Quantitative Control of Early Flowering in White Lupin (Lupinus albus L.)

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Abstract: White lupin (Lupinus albus L.) is a pulse annual plant cultivated from the tropics to temperate regions for its high-protein grain as well as a cover crop or green manure. Wild populations are typically late flowering and have high vernalization requirements. Nevertheless, some early flowering and thermoneutral accessions were found in the Mediterranean basin. Recently, quantitative trait loci (QTLs) explaining flowering time variance were identified in bi-parental population mapping, however, phenotypic and genotypic diversity in the world collection has not been addressed yet. In this study, a diverse set of white lupin accessions (n = 160) was phenotyped for time to flowering in a controlled environment and genotyped with PCR-based markers (n = 50) tagging major QTLs and selected homologs of photoperiod and vernalization pathway genes. This survey highlighted quantitative control of flowering time in white lupin, providing statistically significant associations for all major QTLs and numerous regulatory genes, including white lupin homologs of CONSTANS, FLOWERING LOCUS T, FY, MOTHER OF FT AND TFL1, PHYTOCHROME INTERACTING FACTOR 4, SKI-INTERACTING PROTEIN 1, and VERNALIZATION INDEPENDENCE 3. This revealed the complexity of flowering control in white lupin, dispersed among numerous loci localized on several chromosomes, provided economic justification for future genome-wide association studies or genomic selection rather than relying on simple marker-assisted selection.

Keywords: vernalization; markers; flowering; quantitative trait; selection

1. Introduction

White lupin (Lupinus albus L.) is an annual legume plant cultivated for animal feed and human consumption in Europe, Africa and Australia [1]. It is also a valuable component in crop rotation and organic farming, thanks to its nitrogen fixation via diazotrophic symbiosis and mobilization of soil phosphorus by formation of cluster roots [2–4]. White lupin seeds are a valuable source of protein (38–42% dry weight) and oil (10–13%) with advantageous omega-6 to omega-3 acid ratios [5–7]. Moreover, white lupin consumption has positive nutraceutical impact, reducing hyperglycaemia, hypercholesterolemia and hypertension [8]. For many years, wide use of lupins for human consumption was hampered by their bitter taste caused by a high alkaloid content, but this issue has been fixed in modern cultivars [9]. Considering the agronomic characteristics of white lupin, there is still high potential for further yield increases and drought tolerance improvement by exploitation of existing germplasm resources [10–13].

It is assumed that white lupin originated from the Balkan-Mediterranean area [14]. Native adaptation to environmental conditions of many lupin landraces is the requirement of a prolonged cold period, named vernalization, to initiate flowering. Thus, in late flowering and vernalization-responsive lines, time to flowering is reduced proportionally...
to the length of the vernalization period [15]. White lupin is currently grown under a wide range of climatic conditions, including humid tropics, the Mediterranean basin as well as temperate zones with humid or intermediate continental climates. In some of these regions highly delayed flowering in the absence of vernalization is a very undesired trait.

Moreover, white lupin was revealed to be susceptible to anthracnose, a very devastating disease first reported in Brazil in 1912, then in the USA in 1939, and, finally, until 1997, it appeared in all major regions where white lupin was cultivated [16–19]. Unfortunately, major European early flowering white lupin donors were found to be very susceptible to anthracnose [20,21]. Substantial levels of resistance to anthracnose were found only in several accessions originating from mountainous regions of Ethiopia, particularly in lines P27174, P27175, and P27178 [22]. Marker-based studies revealed that Ethiopian white lupins are relatively closely related to each other and very distinct from the improved germplasm originating from Europe [23]. A reference anthracnose resistance donor, line P27174, was found to carry two major undesired traits, namely a high vernalization requirement and high alkaloid content, and as such was exploited, together with Kiev Mutant carrying opposite traits, for the development of mapping populations and the construction of the linkage map [24]. Following progress in molecular techniques, this linkage map was subsequently improved with novel markers [25,26]. The most recent version of the linkage map served as an anchor for the establishment of two genome sequence assemblies [27,28]. Quantitative trait loci (QTL) mapping, based on the same mapping population, revealed the presence of two major QTLs for anthracnose resistance, located in the linkage groups ALB02 and ALB04, as well as five major QTLs for flowering time, two of them located in the linkage group ALB02 (including one overlapping with the major anthracnose resistance locus), another two in the linkage group ALB13, and one in the linkage group ALB16 [24–26]. The white lupin molecular toolbox has been recently supplemented with PCR-based markers suitable for tracking Ethiopian alleles of major anthracnose resistance loci, alkaloid locus pauper and main candidate genes controlling flowering induction [29–31]. Interestingly, white lupin breeders revealed that joining of early flowering with anthracnose resistance in the progeny descending from the cross between Kiev Mutant and P27174 was ineffective, however, replacement of the Ukrainian donor of early flowering by the other (French) thermoneutral germplasm made this process much more feasible [22,32]. Such an observation highlighted the hypothetical presence of additional genes controlling early flowering in white lupin collection, genetically different than those revealed in population mapping.

Indeed, the lack of knowledge on distribution of alleles controlling flowering time in worldwide white lupin germplasm collection hampers selection of compatible early flowering germplasm for crosses in current breeding approaches. The present study aimed to address this issue by combining phenotyping of time to flowering in diversified germplasm collection with genotyping of marker alleles across all major QTL loci conferring flowering time control as well as some candidate genes from flowering regulation pathways. This approach provided novel evidences for quantitative regulation of flowering induction in white lupin and designated novel candidate donors of earliness and thermoneutrality for further improvement of this species as a crop.

2. Results

2.1. Early Flowering and Thermoneutrality Is Present in Primitive and Domesticated Germplasm

A white lupin germplasm collection (Supplementary Table S1) was phenotyped in greenhouse conditions for time from sowing to flowering without vernalization in three years, 2015, 2018 and 2020. In two years—2015 and 2018—vernalization responsiveness was evaluated as well. Observation data is provided in Supplementary Table S2, whereas calculated mean values with standard deviation are given in Supplementary Table S3. The number of days from sowing to flowering in the absence of vernalization revealed continuous distribution with two major peaks and high variability between genotypes (Figure 1). Observed values ranged from 37.8 ± 0.4 to 90.8 ± 6.5 in 2015 (mean value of
53.9), from 37.1 ± 2.3 to 90.0 ± 0.0 in 2018 (mean value of 52.0) and from 47.2 ± 2.1 to 89.5 ± 1.5 in 2020 (mean value of 63.9).

Figure 1. Distribution of the mean number of days from sowing to flowering in white lupin germplasm collection in years 2015, 2018 and 2020. Plants were cultivated in greenhouse without pre-sowing vernalization. Rectangles visualize mean and standard deviation values.

The general trend of observed variation in time to flowering followed differences in domestication status (Table 1). Improved germplasm (cultivars and cross derivatives) flowered about 3.9–11.1 days earlier than landraces and wild or primitive lines. Mutants were revealed to flower 13.4–18.6 days later than domesticated accessions. Vernalization response was the highest in mutants (−18.3 ± 13.3 and −20.2 ± 13.3 days in 2015 and 2018 experiments, respectively), intermediate in wild/primitive germplasm (−9.8 ± 6.2 and −15.0 ± 7.4) and in landraces (−7.6 ± 6.9 and −14.4 ± 9.6), whereas the lowest in cultivars (−5.2 ± 5.9 and −6.6 ± 3.9) and in cross derivatives (−3.1 ± 2.0 and −7.4 ± 4.1). Interestingly, three primitive accessions from Turkey, namely 95523 “FRA6708B”, 95524 “FRA6712B” and 95525 “FRA6713B”, annotated in Polish database as “wild or primitive” and in Australian database as “landraces”, revealed the earliest time to flowering and full thermoneutrality. Such an observation indicates that early flowering based on thermoneutrality have putatively appeared in white lupin also in primitive germplasm, independently to selective breeding during recent domestication process.

Table 1. Mean number of days from sowing to flowering observed in white lupin germplasm in years 2015, 2018 and 2020.

| Type               | 2020 n 1 | 2018 n | 2018 v 2 | 2015 n | 2015 v |
|--------------------|----------|--------|----------|--------|--------|
| wild or primitive  | 67.4 ± 8.6 3 | 54.3 ± 8.2 | 39.4 ± 2.7 | 56.6 ± 8.8 | 46.8 ± 6.2 |
| landrace           | 64.7 ± 9.3  | 53.5 ± 11.3 | 38.8 ± 2.8 | 53.6 ± 9.4 | 46.0 ± 4.3 |
| mutant             | 74.5 ± 8.6  | 59.5 ± 14.4 | 39.4 ± 1.5 | 65.1 ± 15.5 | 46.8 ± 2.3 |
| cultivar           | 56.3 ± 6.2  | 44.5 ± 6.6  | 38.0 ± 5.5 | 49.7 ± 10.4 | 42.7 ± 2.8 |
| cross derivative   | 58.5 ± 5.8  | 46.1 ± 5.1  | 38.7 ± 3.2 | 46.4 ± 3.6  | 43.3 ± 2.1 |

1 Non-vernalized plants; 2 Vernalized plants; 3 Standard deviation.

2.2. Flowering Induction in White Lupin Germplasm Collection Is under Polygenic Control

In this study, 17 new PCR-based markers were developed using GBS reads aligned to white lupin transcriptome [26,33] and genome [27] assemblies. These markers originated from the flowering time QTLs located in the linkage group ALB02 at position around 100.2/100.5 cM (QTL2, 6 markers), the linkage group ALB13 around 81.0/84.1 cM (QTL3, 4 markers) and 96.2/99.3 cM (QTL4, 3 markers), and the linkage group ALB16 at 0.9/2.2 cM (QTL5, 4 markers). Taking into consideration detection methods, the were 11 CAPS and 6 dCAPS markers developed (Table 2).
Table 2. New molecular markers developed in this study to supplement marker-assisted tracking of flowering time QTLs in white lupin germplasm collection.

| Marker Name | QTL | Primers (5′→3′) | Detection Method | Products Kiev Mutant (bp) | Products P27174 (bp) |
|-------------|-----|-----------------|-----------------|--------------------------|----------------------|
| TP56963     | 2   | TGCTCGAATGCCCCAAATCCATCA TTGATGCTTCGAGTTGAAGGATAA | CAPS, HinfI | 878, 384 | 878, 304, 80 |
| TP235608    | 2   | GTAGCCAAACATGAGAGCAG TCACTGCAATCTT GTCAATCTT | CAPS, AflII | 217 | 179.38 |
| TP94353     | 2   | CAGCATTTATGTTGAGGACA GGAACCTTCGAATTTGATAAAGG | CAPS, Rsal | 60, 51 | 111 |
| TP278885    | 2   | CCATTGGATAGCTGCAAATGCCTTCG CCTTGGATGTTGAAACCTATGC | dCAPS, HpaII | 112 | 86, 26 |
| TP115697    | 2   | TGCTGCCGTATTATGGATCTACA TGAAATTTAGCAAAACTCAGTGTA | dCAPS, Rsal | 111, 24 | 135 |
| TP114357    | 2   | GCCATTCTGGATGGATAACCG TGGAACACCATGCTGACCTTCA | dCAPS, HpaII | 124 | 105, 19 |
| TP100150    | 3   | TATTTCAGCCAATCCCATCCTCT ACCTTCCCATCCATCTTGGACGA | CAPS, HpyCH4V | 87, 38, 30, 5 | 117, 38, 35 |
| TP288840    | 3   | CGCAATTATATCTTAAAGACCATTGT CGTAATCTATATAATGTTT | dCAPS, MboI | 60 | 37, 23 |
| TP3177      | 3   | CGTCGACAACTGTTCACAGG ATCTGGTTGGAAGCTTGGTGGT | CAPS, SspI | 169 | 114, 55 |
| TP360542    | 3   | GAGCCAGAGATAAAGGGTTGGT ACTGGATAGTAAAGACCCCATAGATTACT | dCAPS, TaqI | 113 | 82, 31 |
| TP345457    | 4   | CACAATTCCTACTCACAGATCACC GATTCTCGTCCATCCAGATCTTCTC | CAPS, BseDI | 229, 39, 12 | 143, 84, 39, 12 |
| TP11750     | 4   | AAAACACTGAAAAAGCTTCAACA CAGGGGATAATAAATCTCCATCCA | CAPS, Acl | 209 | 135, 74 |
| TP402859    | 4   | CTGGTGGAAGAAGGCAAGAA AAGGCCAGAAAGCACATTTG | CAPS, HpaII | 198 | 112, 86 |
| TP2488      | 5   | ACCTTTGATTATGATGCTAGCTTCT TGTTAGGAGGCAGCTTGGAAAT | dCAPS, TspE1 | 48, 25, 21 | 48, 46 |
| TP86766     | 5   | CAGCATCGCAAGAAGCCTG TCCTTCCCTCTTCTTCTTCTTT | CAPS, DdeI | 64 | 48, 16 |
| TP47264     | 5   | TAACCATGCACGACTACCAAC TCTGGTTTGGTTAGGTAATGAGGA | CAPS, MboI | 171 | 105, 66 |
| TP30473     | 5   | CAGCACACACAGCGGAATAC TAATACCAGGAATATATGCTTGGT | CAPS, HpyCH4V | 28,25 | 53 |

Together with the previously published markers \[26,29,30\], the final set used for genotyping of white lupin germplasm collection consisted of 50 markers which enabled us to track all major QTLs of flowering time and 29 white lupin homologs of flowering induction pathway genes (Supplementary Tables S4 and S5). Novel alleles (not present in the parental lines of mapping population) were identified for markers FTc1-F4 (16 lines), CO-F1 (2 lines), TP56963 (two lines), MFT-FT3-F1 (one line), and TP114357 (one line). All markers were polymorphic in the analyzed white lupin germplasm collection, however, minor allele frequency (MAF) varied from 0.6% to 48.8% (Supplementary Table S6). Twenty four markers had MAF below 25%, including six markers with MAF below 5% (FRI-F1, ESD4-F8, Fta1-F2, ESD4-F7, SKIP1-F2, and FKF-F2M). Interestingly, two of these markers, Fta1-F2 and SKIP1-F2, were localized in two major QTLs of white lupin flowering time. Moreover, in markers with MAF up to 25.2%, minor alleles were always originating from
the P27174 line. Indeed, in case of several markers, very low frequency of wild allele could make it statistically impossible to find significant association between the genotype and the phenotype in germplasm collection despite differences between flowering time in lines carrying opposite alleles. Nevertheless, an analysis of marker polymorphism pattern in germplasm collection revealed the presence of significant correlation between phenotypes and genotypes for 19 markers (Figure 2). This set included markers tagging all major white lupin QTLs of flowering time and 11 homologs of genes involved in regulation of flowering induction, namely CONSTANS-like (CO, marker CO-F1), EARLY FLOWERING 1 (ELF1, marker ELF1-F1), FLOWERING LOCUS D (FLD, marker FLD-F1), FLOWERING LOCUS T (FTc1, marker FTc1-F4), FY (marker FY-F6), LUMINIDEPENDENS (LD, marker LD-F1), MOTHER OF FT AND TFL1 (MFT, marker MFT-F3-F1), PHYTOCHROME INTERACTING FACTOR 4 (PIF4, marker PIF4-F6), SEPALLATA 3 (SEP3, marker SEP3-F1), SKI-INTERACTING PROTEIN 1 (SKIP1, marker SKIP1-F2), and VERNALIZATION INDEPENDENCE 3 (VIP3, marker VIP3-F2). The highest additive effects were calculated for markers SKIP1-F2 (7.5–9.4), MFT-F3-F1 (4.0–6.6), TP235608 (3.1–7.9), and FTc1-F4 (3.3–6.2).

Figure 2. Correlation between the phenotype (the number of days from sowing to flowering) and the genotype (marker segregation) in white lupin germplasm collection. Color scale visualizes Spearman’s rank correlation coefficient values (Corr). Abbreviations are as follows: LG, linkage group; POS; position in the linkage group in centimorgans; QTL, the number of assigned QTL; AE, additive effect (See Supplementary Table S4). *** p value <0.001; ** p value < 0.01; * p value < 0.05; + p value < 0.1.

2.3. Several White Lupin Subpopulations Differing in Flowering Time and Allelic Composition Were Identified

Data on PCR-based marker polymorphisms obtained for 160 white lupin accessions were exploited for population structure analysis. Minimal cross-entropy values were revealed for six clusters (ancestral populations), followed by a second minimum at four clusters (Figure 3). Based on the results of these calculations, white lupin germplasm collection was divided into six subpopulations. These clusters differed in number of

![Figure 2](https://example.com/figure2)

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accessions, namely five in group V1, 32 in V2, 20 in V3, 53 in V4, 32 in V5, and 18 in V6 (Supplementary Table S1).

![Figure 3. Cross-entropy values obtained for a given number of clusters (ancestral populations).](image)

Molecular analysis of variance (AMOVA) supported the hypothesis that such a grouping into subpopulations is not random (simulated p-value: 9.999 × 10^{-5}). Membership of all genotypes to six subpopulations were visualized by the ancestry matrix (Figure 4).

![Figure 4. Ancestry matrix constructed for six white lupin subpopulations (n = 160).](image)

Plants in these subpopulations showed the following mean values of number of days from sowing to flowering: V1, 64.5 ± 9.5 days; V2, 62.8 ± 12.0; V3, 57.7 ± 9.0; V4, 54.9 ± 10.8; V5, 53.9 ± 8.9; and V6, 55.8 ± 11.5 days. There were significant differences between years (p-value < 2 × 10^{-10}) and groups (p-value 3.7 × 10^{-11}) and no significant interaction between years and groups (p-value 0.996) (Figure 5).

Observed differences in flowering time of early lines between the year 2020 and years 2015 or 2018 may result from the differences in the number of days with maximum temperature above 20 °C recorded during the first 35 days of the experiments, reaching 13 days in 2015, 14 days in 2018 and 4 days in 2020. As the greenhouse was equipped with cooling based only on window opening/closure, high outside temperature directly translated into warmer conditions inside the greenhouse. It is the well-known fact that higher temperature accelerates plant growth and development.

Then, we tested the correlation between distribution of marker polymorphism and observed time to flowering within subpopulations (Supplementary Table S7). Thirteen markers revealed significant correlation (p value ≤ 0.05) with time to flowering in at least one year in one group, nine in two groups and two in three or four groups. This set included one marker from the QTL1, four markers from the QTL2, two markers from the QTL3, five markers from the QTL5, and ten markers anchored in sequences of genes which have not been assigned to any of the QTLs.
Taking into consideration the number of years, ten markers revealed significant correlation ($p$-value $\leq 0.05$) with time to flowering in at least two years within a particular group, including five markers significantly correlated across all three years (Table 3). One marker, TP86766 derived from the QTL5, was significantly correlated with time to flowering in two subpopulations (V4 and V5) across all three years of observations.

**Table 3.** Correlation between the phenotype (the number of days from sowing to flowering) and the genotype (marker segregation) in subpopulations of white lupin germplasm collection.

| Cluster | Marker | Linkage Group | Position (cM) | QTL | 2015 $p$-Value $^1$ | 2018 $p$-Value | 2020 $p$-Value |
|---------|--------|---------------|---------------|-----|---------------------|----------------|----------------|
| V2      | TP56963 | ALB02         | 96.4          | 2   | 0.048 $^3$          | 0.021          | 0.017          |
| V2      | TP235608| ALB02         | 96.8          | 2   | 0.111               | 0.048          | 0.018          |
| V2      | PIF4-F6 | ALB04         | 7.3           | -   | 0.021               | 0.034          | 0.017          |
| V4      | TP86766 | ALB16         | 2.2           | 5   | 0.039               | 2.8 $\times 10^{-4}$ | 4.0 $\times 10^{-3}$ |
| V4      | TP30473 | ALB16         | 6.7           | 5   | 0.044               | 0.019          | 0.003          |
| V5      | FTC1-F4 | ALB14         | 53.9          | -   | 0.051               | 0.016          | 0.008          |
| V5      | TP86766 | ALB16         | 2.2           | 5   | 0.034               | 0.020          | 0.003          |
| V6      | MFTa1-F1| ALB05         | 68.1          | -   | 0.039               | 0.032          | 0.023          |
| V6      | CO-F1   | ALB10         | 15.1          | -   | 0.027               | 0.009          | 0.012          |
| V6      | FRI31-F1| ALB16         | 5.8           | 5   | 0.003               | 0.087          | 0.016          |
| V6      | TP2488  | ALB16         | 0.3           | 5   | 0.062               | 0.029          | 0.039          |

$^1$ $p$-value of Spearman’s rank correlation coefficient; $^2$ Only markers showing significant correlation with time to flowering in at least two years within a particular group were listed here; $^3$ Color scale was used to highlight the statistical significance of obtained values as follows: red, $p \leq 0.001$; orange, $p \leq 0.01$; yellow, $p \leq 0.05$; green, $p \leq 0.1$; white, $p > 0.05$.

Tukey’s tests supported the hypothesis of lack of difference in flowering time between clusters obtained in the population structure analysis for pairs V2-V1, V3-V1, V4-V3, V5-V3, V6-V3, V5-V4, V6-V4, V6-V5, and discarded such a hypothesis for pairs V4-V1, V5-V1, V6-V1, V3-V2, V4-V2, V5-V2, and V6-V2. Therefore, for subpopulation pairs which in Tukey’s test analysis showed a difference in flowering time, $\chi^2$ analysis was performed to test if differences in allele frequencies are significant or not. Between groups V1 and V4 differences in allele frequencies were significant for 16 markers, between V1 and V5 for 21 markers, between V1 and V6 for 11 markers, between V2 and V3 for 17 markers, between V2 and V4 for 16 markers, between V2 and V5 for 19 markers and between V2 and V6 for nine markers.
No marker revealed significant differences in allele frequencies in all (seven) pairwise comparisons, however, two markers (TP402859 and SEP3-F1) showed such differences in six pairs, another two markers (GI-F1 and FLD-F1) in five pairs, and five markers (TP23903, VIP3-F2, TP288840, TP100150, TP3177) in four pairs (Supplementary Table S8). Besides FLD-F1 and VIP3-F2, these markers originated from QTL1, QTL3 and QTL4. Such a result provided additional evidence highlighting hypothetical involvement of genes and QTLs represented by these markers in flowering time control in white lupin.

3. Discussion

3.1. Contribution of FLOWERING LOCUS T Genes to Early Flowering in White Lupin

In white lupin, early flowering accessions were found in both wild/primitive and domesticated germplasm. Vernalization responsiveness in mapping population, as well as in germplasm collection, was revealed as a continuous trait suggesting quantitative (polygenic) regulation. In the \textit{L. albus} mapping population, several QTL loci for time to flowering were identified hitherto, including five QTLs confirmed in at least two years of observations [24–26]. Following the construction of a high-density reference linkage map, sequence-defined markers flanking these QTLs were developed [26,29]. Moreover, recent research highlighted candidate genes for four QTLs, namely \textit{GIGANTEA} (GI) for the QTL1, \textit{FLOWERING LOCUS T} homolog a1 (\textit{FTa1}) for the QTL2, \textit{SEPALLATA 3} (\textit{SEP3}) for the QTL4 and \textit{FRIGIDA3} (\textit{FRI3}) for the QTL5 [30]. In this study, white lupin molecular toolbox was supplemented with another 17 CAPS and dCAPS markers to facilitate efficient PCR-based tracking of all five major QTLs in germplasm which is unrelated to mapping population. Thus, this study confirmed the viability of all major white lupin QTLs for time to flowering, highlighted by statistically significant correlations between the phenotype and the genotype identified by particular markers (see Figure 2 and Table 3).

Correlation coefficients revealed in this study as significant reached the maximum values from 0.32 to 0.38 in the whole white lupin germplasm collection and from 0.35 to 0.95 in the particular sub-populations. Such values are typical for multi-locus traits. Indeed, in our previous study, targeting narrow-leafed lupin, a species with vernalization independence conferred by just a single gene [34], the marker anchored in this gene reached correlation coefficient values from $-0.48$ (yield) to 0.80 (flowering time), whereas PCR-based markers from other genes, also evidenced as statistically significant, reached values from 0.20 to 0.38 [35]. A similar study in soybean revealed, for the marker significantly associated with maturity dates and plant height, correlation coefficient values of $-0.27$ and $-0.51$ [36].

Interestingly, only two (\textit{FRI3} and \textit{SEP3}) from four candidate genes (\textit{FRI3}, \textit{FTa1}, \textit{GI}, and \textit{SEP3}) highlighted by a recent mapping population study [30] revealed significant correlations with time to flowering in this study targeting germplasm collection. The lack of statistical support for the \textit{FTa1} from QTL2 gene may result from very low frequency of the wild allele, which was found only in three accessions: two Ethiopian landraces and one Polish breeding line. Moreover, \textit{FTa1} indel marker is localized in the third exon of this gene, whereas major regulatory components of FT expression in \textit{Arabidopsis thaliana} were evidenced to be localized in the promoter and in the first intron [37–40]. Nevertheless, three other markers from QTL2, namely TP235608, TP94353 and TP56963, were found to be significantly correlated with early phenology but only in one or two years.

The present study evidenced also significant correlations between several other genetic components of flowering regulation pathways, including also another white lupin FT homolog, an \textit{FTc1} gene –both in the whole analyzed collection as well as in one of the subpopulations (V5). In the narrow-leafed lupin, \textit{Lupinus angustifolius} L., early flowering is based on vernalization independence originating from just two natural mutations (named
white lupin. Ku and Jul) discovered in domesticated germplasm in 1960s [41,42]. Ku is based on a large (1.4 kbp) deletion in the promoter region of one of the four L. angustifolius FT homologs, FTc1 [34]. This deletion carries potential binding sites for several transcription factors acting as FT gene repressors in A. thaliana [43]. The second L. angustifolius domesticated early phenology mutation, Jul, was recently revealed to be a 5162 bp deletion in the Ftc1 promoter region, fully encompassing the Ku indel [44]. Recently, a fourth Ftc1 allele, Pal, was found in a wild population originating from Palestine, carrying 1208 bp deletion partially overlapping with Ku [44,45]. Recent gene-based genome-wide association study confirmed that this series of indels in Ftc1 has a major effect on time to flowering and maturity in diversified narrow-leaved germplasm collection [35].

The release of two white lupin high quality genome assemblies [27,28] provided an unprecedented opportunity to search for similar indels in FT homologs in this species, both by whole-genome shotgun and PCR-based approaches. Recent comparative mapping approach performed in yellow lupin, Lupinus luteus L., revealed that a major QTL for vernalization responsiveness in this species is localized in a linkage map segment syntenic to the narrow-leaved lupin genome region carrying the Ftc1 gene [46–48]. All these observations support the conclusion on the highly conserved contribution of the Ftc1 homologs into vernalization pathway in the Old World lupin clade.

The involvement of an Ftc in flowering time control is relatively rare phenomenon in plants, as Ftc homologs appeared only in legumes as a result of whole-genome duplication event(s) [43]. Thus, in the legume model plant, Medicago truncatula L., vernalization responsiveness is more likely conferred by the MtFti1 gene (which is also highly induced by long day conditions), whereas photoperiod pathway by the MtFTb1 and MtFTb2 genes [49–51]. In pea, FT genes showed transcriptional sub-functionalization and one of those, Fti1, underlies the pea GIGAS locus, essential for flowering under long-days and positive for flowering under short-days [50]. In soybean, which is a vernalization independent species, three genes belonging to Fti and Ftc clades confer just the photoperiod response [52–55]. In chickpea, altered expression of a cluster of three FT genes (Fta1, Fta2 and Ftc) is associated with early phenology [56], however, a major QTL for vernalization response in this species is localized in a genome region lacking any FT homolog [57].

3.2. Candidates from Photoperiod, Vernalization, Autonomous and Heat-Responsive Pathways for Flowering Time Control in White Lupin

Besides Ftc1, our study revealed significant correlations between time to flowering and sequence polymorphism in markers anchored in the sequences of the following genes: CO-like, ELF1, FLD, FRI, FY, LD, MFT, PIF4, SEP3, SKIP1, and VIP3. Without any data about linkage disequilibrium decay we could not infer about the size of haplotype blocks represented by these markers. Therefore, we focused on the analysis of hypothetical involvement of genes represented by particular markers in regulation of flowering time in white lupin.

CO-F1 marker was found to be significantly correlated with flowering time of white lupin in all three experiments, both in the whole set of lines as well as in one of the subpopulations (V6). CO gene is a well-known central regulator of photoperiod pathway in A. thaliana [58,59]. In M. truncatula, a homolog of this gene was localized in the major QTL for flowering date and revealed different expression profiles in parental lines contrasting for flowering time [60,61]. The association between CO-like gene and flowering time was also confirmed for Medicago sativa [62]. These observations support the concept of choosing CO gene as one of the targets in artificial selection in white lupin.

ELF1-F1 marker revealed significant correlation in two years (2018 and 2020). The lack of statistical support for correlation in the first experiment might be related with the lower number of white lupin lines analyzed, especially those with late flowering phenotype. ELF1 gene is hypothetically involved in flowering regulation in A. thaliana, however, particular mechanism has not been yet deciphered [63]. ELF1, together with an FLD gene and other six potential flowering time-regulating genes, were highlighted by genome-scale association and QTL mapping of complex flowering time trait in chickpea [64].
FLD-F1 and LD-F1 markers showed significant correlation with flowering time in the last year, whereas FY-F6 marker in all three years. FLD, FY and LD in *A. thaliana* are components of the autonomous pathway that promotes flowering independently of the photoperiod and vernalization [65,66]. It was evidenced that soybean genome encodes a functional copy of an *FLD* gene which promotes flowering when introduced into late flowering *A. thaliana* fld mutant [67].

FRI31-F1 marker revealed significant association with flowering time in one of the subpopulations (V6) in the years 2015 and 2020. Besides the floral repressor **FLOWERING LOCUS C** (*FLC*), **FRI** is a second key component of vernalization pathway in *A. thaliana*, and allelic variation of **FRI** accounts for the majority of natural flowering time variation in germplasm collection of this species [68,69]. **FRI** upregulates expression of the **FLC** by acting in a supercomplex that establishes a local chromosomal environment facilitating production of the **FLC** mRNA [70]. Nevertheless, despite revealed significant association between the **FRI** marker and time to flowering in white lupin, direct translation of this mechanism from *A. thaliana* into this species is not possible, due to the lack of any **FLC** homolog in white lupin and other legume genomes, except soybean [30,71,72]. Interestingly, in soybean, SNP variation in **FRI** sequence was also found to be highly associated with flowering time, shedding new light on potential contribution of this gene into flowering regulation network apart from the vernalization pathway [73].

VIP3-F2 marker, tagging another component of vernalization pathway in *A. thaliana*, was significantly associated with white lupin flowering time in all experiments. It was revealed that loss-of-function mutation in the VIP3 gene suppress the effect of FRI on FLC expression and flowering time in *A. thaliana* [74,75]. To our best knowledge, the present study is the first report on association between VIP3 and flowering time in legumes.

Similarly to VIP3-F2, **SKIP1-F2** marker also revealed significant correlation with white lupin flowering time in all experiments. **SKIP** is a component of flowering regulation network linking alternative splicing and the circadian clock in *A. thaliana* [76]. It works as a splicing factor as part of the spliceosome and as a transcriptional activator interacting with **EARLY FLOWERING 7** (*ELF7*) [77]. **SKIP** is required for the splicing of **serrated leaves and early flowering** (*Sef*) pre-messenger RNA (mRNA) in *A. thaliana* [78]. A **SKIP** homolog in soybean, **GmGBP1**, acts as a positive regulator upstream of **GmFT2a** and **GmFT5a** and promotes flowering on short days. Moreover, natural variation in **GmGBP1** promoter sequence is associated with photoperiod control of soybean flowering time and maturity [79]. Transgenic *A. thaliana* with the ectopic overexpression of **GmGBP** revealed advanced flowering under long days via photoperiodic and gibberellin pathways (including **CO** and **FT** genes) and delayed flowering under short days via autonomous pathway (**SVP** and **FLC**) [80].

**PIF4-F6** marker was significantly correlated with white lupin flowering time in two years (both in the whole set of lines and in the subpopulation V2). **PIF4** is associated in variation of ecologically important traits in *A. thaliana*, including time to flowering [81]. Expression of **PIF4** is induced proportionally to the increase of ambient temperature (at least in the range 12–27 °C), resulting in strong induction of the **FT** gene and overcome of late flowering phenotype of *A. thaliana* [82]. Similar correlation of **PIF4** expression and temperature was also observed in soybean [83]. In our study, **PIF4** marker revealed significant association with flowering time in white lupin collection in 2015 and 2018, and non-significant in 2020. Indeed, the number of days with temperature above 20 °C during first 40 days of the experiment was higher in 2015 and 2018 years than in 2020 (14 and 17 vs. 7). Such an observation may explain observed differences in correlation values and provide some support on the conserved function of **PIF4** as a thermosensory activator of flowering in white lupin.

Two **MFT** markers, **MFTa1-F1** and **MFT-FT3-F1** showed significant correlation with white lupin flowering time in all years, one in the whole collection and the second in the subpopulation V6 (together with **CO** and **FRI**). **MFT** genes constitute the basal clade among phosphatidylethanolamine-binding (PEPB) domain genes and are present in angiosperms,
gymnosperms, lycophytes and bryophytes [84]. Duplication of an ancient MFT-like gene hypothetically contributed to the radiation of seed plants [85]. Overexpression of the MFT gene in *A. thaliana* resulted in slightly early flowering under long days [86]. It was recently evidenced that the MFT functions as a key *A. thaliana* repressor of germination under far-red light conditions [87]. Moreover, MFT is considered a potential candidate gene for a major QTL that alters *A. thaliana* flowering time at elevated CO$_2$ [88]. CO$_2$ concentration in greenhouse at seedling stage can be higher than outside due to the vigorous soil respiration and lower canopy photosynthetic rate [89]. Thus, any of the mentioned functions could be assigned to explain observed significant correlation between white lupin flowering time and MFT gene-based marker polymorphism.

3.3. Perspectives for Molecular-Assisted Breeding of White Lupin

High number of markers which significantly correlated with flowering time suggests that genome-wide selection should be a method of choice for white lupin breeders, rather than the classical approach utilizing marker-assisted selection. Introduction of preferred alleles on one-by-one basis can be inefficient, due to epistatic interactions between major components of flowering regulatory network [90]. Phenotyping studies revealed that white lupin germplasm collection has variability of many agronomic traits, including, besides time to flowering, winter survival, pod fertility, pod wall proportion, number of leaves on the main stem, several yield related traits and anthracnose resistance [11–13,29,91–94]. Despite relatively small genotype sample size (83 landraces and eight varieties), genomic prediction for grain yield, winter plant survival and onset of flowering provided prospective output, manifested by model-based predictive ability values as high as 0.84–0.86 [10]. In that study two models were tested, Bayesian Lasso [95] and ridge regression best linear unbiased prediction (rrBLUP) [96,97], providing similar results. Genomic prediction of grain yield in three European sites with contrasting climate (Mediterranean, subcontinental or oceanic) displayed relatively high intra-environment (up to 0.71) and cross-environment (up to 0.51) predictive abilities, providing economic justification for genomic selection strategy in white lupin breeding [11].

3.4. Concluding Remarks

This study provided novel evidence for quantitative control of flowering time in white lupin by supporting all major QTLs derived from mapping population studies as well as designating new candidate genes from major molecular pathways regulating flowering induction in plants. A very high majority of markers revealed an association of a Kiev Mutant allele with accelerated flowering, indicating that desired alleles have been already introduced into domesticated germplasm. Nevertheless, two important components of flowering regulatory network, *PIF4* and *LD*, showed an opposite association, opening up the development potential for white lupin breeders. Taking into consideration the results of studies obtained for other legume species, such as narrow-leafed lupin or soybean [34,79,98], resequencing of candidate genes in wider germplasm background, including also promoter regions, could be valuable by providing novel alleles conferring early or intermediate phenology.

4. Materials and Methods

4.1. Plant Material

The set of 160 *L. albus* lines, derived from the European Lupin Gene Resources Database maintained by Poznañ Plant Breeders Ltd. station located in Wiatrowo as well as from the National Centre for Plant Genetic Resources: Polish Genebank, Plant Breeding and Acclimatization Institute–National Research Institute, Radzików, 05–870 Blonie, Poland, was used in the study. This germplasm collection contained 63 wild or primitive populations, 51 landraces, 31 cultivars, 12 cross derivatives and three mutants. These lines originated from 23 countries. The information on germplasm donor, country of origin and domestication status was provided in Supplementary Table S1.
4.2. Profiling of Time to Flowering and Vernalization Responsiveness in Controlled Environment

Phenotyping experiments were performed in a greenhouse located at the Institute of Plant Genetics, Polish Academy of Sciences, Poznań, Poland (52°26′ N 16°54′ E) during growing seasons of 2015 (sowing date 10.04), 2018 (sowing date 22.03) and 2020 (sowing date 19.03) under ambient long day photoperiod (12–17 h). To randomize greenhouse-related effects between years and lines, the experiment was performed every year in a different greenhouse and in a different design (Supplementary Figure S1). Automatic heating was used to keep the minimum air temperature above 18 °C. Cooling was maintained by temperature-dependent ventilation system (activated at 22 °C). Vernalization was carried out by placing seeds for 23 days at 5 °C in darkness on moist filter paper in Petri dishes. Non-vernalized plants were sown five days before the end of vernalization period and cultivated at ~23 °C to keep a similar thermal time [99]. Time to flowering was recorded as the number of days from sowing of vernalized plants until the first fully colored petal was developed. Observations were made for each plant separately. The average number of plants with observations in 2015 was 5.1 for non-vernalized variant (min 3, max 9, \( n = 105 \)) and 4.6 for vernalized variant (min 3, max 8, \( n = 104 \)), in 2018 it was 7.9 for non-vernalized variant (min 3, max 10, \( n = 140 \)) and 9.0 for vernalized variant (min 3, max 10, \( n = 138 \)) whereas in 2020 it was 8.8 in non-vernalized variant (min 3, max 10, \( n = 160 \)). Daily mean and maximum air temperature and daily sunshine hours recorded by the nearby localized meteorological station (Poznań-Lawica, 5.1 km away) and theoretical photoperiod hours calculated for this latitude (covering 100 days from sowing date) were provided for reference in Supplementary Table S9.

4.3. Development of Molecular Markers for Flowering Time QTLs

PCR-based molecular markers were developed for single nucleotide polymorphisms (SNPs) localized on the white lupin linkage map in the proximity of all major QTLs for flowering time. The following QTLs [26,30] were explored: linkage group ALB02 (QTL1, position around 2.2 cM), ALB02 (QTL2, 100.2/100.5 cM), ALB13 (QTL3, 81.0/84.1 cM and QTL4, 96.2/99.3 cM), and ALB16 (QTL5, 0.9/2.2 cM). Genotyping-by-sequencing (GBS) reads which were previously mapped in these regions, were aligned to transcriptome sequences of parental lines of \( L. albus \) mapping population [26] as well as the reference transcriptome of \( L. albus \) roots and leaves (https://legumeinfo.org/data/public/Lupinus_albus/ (accessed date 7 April 2021), gene index LAGI01) [33]. BLASTn algorithm [100] implemented in Geneious v8.1 (Biomatters, New Zealand) [101] was used for this alignment with max 2 mismatches allowed per sequence.

To map intron/exon boundaries, matched transcript sequences were extracted and re-aligned to the genome scaffolds [27] using progressive Mauve algorithm with gapped aligner MUSCLE 3.6 [102,103]. Mauve alignments consisting of GBS reads, Kiev Mutant, P27174 and LAGI01 transcripts, and genome scaffolds, were screened for the presence of polymorphic loci. The primers flanking these loci were designed using Primer3Plus [104,105]. Depending on the availability of restriction enzymes, SNPs were transformed the cleaved amplified polymorphic sequence (CAPS) [106] or derived CAPS (dCAPS) [107] markers. Restriction sites and dCAPS primers were identified using dCAPS Finder 2.0 and SNP2dCAPS [108,109].

4.4. Genotyping of White Lupin Germplasm with PCR-Based Markers

The white lupin germplasm collection which was subjected to phenotyping of flowering time and vernalization responsiveness, was also genotyped with PCR-based markers. The set of markers included markers for flowering time QTLs developed in this study as well as previously published markers designed for white lupin homologs of flowering control genes [30] and those developed for the anthracnose resistance QTL localized at the linkage group ALB02 overlapping with the flowering time QTL1 [29]. Young 5 week-old leaves were collected from plants cultivated in greenhouse and immediately frozen under liquid nitrogen. Frozen plant tissue (50–100 mg) was homogenized using TissueLyser II
(Qiagen, Hilden, Germany) and two stainless steel beads (ø 5 mm) placed in a 2 mL tube (Eppendorf, Hamburg, Germany). DNA isolation was performed using DNeasy Plant Mini Kit (Qiagen). PCR products were amplified using GoTaq G2 Flexi DNA Polymerase (Promega, Mannheim, Germany). Restriction enzyme digestion was performed for 12 h in 20 µL of mixture including 5 µL of post-PCR mixture, one unit of enzyme and appropriate amount of water and buffer to reach concentration recommended by the enzyme manufacturer. Restriction products were separated by agarose gel electrophoresis, with the agarose concentration (1–3%) adjusted to follow the size of the expected digestion products. Wide range agarose (Serva, Heidelberg, Germany) was used for most applications, however, markers yielding polymorphic cut products shorter than 100 bp were resolved using high resolution 3:1 agarose (Serva).

The list of markers used in the study with localization on linkage map, QTL assignment, accession numbers and sequences of both alleles is provided in Supplementary Table S4. Information on PCR annealing temperature, polymorphism detection method, expected products and agarose gel concentration is provided in Supplementary Table S5. Two biological replicates were analyzed per line. Kiev Mutant marker alleles were assigned “1”, P27174 marker alleles were assigned “2”, whereas heterozygotes “1.5”. Additional alleles (different than those observed in Kiev Mutant or P27174) were assigned “C” or “3”.

4.5. Population Structure Analysis

Population structure analysis was performed with LEA R package [110]. Obtained marker data were converted to multi-allele geno format using a custom Python script converting every allele into a specific character symbol. Next, the optimal number of clusters (k) in the investigated population was determined using admixture analysis [111,112] with smnf function from the same package. The entropy criterion evaluating the quality fit of the statistical model to the data was calculated in 1000 repetitions and 10,000 iterations. The optimal number of clusters was set as k with minimal cross-entropy values [113,114]. The ancestry matrix for optimal k = 6 was visualized with the barchar function from LEA package.

4.6. Statistical Analysis

All statistical analysis were preferred in R software (R Core Team 2013, Vienna, Austria) using base function. Simple regression was used to estimate QTL/marker additive effects. The correlation between alleles in all 50 loci and flowering time was calculated using Spearman’s rank method [115]. The calculation was made for the whole population and separately for each cluster obtained by population structure analysis. Homozygote wild (P27174) got rank 3, homozygote cultivated (Kiev Mutant)–rank 1, and heterozygote–rank 2. Novel alleles, which occurred only for some markers and few lines, were treated as missing values. Giving rank 1 or rank 3 for novel alleles had negligible effect on obtained correlation values, highlighted by ~0.99 correlation of the results. The two-way variance analysis was performed to test the hypothesis of lack of difference in flowering time between clusters obtained in the population structure analysis. The variance model was given by the following formula:

\[ Y_{ijk} = \mu + \alpha_i + \beta_j + \epsilon_{ijk} \]

where \( Y_{ijk} \) is flowering time of \( k \)-th genotype in \( j \)-th year and \( i \)-th cluster, \( \alpha_i \) is \( i \)-th cluster, \( \beta_j \) is \( j \)-th year, \( \mu \) is cluster \( \times \) year interaction and \( \epsilon_{ijk} \) is the random effect component. The true difference between the mean was tested using Tukey’s ‘Honest Significant Difference’ method [116]. The \( \chi^2 \) test was used to test the hypothesis of lack of difference in allele frequency between clusters, which in ANOVA analysis showed a difference in flowering time.
Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/jms22083856/s1, Table S1: White lupin lines used in the study, Table S2: Number of days from sowing to flowering observed for white lupin lines used in the study, Table S3: Mean values and standard deviation of flowering time observed for white lupin lines used in the study, Table S4: The list of markers used in the study with localization on linkage map, QTL assignment, accession numbers and sequences of both alleles, Table S5: PCR annealing temperature, polymorphism detection method, expected products and agarose gel concentration for markers used in the study, Table S6: Results of genotyping of white lupin germplasm collection with PCR-based markers, Table S7: p-values of Spearman’s rank correlation coefficient calculated for flowering time and marker distribution in white lupin germplasm collection, Table S8: p-values of Chi-square test of differences in allele frequencies between subpopulations, Table S9: Meteorological conditions recorded by the nearby localized meteorological station (Poznań-Lawica, 5.1 km) and theoretical photoperiod hours calculated for this latitude (covering 100 days from sowing date). Figure S1. Orientation of greenhouses and arrangement of tables in plant phenotyping experiments.

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References
1. Gladstones, J.S. Lupins as crop plants. *Field Crop Abstr.* **1970**, *23*, 26.
2. Lambers, H.; Clements, J.C.; Nelson, M.N. How a phosphorus-acquisition strategy based on carboxylate exudation powers the success and agronomic potential of lupines (*Lupinus*, Fabaceae). *Am. J. Bot.* **2013**, *100*, 263–288. [CrossRef]
3. Watt, M.; Evans, J.R. Phosphorus acquisition from soil by white lupin (*Lupinus albus*) and soybean (*Glycine max*), species with contrasting root development. *Plant Soil* **2003**, *248*, 271–283. [CrossRef]
4. Schulze, J.; Temple, G.; Temple, S.J.; Beschow, H.; Vance, C.P. Nitrogen fixation by white lupin under phosphorus deficiency. *Ann. Bot.* **2006**, *98*, 731–740. [CrossRef] [PubMed]
5. Papineau, J.; Huyghe, C. *Le Lupin Dous Protéagineux*; France Agricole: Paris, France, 2004; p. 176.
6. Annicchiarico, P.; Manunza, P.; Arnoldi, A.; Boschin, G. Quality of *Lupinus albus* L. (white lupin) seed: Extent of genotypic and environmental effects. *J. Agric. Food Chem.* **2014**, *62*, 6539–6545. [CrossRef]
7. Boschin, G.; D’Agostina, A.; Annicchiarico, P.; Arnoldi, A. The fatty acid composition of the oil from *Lupinus albus* cv. Luxe as affected by environmental and agricultural factors. *Eur. Food Res. Technol.* **2007**, *225*, 769–776. [CrossRef]
8. Arnoldi, A.; Greco, S. Nutritional and nutraceutical characteristics of lupin protein. *Nutrafoods* **2011**, *10*, 23–29. [CrossRef]
9. Kroc, M.; Rybiński, W.; Wilczura, P.; Kamel, K.A.; Kaczmarek, Z.; Barzyk, P.; Święcicki, W. Quantitative and qualitative analysis of alkaloids composition in the seeds of a white lupin (*Lupinus albus* L.) collection. *Genet. Resour. Crop Evol.* **2017**, *64*, 1853–1860. [CrossRef]
10. Annicchiarico, P.; Nazzicari, N.; Ferrari, B. Genetic and genomic resources in white lupin and the application of genomic selection. In *The Lupin Genome*; Singh, K.B., Kamphuis, L.G., Nelson, M.N., Eds.; Springer International Publishing: Cham, Switzerland, 2020; pp. 139–149.
11. Annicchiarico, P.; Nazzicari, N.; Ferrari, B.; Harzic, N.; Carroni, A.M.; Romani, M.; Pecetti, L. Genomic prediction of grain yield in contrasting environments for white lupin genetic resources. *Mol. Breed.* **2019**, *39*, 142. [CrossRef]
12. Annicchiarico, P.; Romani, M.; Pecetti, L. White lupin (Lupinus albus) variation for adaptation to severe drought stress. Plant Breeding 2018, 137, 782–789. [CrossRef]
13. Annicchiarico, P.; Harizic, N.; Carroni, A.M. Adaptation, diversity, and exploitation of global white lupin (Lupinus albus L.) landrace genetic resources. Field Crops Res. 2010, 119, 114–124. [CrossRef]
14. Gladstones, J.S. The Mediterranean white lupin. J. Dep. Agric. West. Aust. Ser. 4 1976, 17, 70–74.
15. Adhikari, K.N.; Buichell, B.J.; Sweetingham, M.W. Length of vernalization period affects flowering time in three lupin species. Plant Breed. 2012, 131, 631–636. [CrossRef]
16. Weimer, J.L. Anthracnose of lupines. Phytopathology 1943, 33, 249–252.
17. Talhinhas, P.; Baronecelli, R.; Floch, G.L. Anthracnose of lupins caused by Colletotrichum lupini: A recent disease and a successful worldwide pathogen. J. Plant Pathol. 2016, 98, 5–14.
18. Fencel, I.M. Report on first detection of anthracnose (Colletotrichum gloeosporioides) on lupins in Poland. Plant Dis. 1998, 82, 350. [CrossRef]
19. Gondran, J.; Bateman, G.L.; Milford, G.F.J.; Bayer, J.; Beerepoot, L.; Caligari, P.D.S.; Carrasco-Lopez, J.M.; Crowley, J.G.; da Rocha, J.P.; et al. Anthracnose of white lupin (Lupinus albus): European prospects for a future sustainable crop. In Towards the 21st Century. Proceedings of the 8th International Lupin Conference; Hill, G.D., Ed.; International Lupin Association: Asti, CA, USA, 1996; pp. 512–518.
20. Thomas, G.J.; Sweetingham, M.W. Cultivar and environment influence the development of lupin anthracnose caused by Colletotrichum lupini. Australas. Plant Path. 2004, 33, 571–577. [CrossRef]
21. Sweetingham, M.W.; Fencel, I.; Buichell, B.J.; Barzyk, P.; Lewartowska, E.; Yang, H.A.; Ponomaryova, L.; Yakasheva, A.; Adhikari, K.N.; Francis, C.M.; et al. Collaborative anthracnose resistance screening in Poland, Russia and Australia. México, where old and new world lupins meet. In Proceedings of the 11th International Lupin Conference, Guadalajara, Mexico, 4–9 May 2005; International Lupin Association: Canterbury, New Zealand, 2006.
22. Adhikari, K.N.; Buichell, B.J.; Thomas, G.J.; Sweetingham, M.W.; Yang, H. Identification of anthracnose resistance in Lupinus albus L. and its transfer from landraces to modern cultivars. Crop Pasture Sci. 2009, 60, 472–479. [CrossRef]
23. Atnaf, M.; Yao, N.; Martina, K.; Daghe, K.; Wegary, D.; Tesfaye, K. Molecular genetic diversity and population structure of Ethiopian white lupin landraces: Implications for breeding and conservation. PLoS ONE 2017, 12, e0188696. [CrossRef]
24. Phan, H.T.T.; Ellwood, S.R.; Adhikari, K.; Nelson, M.N.; Oliver, R.P. The first genetic and comparative map of white lupin (Lupinus albus L.): Identification of QTLs for anthracnose resistance and flowering time, and a locus for alkaloid content. DNA Res. 2007, 14, 59–70. [CrossRef]
25. Vipin, C.A.; Luckett, D.J.; Harper, J.D.I.; Ash, G.J.; Kilian, A.; Ellwood, S.R.; Phan, H.T.T.; Raman, H. Construction of integrated linkage map of a recombinant inbred line population of white lupin (Lupinus albus L.). Breed. Sci. 2013, 63, 292–300. [CrossRef] [PubMed]
26. Książkiewicz, M.; Nazzicari, N.; Yang, H.; Nelson, M.N.; Renshaw, D.; Rychel, S.; Ferrari, B.; Carelli, M.; Tomaszewska, M.; Stawirski, S.; et al. A high-density consensus linkage map of white lupin highlights synteny with narrow-leaved lupin and provides markers tagging key agronomic traits. Sci. Rep. 2017, 7, 15335. [CrossRef]
27. Hufnagel, B.; Marques, A.; Soriano, A.; Marquès, L.; Divol, F.; Doumas, P.; Sallet, E.; Mancinotti, D.; Carrere, S.; Marande, W.; et al. High-quality genome sequence of white lupin provides insight into soil exploration and seed quality. Nat. Commun. 2020, 11, 492. [CrossRef] [PubMed]
28. Xu, W.; Zhang, Q.; Yuan, W.; Xu, F.; Muhammad Aslam, M.; Miao, R.; Li, Y.; Wang, Q.; Li, X.; Zhang, X.; et al. The genome evolution and low-phosphorus adaptation in white lupin. Nat. Commun. 2020, 11, 1069. [CrossRef] [PubMed]
29. Rychel-Bielska, S.; Nazzicari, N.; Plewiński, P.; Bielski, W.; Annicchiarico, P.; Książkiewicz, M. Development of PCR-based markers and whole-genome selection model for anthracnose resistance in white lupin (Lupinus albus L.). J. Appl. Genet. 2020, 61, 531–545. [CrossRef]
30. Rychel, S.; Książkiewicz, M.; Tomaszewska, M.; Bielski, W.; Wolko, B. FLOWERING LOCUS T, GIGANTEA, SEPELLATA and FRIGIDA homologs are candidate genes involved in white lupin (Lupinus albus L.) early flowering. Mol. Breed. 2019, 39, 43. [CrossRef]
31. Rychel, S.; Książkiewicz, M. Development of gene-based molecular markers tagging low alkaloid pauper locus in white lupin (Lupinus albus L.). J. Appl. Genet. 2019, 60, 269–281. [CrossRef] [PubMed]
32. Adhikari, K.N.; Thomas, G.; Diepeveen, D.; Trethowan, R. Overcoming the barriers of combining early flowering and anthracnose resistance in white lupin (Lupinus albus L.) for the Northern Agricultural Region of Western Australia. Crop Pasture Sci. 2013, 64, 914–921. [CrossRef]
33. O’Rourke, J.A.; Yang, S.S.; Miller, S.S.; Bucciarelli, B.; Liu, J.; Rydeen, A.; Bozsoki, Z.; Udhe-Stone, C.; Tu, Z.J.; Allan, D.; et al. An RNA-Seq transcriptome analysis of orthophosphate-deficient white lupin reveals novel insights into phosphorus acclimation in plants. Plant Physiol. 2013, 161, 705–724. [CrossRef]
34. Nelson, M.N.; Książkiewicz, M.; Rychel, S.; Besharat, N.; Taylor, C.M.; Wyrwa, K.; Jost, R.; Erskine, W.; Cowling, W.A.; Berger, J.D.; et al. The loss of vernalization requirement in narrow-leaved lupin is associated with a deletion in the promoter and de-repressed expression of a Flowering Locus T (FT) homologue. New Phytol. 2017, 213, 220–232. [CrossRef]
35. Plewinski, P.; Cwik-Kupczyńska, H.; Rudy, E.; Bielski, W.; Rychel-Bielska, S.; Stawiski, S.; Barzyk, P.; Krajewski, P.; Naganowska, B.; Wólko, B.; et al. Innovative transcriptome-based genotyping highlights environmentally responsive genes for phenology, growth and yield in a non-model grain legume. Plant Cell Environ. 2020, 43, 2680–2698. [CrossRef]

36. Zhang, J.; Song, Q.; Cregan, P.B.; Nelson, R.L.; Wang, X.; Wu, J.; Jiang, G.-L. Genome-wide association study for flowering time, maturity dates and plant height in early maturing soybean (Glycine max) germplasm. BMC Genom. 2015, 16, 217. [CrossRef] [PubMed]

37. Cao, S.; Kumimoto, R.W.; Gnesutta, N.; Calogero, A.M.; Mantovani, R.; Holt, B.F., 3rd. A distal CCAAT/NUCLEAR FACTOR Y complex promotes chromatin looping at the FLOWERING LOCUS T promoter and regulates the timing of flowering in Arabidopsis. Plant Cell 2014, 26, 1009–1017. [CrossRef] [PubMed]

38. Liu, L.; Adrian, J.; Pankin, A.; Hu, J.; Dong, X.; von Korff, M.; Turck, F. Induced and natural variation of promoter length modulates the photoperiodic response of FLOWERING LOCUS T. Nat. Commun. 2014, 5, 4558. [CrossRef]

39. Turck, F.; Fornara, F.; Coupland, G. Regulation and identity of florigen: FLOWERING LOCUS T moves center stage. Annu. Rev. Plant Biol. 2008, 59, 573–594. [CrossRef]

40. Taylor, C.M.; Kamphuis, L.G.; Zhang, W.; Garg, G.; Berger, J.D.; Mousavi-Derazmahalleh, M.; Bayer, P.E.; Edwards, D.; Singh, K.B.; Cowling, W.A.; et al. IND1 variation in the regulatory region of the major flowering time gene LanFTc1 is associated with vernalization response and flowering time in narrow-leaved lupin (Lupinus L.). Plant Cell Environ. 2019, 42, 174–187. [CrossRef] [PubMed]

41. Lichtin, N.; Salvo-Garrido, H.; Till, B.; Caligari, P.D.S.; Rupayan, A.; Westermeyer, F.; Olivos, M. Genetic and comparative mapping of Lupinus luteus L. highlight syntenic regions with major orthologous genes controlling anthracnose resistance and flowering time. Sci. Rep. 2020, 10, 19174. [CrossRef] [PubMed]

42. Iqbal, M.M.; Erskine, W.; Berger, J.D.; Nelson, M.N. Phenotypic characterisation and linkage mapping of domestication syndrome genes in the narrow-leaved lupin (Lupinus angustifolius L.). Front. Plant Sci. 2020, 11, 572135. [CrossRef]

43. Liu, L.; Adrian, J.; Pankin, A.; Hu, J.; Dong, X.; von Korff, M.; Turck, F. Induced and natural variation of promoter length modulates the photoperiodic response of FLOWERING LOCUS T. Nat. Commun. 2014, 5, 4558. [CrossRef] [PubMed]

44. Iqbal, M.M.; Huynh, M.; Udall, J.A.; Kilian, A.; Adhikari, K.N.; Berger, J.D.; Erskine, W.; Nelson, M.N. The first genetic map for yellow lupin enables genetic dissection of adaptation traits in an orphan grain legume crop. BMC Genet. 2019, 20, 68. [CrossRef] [PubMed]

45. Rychel-Bielska, S.; Plewinski, P.; Kozak, B.; Galek, R.; Książkiewicz, M. Photoperiod and vernalization control of flowering-related genes: A case study of the narrow-leaved lupin (Lupinus angustifolius L.). Plant Physiol. 2017, 17, 820. [CrossRef] [PubMed]

46. Takeshima, R.; Hayashi, T.; Zhu, J.; Zhao, C.; Xu, M.; Yamaguchi, N.; Sayama, T.; Ishimoto, M.; Kong, L.; Shi, X.; et al. A soybean homolog necessary for graft-transmissible specification of flowering time in narrow-leaved lupin (Lupinus L.). Plant Cell Environ. 2019, 2019, 217. [CrossRef] [PubMed]

47. Kong, F.; Liu, B.; Xia, Z.; Sato, S.; Kim, B.M.; Watanabe, S.; Yamada, T.; Tabata, S.; Kanazawa, A.; Harada, K.; et al. Two coordinately regulated homologs of FLOWERING LOCUS T are involved in the control of photoperiodic flowering in soybean. Plant Physiol. 2010, 154, 1220–1231. [CrossRef]

48. Chen, L.; Cai, Y.; Qu, M.; Wang, L.; Sun, H.; Jiang, B.; Wu, T.; Liu, L.; Sun, S.; Wu, C.; et al. Soybean adaption to high-latitude germplasm. BMC Genom. 2015, 16, 217. [CrossRef] [PubMed]

49. Liu, L.; Adrian, J.; Pankin, A.; Hu, J.; Dong, X.; von Korff, M.; Turck, F. Induced and natural variation of promoter length modulates the photoperiodic response of FLOWERING LOCUS T. Nat. Commun. 2014, 5, 4558. [CrossRef]

50. Taylor, C.M.; Kamphuis, L.G.; Zhang, W.; Garg, G.; Berger, J.D.; Mousavi-Derazmahalleh, M.; Bayer, P.E.; Edwards, D.; Singh, K.B.; Cowling, W.A.; et al. IND1 variation in the regulatory region of the major flowering time gene LanFTc1 is associated with vernalization response and flowering time in narrow-leaved lupin (Lupinus L.). Plant Cell Environ. 2019, 2019, 217. [CrossRef] [PubMed]

51. Thomson, G.; Taylor, J.; Putterill, J. The transcriptomic response to a short day to long day shift in leaves of the reference legume Medicago truncatula. PeerJ 2019, 7, e6626. [CrossRef]

52. Kong, F.; Liu, B.; Xia, Z.; Sato, S.; Kim, B.M.; Watanabe, S.; Yamada, T.; Tabata, S.; Kanazawa, A.; Harada, K.; et al. Two coordinately regulated homologs of FLOWERING LOCUS T are involved in the control of photoperiodic flowering in soybean. Plant Physiol. 2010, 154, 1220–1231. [CrossRef]

53. Chen, L.; Cai, Y.; Qu, M.; Wang, L.; Sun, H.; Jiang, B.; Wu, T.; Liu, L.; Sun, S.; Wu, C.; et al. Soybean adaption to high-latitude germplasm. BMC Genom. 2015, 16, 217. [CrossRef] [PubMed]

54. Liu, L.; Adrian, J.; Pankin, A.; Hu, J.; Dong, X.; von Korff, M.; Turck, F. Induced and natural variation of promoter length modulates the photoperiodic response of FLOWERING LOCUS T. Nat. Commun. 2014, 5, 4558. [CrossRef] [PubMed]

55. Rychel-Bielska, S.; Plewinski, P.; Kozak, B.; Galek, R.; Książkiewicz, M. Photoperiod and vernalization control of flowering-related genes: A case study of the narrow-leaved lupin (Lupinus angustifolius L.). Plant Physiol. 2017, 17, 820. [CrossRef] [PubMed]

56. Lichtin, N.; Salvo-Garrido, H.; Till, B.; Caligari, P.D.S.; Rupayan, A.; Westermeyer, F.; Olivos, M. Genetic and comparative mapping of Lupinus luteus L. highlight syntenic regions with major orthologous genes controlling anthracnose resistance and flowering time. Sci. Rep. 2020, 10, 19174. [CrossRef] [PubMed]

57. Laurie, R.E.; Diwadkar, P.; Jaudal, M.; Zhang, L.; Hecht, V.; Wen, J.; Talege, M.; Mysore, K.S.; Putterill, J.; Weller, J.L.; et al. The Medicago FLOWERING LOCUS T homolog, MtFTa1, is a key regulator of flowering time. Plant Physiol. 2011, 156, 2207–2224. [CrossRef] [PubMed]

58. Hecht, V.; Laurie, R.E.; Vander Schoor, J.K.; Rideg, S.; Knowles, C.L.; Liew, L.C.; Sussmilch, F.C.; Murfet, I.C.; Macknight, R.C.; Weller, J.L. The pea GIGAS gene is a FLOWERING LOCUS T homolog necessary for graft-transmissible specification of flowering but not for responsiveness to photoperiod. Plant Cell 2011, 23, 147–161. [CrossRef]
57. Samineni, S.; Kamatam, S.; Thudi, M.; Varshney, R.K.; Gaur, P.M. Vernalization response in chickpea is controlled by a major QTL. *Euphytica* 2016, 207, 453–461. [CrossRef]
58. Valverde, F.; Mouradov, A.; Sparre, W.; Ravenscroft, D.; Samach, A.; Coupland, G. Photoreceptor regulation of CONSTANS protein in photoperiodic flowering. *Science* 2004, 303, 1003–1006. [CrossRef]
59. Valverde, F. CONSTANS and the evolutionary origin of photoperiodic timing of flowering. *J. Exp. Bot.* 2011, 62, 2453–2463. [CrossRef] [PubMed]
60. Pierre, J.B.; Bogard, M.; Herrmann, D.; Huyghe, C.; Julier, B. Detection of QTLs for flowering date in three mapping populations of the model legume species *Medicago truncatula*. *Theor. Appl. Genet.* 2008, 117, 609–620. [CrossRef] [PubMed]
61. Pierre, J.B.; Lister, C.; Nordborg, M.; Dean, C. Role of photoperiod sensing in determining variation in flowering time of *Arabidopsis thaliana*. *Science* 2003, 302, 1751–1754. [CrossRef] [PubMed]
62. Lyu, J.; Cai, Z.; Li, Y.; Suo, H.; Li, R.; Zhang, S.; Nian, H. The floral repressor GmGBP1, a soybean homolog of the autonomous pathway gene FLOWERING LOCUS D, promotes flowering in *Arabidopsis thaliana*. *BMC Plant Biol.* 2014, 14, 263. [CrossRef] [PubMed]
63. Johanson, U.; West, J.; Lister, C.; Michaels, S.; Amasino, R.M.; Dean, C. Molecular analysis of FRIGIDA, a major determinant of natural variation in *Arabidopsis* flowering time. *Science* 2000, 290, 344–347. [CrossRef]
64. Upadhyaya, H.D.; Bajaj, D.; Das, S.; Saxena, M.S.; Badoni, S.; Kumar, V.; Tripathi, S.; Gowda, C.L.; Sharma, S.; Tyagi, A.K.; et al. A genome-scale integrated approach aids in genetic dissection of complex flowering time trait in chickpea. *Plant Mol. Biol.* 2015, 89, 403–420. [CrossRef]
65. He, Y.; Michaels, S.D.; Amasino, R.M. Regulation of flowering time by histone acetylation in *Arabidopsis*. *Science* 2003, 302, 1751–1754. [CrossRef] [PubMed]
66. Mouradov, A.; Cremer, F.; Coupland, G. Control of flowering time. *Plant Cell* 2002, 14, S111. [CrossRef] [PubMed]
67. Hu, Q.; Jin, Y.; Shi, H.; Yang, W. GmFLD, a soybean homolog of the autonomous pathway gene FLOWERING LOCUS D, promotes flowering in *Arabidopsis thaliana*. *BMC Plant Biol.* 2014, 14, 263. [CrossRef] [PubMed]
68. Li, Z.; Jiang, D.; He, Y. FRIGIDA establishes a local chromosomal environment for FLOWERING LOCUS C mRNA production. *Nat. Plants* 2018, 4, 836–846. [CrossRef]
69. Hecht, V.; Foucher, F.; Ferrandiz, C.; Macknight, R.; Navarro, C.; Morin, J.; Vardy, M.E.; Ellis, N.; Beltran, J.P.; Rameau, C.; et al. Conservation of Arabidopsis flowering genes in model legumes. *Plant Physiol.* 2005, 137, 1420–1434. [CrossRef]
70. Lyu, J.; Cai, Z.; Li, Y.; Suo, H.; Li, R.; Zhang, S.; Nian, H. The floral repressor GmFLC-like is involved in regulating flowering time mediated by low temperature in soybean. *Int. J. Mol. Sci.* 2020, 21, 1322. [CrossRef]
71. Li, M.; Liu, Y.; Tao, Y.; Xu, C.; Li, X.; Zhang, X.; Han, Y.; Yang, X.; Sun, J.; Li, W.; et al. Identification of genetic loci and candidate genes related to soybean flowering through genome wide association study. *BMC Genom.* 2019, 20, 987. [CrossRef] [PubMed]
72. Michaels, S.D.; Bezierra, I.C.; Amasino, R.M. FRIGIDA-related genes are required for the winter-annual habit in *Arabidopsis*. *Proc. Natl. Acad. Sci. USA* 2001, 104, 3281–3285. [CrossRef]
73. Zhu, M.L.; Yang, C.H. Late-flowering genes interact with early-flowering genes to regulate flowering time in *Arabidopsis thaliana*. *Plant Cell Physiol.* 1999, 40, 702–708. [CrossRef]
74. Pierre, J.B.; Huguet, T.; Barre, P.; Huyghe, C.; Julier, B. Detection of QTLs for flowering date in three mapping populations of the model legume species *Medicago truncatula*. *Theor. Appl. Genet.* 2008, 117, 609–620. [CrossRef] [PubMed]
75. Zhang, H.; Ransom, C.; Ludwig, P.; van Nocker, S. Genetic analysis of early flowering mutants in *Arabidopsis*. *New Phytol.* 2017, 21, 1420–1434. [CrossRef] [PubMed]
76. Wang, X.; Wu, F.; Xie, Q.; Wang, H.; Wang, Y.; Yue, Y.; Gahirra, O.; Ma, S.; Liu, L.; Cao, Y.; et al. SKIP controls flowering time via the alternative splicing of SEF pre-mRNA in *Arabidopsis*. *BMC Biol.* 2017, 15, 80. [CrossRef] [PubMed]
77. Zhao, L.; Li, M.; Xu, C.; Yang, X.; Li, D.; Zhao, X.; Wang, K.; Li, Y.; Zhang, X.; Liu, L.; et al. Natural variation in GmGBP1 promoter affects photoperiod control of flowering time and maturity in soybean. *Plant J.* 2018, 96, 147–162. [CrossRef] [PubMed]
78. Zhang, Y.; Zhao, L.; Li, H.; Gao, Y.; Li, Y.; Wu, X.; Teng, W.; Han, Y.; Zhao, X.; Li, W. GmGBP1, a homolog of human ski interacting protein in soybean, regulates flowering and stress tolerance in *Arabidopsis*. *BMC Plant Biol.* 2013, 13, 21. [CrossRef]
79. Brock, M.T.; Maloof, J.N.; Weining, C. Genes underlying quantitative variation in ecologically important traits: PIF4 (phytochrome interacting factor 4) is associated with variation in internode length, flowering time, and fruit set in *Arabidopsis thaliana*. *Mol. Ecol.* 2010, 19, 1187–1199. [CrossRef] [PubMed]
80. Kumar, S.V.; Lucyshyn, D.; Jaeger, K.E.; Alós, E.; Alvey, E.; Harberd, N.P.; Wigge, P.A. Transcription factor PIF4 controls the thermosensory activation of flowering. *Nature* 2012, 484, 242–245. [CrossRef] [PubMed]
81. Arya, H.; Singh, M.B.; Bhalla, P.L. Genomic analysis of conserved and unique features of soybean *PIF4*. *Sci. Rep.* 2018, 8, 12569. [CrossRef]
82. Karlgren, A.; Cyllenstrand, N.; Kallman, T.; Sundstrom, J.F.; Moore, D.; Lascoux, M.; Lagercrantz, U. Evolution of the PEBP gene family in plants: Functional diversification in seed plant evolution. *Plant Physiol.* 2011, 156, 1967–1977. [CrossRef]
85. Hedman, H.; Kallman, T.; Lagercrantz, U. Early evolution of the MFT-like gene family in plants. Plant Mol. Biol. 2009, 70, 359–369. [CrossRef] [PubMed]

86. Yoo, S.Y.; Kardailsky, I.; Lee, J.S.; Weigel, D.; Ahn, J.H. Acceleration of flowering by overexpression of MFT (MOTHER OF FT AND TFL1). Mol. Cells 2004, 17, 95–101.

87. Vaistij, F.E.; Barros-Galvão, T.; Cole, A.F.; Gilday, A.D.; He, Z.; Li, Y.; Harvey, D.; Larson, T.R.; Graham, I.A. MOTHER-OF-FT-AND-TFL1 represses seed germination under far-red light by modulating phytohormone responses in Arabidopsis thaliana. Proc. Natl. Acad. Sci. USA 2018, 115, 8442–8447. [CrossRef]

88. Ward, J.K.; Samanta Roy, D.; Chatterjee, I.; Bone, C.R.; Springer, C.J.; Kelly, J.K. Identification of a major QTL that alters flowering time at elevated [CO2] in Arabidopsis thaliana. PLoS ONE 2012, 7, e49028. [CrossRef] [PubMed]

89. Wei, M.; Xing, Y.; Wang, X.; Ma, H. [Variation of CO2 concentration in solar greenhouse in Northern China]. J. Appl. Ecol. 2003, 14, 354–358.

90. Caicedo, A.L.; Stinchcombe, J.R.; Olsen, K.M.; Schmitt, J.; Purugganan, M.D. Epistatic interaction between Arabidopsis FRI and FLC flowering time genes generates a latitudinal cline in a life history trait. Proc. Natl. Acad. Sci. USA 2004, 101, 15670–15675. [CrossRef] [PubMed]

91. Julier, B.; Huyghe, C.; Papineau, J.; Milford, G.F.J.; Day, J.M.; Billot, C.; Mangin, P. Seed yield and yield stability of determinate and indeterminate autumn-sown white lupins (Lupinus albus L.) grown at different locations in France and the UK. J. Agric. Sci. 1993, 121, 177–186. [CrossRef]

92. Julier, B.; Huyghe, C. Description and model of the architecture of four genotypes of determinate autumn-sown white lupin (Lupinus albus L.) as influenced by location, sowing date and density. Aust. J. Agric. Res. 1993, 44, 493–501. [CrossRef] [PubMed]

93. Alkemade, J.; Messmer, M.; Arncken, C.; Leska, A.; Anniciaricco, P.; Nazzari, N.; Kuijikwicz, M.; Voegele, R.T.; Finckh, M.; Hohmann, P. A high-throughput phenotyping tool to identify field-relevant anthracnose resistance in white lupin. Plant Dis. 2020, 104, 95–101. [CrossRef]

94. Altschul, S.F.; Gish, W.; Miller, W.; Myers, E.W.; Lipman, D.J. Basic local alignment search tool. J. Mol. Biol. 1990, 215, 403–410. [CrossRef]

95. Searle, S.R.; Casella, G.; McCulloch, C.E. Variance Components; John Wiley & Sons: New York, NY, USA, 2008.

96. Meuwissen, T.H.; Hayes, B.J.; Goddard, M.E. Prediction of total genetic value using genome-wide dense marker maps. Genetics 2001, 157, 1819–1829.

97. Darling, A.C.; Mau, B.; Blattner, F.R.; Perna, N.T. Mauve: Multiple alignment of conserved genomic sequence with rearrangements. Genome Res. 2004, 14, 1394–1403. [CrossRef] [PubMed]

98. Edgar, R.C. MUSCLE: Multiple sequence alignment with high accuracy and high throughput. Nucleic Acids Res. 2004, 32, 1792–1797. [CrossRef]

99. Frichet, E.; François, O. LEA: An R package for landscape and ecological association studies. Methods Ecol. Evol. 2015, 6, 925–929. [CrossRef] [PubMed]

100. Patterson, N.; Price, A.L.; Reich, D. Population structure and eigenanalysis. PLoS Genet. 2006, 2, e190. [CrossRef] [PubMed]

101. Pritchard, J.K.; Stephens, M.; Donnelly, P. Inference of population structure using multilocus genotype data. Genetics 2000, 155, 945–959. [PubMed]
113. Frichot, E.; Mathieu, F.; Trouillon, T.; Bouchard, G.; François, O. Fast and efficient estimation of individual ancestry coefficients. *Genetics* **2014**, *196*, 973–983. [CrossRef]

114. Alexander, D.H.; Lange, K. Enhancements to the ADMIXTURE algorithm for individual ancestry estimation. *BMC Bioinform.* **2011**, *12*, 246. [CrossRef]

115. Kendall, M.G. A New Measure of Rank Correlation. *Biometrika* **1938**, *30*, 81–93. [CrossRef]

116. Miller, R.G. Normal Univariate Techniques. In *Simultaneous Statistical Inference*, Springer: New York, NY, USA, 1981; pp. 37–108.