes, concerning split regions on A08, A09, A10, C08 and C09 (Table 3). The regions on A09/C08 (containing copies of Bna.PHYA and Bna.GIBBERELLIN 3 OXIDASE 1 (GA3ox1)) and A10/C09 (containing copies of Bna.FLC) are homeologous to each other. Some of these regions, particularly those on C08, appear to involve larger homeologous exchanges that probably affect not only the detected genes.
For the winter-spring split, we found 234 genes with flowering-related gene ontology terms within 1 Mb of one of the diagnostic split markers. Examples are shown in Table 4. In the 13 regions showing a split between swedes and non-swarb, we found 260 candidate genes within 1 Mb. In this analysis, several split markers lay directly in candidate genes covered by deep sequencing, for example *Bna.VIN3*, *Bna.PHYA* (2 copies), *Bna.TEMPRA*N*1* (TEM1), *Bna.ELF7*, *Bna.CONSTANS* (CO), *Bna.CO-like* or *Bna.CINNAMOYL COA REDUCTASE 1* (CCR1) (Table 5). Some of those markers were non-synonymous SNPs (in *Bna.CCR1*, *Bna.TEM1*, *Bna.CO*, *Bna.CO-like*), whereas others were either synonymous or located in introns or untranslated regions (UTRs).

**Vernalisation trials.** In order to test if the swede-specific pattern of *Bna.FLC* deletions and duplications would affect vernalisation dependency, we conducted a vernalisation trial with a reduced set of lines (11 lines each from the winter, spring and swede panels). These were selected to represent either lines without CNV in *Bna.FLC* (as a control), or lines which have alternative patterns of deletion and duplication in *Bna.FLC*. The plants were subjected to either 6 or 12 weeks of vernalisation, or were not vernalised. We then scored the time until opening of the first flower. We found that all spring lines and one winter line were vernalisation independent. The vernalisation-independent winter line was one of the two genotypes carrying a strongly divergent *Bna.FT* promoter on C02_random. All swedes and three winter lines were found to be strongly vernalisation dependent, meaning that no plants flowered after mild vernalisation. At the end of the experiment, one winter line and 8 swede lines did not flower at all, meaning that 12 weeks of vernalisation were not sufficient to induce flowering (Fig. 8).

For the spring types, we found that lines with altered *Bna.FLC* patterns flower significantly later than lines without such changes (Fig. 9). All the same, spring lines carrying a swede pattern in *Bna.FLC* were not vernalisation dependent, indicating that this pattern is not sufficient to induce vernalisation.

**Discussion**

Our study aimed at identifying genetic variants which are responsible for the separation of the different morphotype pools in *B. napus*. According to population structure analyses performed in this and other studies,33-34,47, winter-type *B. napus* accessions tend to separate almost completely from other accessions, while some spring types along with semi-winter and swede material are more diverse. Here, we defined a total of 12 variant haplotypes which are diagnostic for the winter-spring split, and 13 variant haplotypes for the swede-non-swarb split. Moreover, we found one winter type without vernalisation requirement that was nevertheless winter-hardy, and spring types with some degree of vernalisation responsiveness. Swedes were found to be extremely vernalisation dependent, with some variation among the accessions, presumably because swede forms have been bred to maintain their vegetative state for as long as possible. Vernalisation is a quantitative process, so there is natural variation in responsive temperature range and vernalisation duration21–25. Markers for such life cycle traits are extremely important in order to introgress desirable traits between ecogeographical or morphotype gene pools, for example seed quality traits from spring to winter oilseed forms23 or resistance traits from swede to non-swarb material51. An R10P mutation in the MADS box domain of *Bna.FLC.A10* was revealed as one candidate for the winter-spring split in *B. napus*, however our data shows that this mutation is neither the only candidate nor the best one. Neither the other *Bna.FLC* homologues, nor the detected copies of *Bna.FRI*, showed an appropriate variant distribution to explain the winter-spring split. Similar results were found for natural variation in flowering time for *A. thaliana*54,55. This excludes the possibility that genetic variation within *Bna.FLC* gene sequences, besides *Bna.FLC.A10*, are causal for vernalisation requirement, thus indicating that other *Bna.FLC* copies are either not responsible for vernalisation or there is variation in cis-regulatory elements. As shown by the vernalisation trials, spring-type plants without CNV in *Bna.FLC* copies flower earlier, indicating that *Bna.FLC* still plays a role in modulating flowering time in the absence of vernalisation requirement. Spring genotypes with a high *Bna.FLC* copy number showed accelerated flowering under vernalisation, indicating that they established weak vernalisation responsiveness. *FLC* is known to bind many other genes in *A. thaliana*65, and it also regulates other developmental processes like germination59, hence additional copies might be assumed to underlie strong selection. The differential degree of conservation between the copies suggests sub-functionalisation, whereby conserved *Bna.FLC.A02* and *Bna.FLC.C02* presumably retain more general roles, whereas *Bna.FLC.A10* might be more specialized towards flowering regulation. Sub-functionalisation events are characteristic for the evolution of MADS box transcription factors57–59.

On the other hand, *Bna.SRR1.A02*, *Bna.VIN3.A02*, *Bna.AGL71.A03*, *Bna.CCR1.A09_random* and the *Bna.FT* promoter on C02_random represent further candidates for the morphotype split. SRR1 is a clock-associated gene found to regulate CO, FT and CYCLING DOF FACTOR 1 (CDF1) in *A. thaliana*59. Moreover, *A. thaliana srr1* mutants have reduced levels of FLC and respond only weakly to vernalisation59. Similarly, *Bna.VIN3* represents a copy of another vernalisation candidate upstream of *Bna.FLC*54. In *Arabidopsis*, *VIN3* is expressed during cold and associates to the PRC2 complex to downregulate FLC gene activity22–24. AGAMOUS-LIKE 71 (AGL71) is closely related to the flowering integrator SUPPRESSOR OF CONSTANS 1 (SOC1) and seems to be involved in gibberellin-dependent flowering pathways61. Its promoter contains a CArG box for FLC binding in *A. thaliana*62. CCR1 is a biosynthetic enzyme in lignin production and leaf development, which is known to regulate the concentration of the antioxidative compound ferulic acid63. This could be related to cold perception, as cold is partly perceived via the redox state64. Although all observed candidate SNPs are synonymous or silent, they may still have strong potential consequences for cis-regulatory elements, methylation, small RNA regulation and chromatin structure, and associated changes in the promoter. Moreover, alternative splicing was found to occur abundantly in resynthesized *B. napus*, also for copies of *Bna.FLC*. On the other hand, we also cannot fully exclude that the observed effects are caused by linkage to additional genes in the neighbourhood of the investigated flowering-time regulators.
We also found patterns on the Bna.FT promoter on C02_random associating with the winter-spring split. Haplotype analysis showed that only two winter lines showed a strongly varying pattern, resembling most of the spring morphotypes. One of these was an exotic line, while the other was found to be vernalisation independent. This indicates that a functional promoter sequence for this gene is necessary to build up vernalisation requirement. A change in vernalisation requirement through variation in an FT promoter was already found in different Brassicas, narrow-leaved lupine and litchi. In A. thaliana, FLC was shown to bind to a CArG box located in the first intron of FT59, although in cereals the binding site lies in the promoter60. Indeed, the Bna.FT copy on C02_random contains a CArG box motif. It is possible that both the promoter and the intron are responsible for Bna.FLC binding. All the same, in both winter and spring types it has been reported that the A02 copy in B. napus is constitutively expressed and the C02 copy is completely silenced, whereas the copies on A07 and C06 appear to be specifically silenced in winter morphotypes but transcribed in spring morphotypes61. As our Bna.FT copy on C02_random corresponds to the C02 copy reported in the aforementioned study62, we assume that either there was a problem with the RT-PCR due to allelic variation, or the regulatory mechanism is more complex. However, in an independent transcriptome study we were unable to detect the constitutive expression of Bna.FT.A02, nor of any other Bna.FT copy, in winter-type B. napus before vernalisation (C. Obermeier, unpublished data).

Genomic rearrangements are common in B. napus63–75. They are particularly predominant in the first generations after allopolyploidisation76, but the process is ongoing and believed to have an important role in speciation77. Different studies found indications for genomic rearrangements between B. napus morphotypes6,12,28. In the present study we found CNVs concerning copies of Bna.FLC, Bna.PHYA and Bna.GA3ox1 to involve duplications in the A subgenome and corresponding homoeologous deletions in the C subgenome. This indicates replacement of the C-subgenome regions by the respective A-subgenome regions, a process known as homoeologous non-reciprocal translocation (HNRT)77. A de novo HNRT will erase any sub-functionalisation which may have occurred prior to the rearrangement. Our data concur with the hypothesis27 that the Bna.FLC.A10 copy is most specifically involved in flowering regulation. A duplication in Bna.FLC.A10 would therefore increase vernalisation requirement. This hypothesis fits with the strong vernalisation requirement we observed in lines carrying this duplication. Differential expression of Bna.FLC in highly rearranged, resynthesized rapeseed was observed before26. All the same, this pattern can only be effective when the vernalisation system is functional, as two spring lines with the same pattern are not vernalisation dependent. One of these presumably has a defective Bna.FT promoter on C02_random, whereas neither of them carries the swede-specific Bna.VIN3.A03 marker. Other genes affected by such HNRTs are Bna.PHYA and Bna.GA3ox1. PHYA is a red/ far-red perceiving photoreceptor which has a stabilising role for CO under long days78. This might represent a necessary co-adaptation of the photoperiodic pathway due to the strong vernalisation requirement, as later flowering means that the day length is longer at the time of flowering. This assumption is underlined by the finding that a D94G mutation in a copy of Bna.CO-like is a candidate for the swede split. GA3ox1, a biosynthetic key gene involved in GA production, is regulated by PHYB and by feedback mechanisms of downstream pathways79. GA also affects other developmental processes like seed germination, hypocotyl elongation and fruit set80. Bna.GA3ox1 is therefore also a candidate for the swede morphotype, which is characterized by an enlarged hypocotyl and low seed-set. This might also apply to Bna.CCR1 as a candidate for the swede split. In Arabidopsis CCR1 is involved in lignin biosynthesis, leaf development regulation and regulation of the redox state81 (see above).

All swede lines share a silent mutation in Bna.VIN3.A03, which is not shared by any other line. Although the consequence of this mutation is unclear, Bna.VIN3 is a strong candidate for vernalisation requirement, particularly because another copy is a candidate for the winter-spring split (see above). Copies of VIN3 have previously been named as candidates for vernalisation requirement and flowering time in A. thaliana and B. napus82,83. In B. oleracea, it was found that BoVIN3 was upregulated much faster than A. thaliana VIN383, indicating that the expression is more sensitive to cold. Another upstream candidate for the strong vernalisation requirement is Bna.ELF7. ELF7 is involved in chromatin remodeling of FLC during vernalisation84. A further gene variant which was not found outside the swede population lay in Bna.TEM1. TEM1 encodes another repressor of FT, which competes with CO for the same genetic region to fulfill their function85. The variant is a non-synonymous T167R mutation that potentially affects binding to the UTR of FT. A further candidate, Bna.FD, possibly modulates FT protein effectiveness, as FD is a direct and essential interaction partner of FT in the shoot apex86.

These results represent an excellent base for further experiments to transfer morphotype features between B. napus genetic pools. Moreover, they also shed light on the evolution of major flowering time genes in the aftermath of allopolyploidisation, and their role in morphotype diversification and ecogeographical adaptation. We clearly demonstrate that different copies of important flowering regulators play different regulatory roles across the vernalisation and flowering pathways. The scarcity of non-synonymous mutations, along with the observed variation in the Bna.FT promoter, underline the importance of cis-regulatory mechanisms in flowering time regulation.

References
1. Adams, K. L. & Wendel, J. F. Polyploidy and genome evolution in plants. Curr Opin Plant Bio 8, 135–141 (2005).
2. Solits, P. S. & Solits, D. E. The role of hybridization in plant speciation. Annu. Rev. Plant Biol. 60, 561–588 (2009).
3. Arrigo, N. & Barker, M. S. Rarely successful polyploids and their legacy in plant genomes. Curr Opin Plant Bio 15, 140–146 (2012).
4. Brenchley, R. et al. Analysis of the bread wheat genome using whole-genome shotgun sequencing. Nature 491, 705–710 (2012).
5. The Potato Genome Sequencing Consortium. Genome sequence and analysis of the tuber crop potato. Nature 475, 189–195 (2011).
6. Chalhoub, B. et al. Early allopolyploid evolution in the post-Neolithic Brassica napus oilseed genome. Science 345, 950–953 (2014).
7. Edwards, Dave, Butley, Jacqueline, Parkin, Isobel & Chittaranjan, Kole (ed.) Genomics and Breeding of Oilseed Brassicas (CRC Press, 2011).
8. Bundessortenamt. Beschreibende Sortenliste 2016. Getreide, Mais Öl- und Faserpflanzen Leguminosen Rüben Zwischenfrüchte (2016).
9. Allender, C. J. & King, G. J. Origins of the amphiphilic species Brassica napus L. investigated by chloroplast and nuclear molecular markers. BMC Plant Biol 10, 54 (2010).
59. Dittmar, E. L., Oakley, C. G., Agren, J. & Schemske, D. W. Flowering time QTL in natural populations of Bromus tectorum. Front. Plant. Sci. 5, 404 (2014).
60. Coustham, V. 
61. Meyer, S. E., Nelson, D. L. & Carlson, S. L. Ecological genetics of vernalization response in Arabidopsis thaliana. Mol Ecol 23, 1294–1296 (2014).
62. Díaz, A., Schiessl, S., Iniguez-Luy, F., Qian, W. & Snowdon, R. J. Diverse regulatory factors associate with flowering time and yield responses in maize. Mol Ecol 23, 757–772 (2014).
63. Schmutzer, T. et al. The FRIGIDA complex activates transcription of FLC, a strong flowering repressor in Arabidopsis, by recruiting chromatin modification factors. The Plant Cell 23, 289–303 (2011).
64. Bus, A., Hecht, J., Huettel, B., Reinhardt, R. & Stich, B. High-throughput polymorphism detection and genotyping in Brassica napus using next-generation RAD sequencing. BMC Genomics (2012).
65. Corber, N. E. et al. Seedling development in a Brassica napus diversity set and its relationship to agronomic performance. Theor Appl Genet 125, 1275–1287 (2012).
66. Quijada, P. A., Udall, J. A., Lambert, B. & Osborn, T. C. Quantitative trait analysis of seed yield and other complex traits in hybrid spring rapeseed (Brassica napus L.). 137, 537–554 (2005).
67. Nelson, M. N. et al. Quantitative trait loci for thermal time to flowering and photoperiod responsiveness discovered in summer annual-type Brassica napus L. PLoS ONE 9, e102611 (2014).
68. Raman, H. et al. Genetic and physical mapping of flowering time loci in canola (Brassica napus L.). Theor Appl Genet 126, 119–132 (2013).
69. Fletcher, R. S., Mullen, J. L., Heiliger, A. & McKay, I. K. QTL analysis of root morphology, flowering time, and yield reveals trade-offs in response to drought in Brassica napus. Ex Bot J (2014).
70. Bus, A., Hecht, J., Huettel, B., Reinhardt, R. & Stich, B. High-throughput polymorphism detection and genotyping in Brassica napus using next-generation RAD sequencing. BMC Genomics (2012).
71. McKenna, A. et al. The genome analysis toolkit: a mapreduce framework for analyzing next-generation DNA sequencing data. Genome Res 20, 1297–1303 (2010).
72. Danecek, P. et al. The variant call format and VCFTools. Bioinformatics 27, 2156–2158 (2011).
73. Schiessl, S., Iniguez-Luy, F., Qian, W. & Snowdon, R. J. Diverse regulatory factors associate with flowering time and yield responses in winter-type Brassica napus. BMC Genomics 16, 737 (2015).
74. Aulchenko, Y. S., Ripke, S., Isaacs, A. & van Duijn, C. M. GenABEL: an R library for genome-wide association analysis. Bioinformatics 23, 1324–1326 (2007).
75. Deng, W. et al. FLOWERING LOCUS C (FLC) regulates developmental pathways throughout the life cycle of Arabidopsis. PNAS 108, 6680–6685 (2011).
76. Bus, A., Corber, N., Snowden, R. J. & Stich, B. Patterns of molecular variation in a species-wide germplasm set of Brassica napus. Theor Appl Genet 123, 1433–1437 (2011).
77. Song, J., Irwin, J. & Dean, C. Remembering the prolonged cold of winter. Current Biology 23, R807–R811 (2013).
78. Duncan, S. et al. Seasonal shift in timing of vernalization as an adaptation to extreme winter. elife (2015).
79. Coustham, V. et al. Quantitative modulation of polycomb silencing underlies natural variation in vernalization. Science 337, 584–587 (2012).
80. Meyer, S. E., Nelson, D. L. & Carlson, S. L. Ecological genetics of vernalization response in Bromus tectorum L. (Poaceaee). Ann Bot 93, 653–663 (2004).
81. Wu, G., Wu, Y., Xiao, L., Li, X. & Lu, C. Zero erucic acid trait of rapeseed (Brassica napus L.) results from a deletion of four base pairs in the fatty acid elongase 1 gene. Theor Appl Genet 116, 491–499 (2008).
82. Hasan, M. J. & Rahman, H. Genetics and molecular mapping of resistance to Plasmopara brassicae pathotypes 2, 3, 5, 6, and 8 in rapeseed as a whole genome panel. Is Caronav. 50, 805–815 (2016).
83. Dittmar, E. L., Oakley, C. G., Agren, J. & Schemske, D. W. Flowering time QTL in natural populations of Arabidopsis thaliana and implications for their adaptive value. Mol Ecol 23, 4291–4303 (2014).
84. Grillo, M. A., Li, C., Hammond, M., Wang, L. & Schemske, D. W. Genetic architecture of flowering time differentiation between locally adapted populations of Arabidopsis thaliana. New Phytol 197, 1321–1331 (2013).
85. Chiang, G. K. C., Barua, D., Kramer, E. M., Amin, R. M. & Donohue, K. Major flowering time gene, FLOWERING LOCUS C, regulates seed germination in Arabidopsis thaliana. PNAS 106, 11661–11666 (2009).
86. Becker, A. The major clades of MADS-box genes and their role in the development and evolution of flowering plants. Mol Phylogen Evol 29, 464–489 (2003).
87. Dreni, L. & Kater, M. M. MADS reloaded: evolution of the AGAMOUS subfamily genes. New Phytol 201, 717–732 (2014).
88. Johannson, M. & Staal, D. SSR1 is essential for repressing flowering in non-inductive conditions in Arabidopsis thaliana. Ex Bot J 65, 5811–5822 (2014).
60. Wood, C. C. et al. *Arabidopsis thaliana* vernalization response requires a polycomb-like protein complex that also includes VERNALIZATION INSENSITIVE 3. PNAS 103, 14631–14636 (2006).

61. Dorca-Fornell, C. et al. The *Arabidopsis* SOC1-like genes AGL42, AGL71 and AGL72 promote flowering in the shoot apical and axillary meristems. *Plant J.* 67, 1006–1017 (2011).

62. Hellwell, C. A., Wood, C. C., Robertson, M., James Peacock, W. & Dennis, E. S. The *Arabidopsis* FLC protein interacts directly in vivo with SOC1 and FT chromatin and is part of a high-molecular-weight protein complex. *Plant J.* 46, 183–192 (2006).

63. Xue, J. et al. CCR1, an enzyme required for lignin biosynthesis in *Arabidopsis*, mediates cell proliferation exit for leaf development. *Plant J.* 83, 375–387 (2015).

64. Kovi, M. R., Ergon, A. & Rognli, O. A. Freezing tolerance revisited: effects of variable temperatures on gene regulation in temperate grasses and legumes. *Curr Opin Plant Bio.* 33, 140–146 (2016).

65. Zhou, R., Moshtaghi, N. & Adams, K. L. Extensive changes to alternative splicing patterns following allopolyploidy in natural and resynthesized polyploids. *PNAS* 108, 16122–16127 (2011).

66. Wang, J. et al. Promoter Variation and Transcript Divergence in Brassicaceae Lineages of FLOWERING LOCUS T. *PLoS ONE* 7, e47127 (2012).

67. Nelson, M. N. et al. The loss of vernalization requirement in narrow-leafed lupin is associated with a deletion in the promoter and de-repressed expression of a *FLOWERING LOCUS T* (*FT*) homologue. *New Phytol.* (2016).

68. Ding, F. et al. Promoter difference of *LaFT1* is a leading cause of natural variation of flowering timing in different *litchi* cultivars (*Litchi chinensis* Sonn.). *Plant science* 241, 128–137 (2015).

69. Searle, J. et al. The transcription factor FLC confers a flowering response to vernalization by repressing meristem competence and systemic signaling in *Arabidopsis*. *Genes & Dev.* 20, 898–912 (2006).

70. Deng, W. et al. Direct links between the vernalization response and other key traits of cereal crops. *Nat Comms* 6, 5882 (2015).

71. Clarke, W. E. et al. A high-density SNP genotyping array for *Brassica napus* and its diploid diploid species based on optimised selection of single-locus markers in the allotetraploid genome. *Theor Appl Genet* 129, 1887–1899 (2016).

72. Gaeta, R. T., Pires, J. C., Insiguez-Luy, E., Leon, E. & Osborn, T. C. Genomic Changes in Resynthesized *Brassica napus* and Their Effect on Gene Expression and Phenotype. *The Plant Cell* 19, 3403–3417 (2007).

73. Nicolas, S. D., Monod, H., Eber, F., Chevre, A. M. & Jenczewski, E. Non-random distribution of extensive chromosome rearrangements in *Brassica napus* depends on genome organization. *Plant J.* 79, 991–703 (2012).

74. Udall, J. A., Quijada, P. A. & Osborn, T. C. Detection of Chromosomal Rearrangements Derived From Homoeologous Recombination in Four Mapping Populations of *Brassica napus* L. *Genetics* 169, 967–979 (2005).

75. Szadkowski, E. et al. Polyploid formation pathways have an impact on genetic rearrangements in resynthesized *Brassica napus*. *New Phytol.* 191, 884–894 (2011).

76. Faria, R. & Navarro, A. Chromosomal speciation revisited: rearranging theory with pieces of evidence. *Trends ecol evol* 25, 660–669 (2010).

77. Gaeta, R. T. & Pires, J. C. Homoeologous recombination in allopolyploids: the polyploid ratchet. *New Phytol.* 186, 1–8 (2010).

78. Valverde, F. et al. Photoreceptor regulation of CONSTANS protein in photoperiodic flowering. *Current Biology* 18, 1338–1343 (2008).

79. Gallego-Giraldo, C. et al. Role of the gibberellin receptors GID1 during fruit-set in *Arabidopsis*. *Plant J.* 79, 1020–1032 (2014).

80. Shi, J. et al. Unraveling the Complex Trait of Crop Yield With Quantitative Trait Loci Mapping in *Brassica napus*. *Genetics* 182, 851–861 (2009).

81. Ridge, S., Brown, P. H., Hecht, V., Driessen, R. G. & Weller, J. L. The role of *BoFBC2* in cauliflower (*Brassica oleracea var. botrytis* L.) reproductive development. *Ex Bot J.* 166, 125–135 (2015).

82. He, Y., Doyle, M. R. & Amasino, R. M. PAF1-complex-mediated histone methylation of *FLOWERING LOCUS C* chromatin is required for the vernalization-responsive, winter-annual habit in *Arabidopsis*. *Genes & Dev.* 18, 2774–2784 (2004).

83. Castilho, C. & Pelaz, S. The Balance between CONSTANS and TEMPRANILLO Activities Determines FT Expression to Trigger Flowering. *Current Biology* 18, 1338–1343 (2008).

84. Abe, M. et al. FD, a bZIP Protein Mediating Signals from the Floral Pathway Integrator FT at the Shoot Apex. *Science* 309, 1052–1056 (2005).

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**Author Contributions**

S.S. and R.S. planned the experiments and wrote the manuscript. S.S. performed DNA extraction, data analysis and phenotyping. B.H., D.K. and R.R. performed library preparation and sequencing. SS and RS prepared the manuscript.

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