Isolation and Functional Analysis of a PISTILLATA-like MADS-Box Gene from Argan Tree (Argania spinosa)

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Abstract: Argan trees (Argania spinosa) belong to a species native to southwestern Morocco, playing an important role in the environment and local economy. Argan oil extracted from kernels has a unique composition and properties. Argan trees were introduced in Tunisia, where hundreds of trees can be found nowadays. In this study, we examined reproductive development in Argan trees from four sites in Tunisia and carried out the functional characterization of a floral homeotic gene in this non-model species. Despite the importance of reproductive development, nothing is known about the genetic network controlling flower development in Argania spinosa. Results obtained in several plant species established that floral organ development is mostly controlled by MADS-box genes and, in particular, APETALA3 (AP3) and PISTILLATA (PI) homologs are required for proper petal and stamen identity. Here, we describe the isolation and functional characterization of a MADS-box gene from Argania spinosa. Phylogenetic analyses showed strong homology with PI-like proteins, and the expression of the gene was found to be restricted to the second and third whorls. Functional homology with Arabidopsis PI was demonstrated by the ability of AsPI to confer petal and stamen identity when overexpressed in a pi-1 mutant background. The identification and characterization of this gene support the strong conservation of PI homologs among distant angiosperm plants.

Keywords: Argania spinosa; Argan tree; PISTILLATA; flower; MADS-box gene; Tunisia

1. Introduction

Argania spinosa L. Skeels is an evergreen xerophyte, agroforestry species that belongs to the tropical family of Sapotaceae [1]. This species is known to be endemic to Morocco and Algeria [2]. Despite the rarity of this species outside of these two countries, Argan trees have been spotted in different stations in Tunisia. This can be explained by several introductory trials in a few stations in Tunisia between 1963 and 1968 [3]. However, long before these introductory attempts, Emberger (1925) [4] mentioned the existence of Argan trees in Tunisia, around Kairouan, a station not validated by botanists at the time [4]. Taking into consideration the geographical location of Tunisia, the hypothesis of the endemism of the Argan tree in Tunisia remains probable.

Since the past centuries, this species has proved to be essential for the economy of Morocco due to the production of Argan oil [5]. Moreover, the leaves and fruits of the Argan tree can be used as forage, and the plant is considered multipurpose, which increases its socioeconomic value [6]. Other than their considerable socioeconomic value, Argan trees have a great ecological interest. This species has no requirements for the type of soil and tolerates a wide pH range and high concentration of limestone [7]. Furthermore, Argan trees improve the quality of soil and water around them [8], and the root structures of this
species and its high tolerance to aridity and high temperature make it a unique tool to counteract desertification [6]. The study of the argan tree population from Tunisia revealed large variability both within and between the studied sites [7]. The characterization and preservation of these populations could improve biodiversity in arid and semi-arid areas of Tunisia. In the long term, the effect on the ecosystem would be beneficial, and the establishment of the Argan tree can help the development of agriculture and the fight against wind erosion.

While many studies have been previously carried out to characterize the fruits, kernels, and leaves of the Argan [7,9–11], there is limited information concerning flower development [12–15]. Even more, to date, there is no available information of genes specifically involved in flower or fruit development in these species.

Flower development has been extensively studied in Arabidopsis and several model and non-model species, including crops. Genetics and molecular studies firmly established that floral organ development is genetically controlled by transcription factors encoded by genes from the MADS-box family [16,17]. Floral organs are arranged in concentric circles or whorls, and the number and disposition of the organs are characteristic of each species. MADS-box proteins interact in a combinatorial fashion to specify the identity of a particular floral organ. Thirty years ago, the ABC model was proposed and established three homeotic gene classes: A, B, and C [18]. Class A genes alone produce sepals, and in combination with B-class genes, regulate petal identity. Similarly, B and C function genes are required to establish stamen identity, and C functions alone to produce the carpel. Some years later, additional MADS-box genes were identified, resulting in two additional classes: D-class genes involved in carpel and ovule development [19,20] and E-class genes SEPALLATA genes that act redundantly in the control of all floral organs identities [21,22]. Altogether, the current model is known as the ABCDE model for flower development.

In Arabidopsis, petal and stamen development requires the presence of two B-class genes, APETALA3 (AP3) and PISTILLATA (PI). Accordingly, mutations in either of these genes result in similar phenotypes with homeotic transformations of petals into sepals and stamens into carpels [23,24]. These two proteins, encoded by AP3 and PI, function as obligate heterodimers to bind DNA and directly regulate different sets of genes at distinct phases of flower development [25]. Several B-class gene orthologs have been identified and functionally analyzed in ferns, gymnosperms, and different types of angiosperms, and the biochemical aspects of AP3 and PI function appear to be conserved [26]. In the basal angiosperms, the study of AP3 and PI lineages shows dynamic patterns of gene lineage evolution with complex events of gene duplication [27]. It seems as though the evolution of the floral MADS-box gene, and in particular, AP3/PI lineages, is connected to the evolution of the flower itself and a source of major innovation during land plant evolution and angiosperm diversification [26].

Here, we studied reproductive development in the non-model species Argania spinosa. First, we examined the floral diversity of a collection of Argan trees from different geographical locations in Tunisia, and second, we isolated a PI-like B-class gene that we show is involved in petal and stamen identity. At present, there are no functional analyses of genes in Argan, mostly due to the absence of a reference genome publicly available. The molecular characterization of this gene and the generation of overexpression Arabidopsis lines provide the first functional analysis of a PISTILLATA MADS-box gene in this species.
2. Results

2.1. Argan Flower Anatomy and Development

We took advantage of the available Argan population in Tunisia to study the diversity of flower disposition and number. Characterizing flower number variability in Argan populations could be interesting for the selection of desirable genotypes for future breeding programs in Tunisia. Observation of Argan trees at the four stations revealed a very high level of phenotypic variability during reproductive development. Four types of flower arrangements were observed: covering the whole shoot, helicoid, grouped as glomeruli in the axils of the leaves, and scattered (Figure 1a–d). The flowers appear on the axils of leaves or spines or on the nodes of mature wood. Argan tree is a monoecious species, with zygomorphic, bisexual, pentameric hermaphrodite flowers that have a pleasant smell [14]. The outermost whorls are composed of five hairy, rounded brownish sepal and five rounded greenish-yellow or white petals (Figure 1). The androecium is composed of five stamens forming a ring around the carpel that protrudes before flower opening (Figure 1e). Dissection of Argan flowers showed that the petals are attached to each other and to the stamens at the base and revealed the presence of five short staminodes (Figure 1f,h). Histological sections of flowers from different sites showed pistils with a single ovule, two ovules, or even three ovules (Figure 1j–m). The most frequent number of ovules per flower was two (average number of ovule per flower = 1.9 for botanical garden; 2.27 for Korbous; 1.66 for Sousse and 2.21 for Sfax) independently of the population analyzed.

In contrast, the four stations showed variability in terms of the number of flowers per tree and the disposition of flowers in the branches (Table 1). Regarding flower frequency per branch, we considered four phenotypical classes: high (>150), medium (50–150), low (1–50), and none (0). The most frequent number of flowers per branch in the entire tree population was medium (50–150 flowers/branch) and also in the individual sites. The exception was the trees from the Korbous site that showed more than 150 flowers/branch (Table 1). Regarding flower disposition, 43% of the trees analyzed showed flowers grouped as glomeruli, although we observed differences in flowers disposition inter and intra population. Helicoidal disposition of flowers is more frequent in trees from the Botanic Garden and Korbous. Remarkably, the percentage of scattered single flowers is higher in Sfax and Sousse than in Korbous and the botanical garden. The presence of flowers covering the whole shoot was infrequent (Table 1).

Table 1. Flowering phenology among trees from the four sites. Number of trees analyzed per site n = 15. The most frequent classes for every site and for the entire population are indicated in bold.

| Site                   | Flower Disposition (% of Trees) | Flower Frequency/Branch (% of Trees) |
|------------------------|---------------------------------|-------------------------------------|
|                        | Covering the Whole Shoot        | Helicoidal Disposition              | Grouped as Glomeruli | Scattered | >150 | 50–150 | 1–50 | 0    |
| Botanical Garden (BG)  | 13.3                            | 33.3                                | 40.0                  | 13.3      | 33.3 | 53.3   | 6.7  | 6.7  |
| Korbous (A)            | 13.3                            | 20.0                                | 46.7                  | 20.0      | 60.0 | 26.7   | 13.3 | 6.7  |
| Sousse (H)             | 0.0                             | 13.3                                | 46.7                  | 40.0      | 20.0 | 53.3   | 20.0 | 6.7  |
| Sfax (S)               | 6.7                             | 0.0                                 | 40.0                  | 53.3      | 20.0 | 33.3   | 33.3 | 13.3 |
| All trees              | 8.3                             | 16.7                                | 43.3                  | 31.7      | 33.3 | 41.7   | 18.3 | 8.3  |
2.2. Cloning and Sequence Analysis of a PISTILLATA Gene from Argania spinosa

To identify the PISTILLATA homolog in Argania spinosa, we searched DNA databases for PI coding sequences from trees or woody plants. We selected a subset of seven PI-like sequences previously characterized as B-function genes [28–32]. The alignment of the
sequences identified a fragment over 300 bp highly conserved among all sequences starting from the start codon and another shorter but similarly conserved region near the C-terminal end (Supplementary Figure S1). We then designed degenerated primers to amplify the first region, both primers containing variations in four of the positions (Supplementary Table S1). Using these primers and cDNA obtained from Argan flowers, we amplified and cloned a DNA fragment corresponding to a partial coding sequence of the AsPI (Argania spinosa PISTILLATA) gene.

By means of 5′ and 3′ RACE experiments, we obtained the full coding sequence of the AsPI gene (GenBank MW650858) that encoded a deduced protein of 208 aa (Supplementary Figure S2a). The predicted amino acid sequence contains three conserved regions annotated as MADS-box domain (1–61 aa), I-box (62–83 aa) and K-box domain (84–170 aa), and the consensus PI-motif (MPFxFRVQPxQNPxLQE) in the C-terminal region (Supplementary Figure S3), which is specifically found in PI orthologs [33]. Additionally, we obtained a collection of sequences of variable length corresponding to the 3′ UTR region (Supplementary Figure S2b). Arabidopsis and Argan PI proteins showed 62.0% total amino acid identity (85.6% similarity).

We used the coding sequence of the PI gene from Argan to perform phylogenetic analysis with AP3 and PI proteins from different plant species. We included functionally characterized PI proteins from distant phylogenetic plants. The topology of the tree showed that the AsPI gene is closely related to members of the PI/GLO subfamily (Figure 2). AsPI falls within the well-supported PI/GLO clade (bootstrap value of 100%). This clade is well defined and separated from the AP3/DEF clade (Figure 2).

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

**Figure 2.** AsPI is a PISTILLATA-like MADS-box protein. Phylogenetic reconstruction using the maximum likelihood (ML) method based on PI/GLO- and AP3/DEF-related MADS-box proteins from different species. The PI/GLO clade has been highlighted with a gray square. Argania spinosa PI (AsPI) protein is labeled in red. The number close to the nodes indicates the bootstrap support values from 10,000 pseudo-replicates. The accession numbers of the sequence data used are listed in Supplementary Table S2.

### 2.3. Expression Profiling of AsPI in Argan Flowers and Leaves

MADS-box genes involved in floral organ identity are expressed in distinct floral whorls. Class A genes are expressed in developing organ primordia of sepals and petals,
class B genes in petals and stamens, and class C genes in stamens and carpels. Therefore, we analyzed the expression pattern of AsPI in different floral organs and leaves by quantitative RT-PCR. In agreement with the expected expression pattern for a B-function gene, AsPI was expressed in petals and stamens/staminoids and barely detectable in sepals and carpels, and was not expressed in leaves (Figure 3), showing that AsPI expression is flower-specific. In summary, AsPI sequence analyses and the expression pattern of the gene correspond to a typical PISTILLATA MADS-box B-class gene.

Since PI proteins are transcription factors, they are expected to be localized into the nucleus. To investigate the subcellular localization of the AsPI protein, we produced a translational fusion of AsPI with the YFP fluorescent protein. The expression of the fusion protein was driven by a constitutive 35S promoter and transiently expressed in epidermal cells of N. benthamiana leaves (Figure 4). Plants expressing the 35S::GFP control showed strong fluorescence in the cytoplasm and nucleus. In contrast, we observed intense fluorescence of the AsPI-YFP protein exclusively in the nuclei (Figure 4). The analysis in silico of the protein indicated that the AsPI protein carries potential nuclear localization signals in its N-terminus (Supplementary Figure S2a).

**Figure 3.** Relative expression of AsPI in leaves and flower organs analyzed by qRT-PCR. Data were normalized to the expression of the ycf2 gene and correspond to the mean (±SD) of three replicates.

**Figure 4.** Subcellular localization of AsPI protein in Nicotiana benthamiana leaves as observed by confocal microscopy. Upper panel: cytoplasmic and nuclear localization of the 35S::GFP control construct. Lower panel: nuclear localization of the 35S::AsPI-YFP construct. From right to left, the three independent images correspond to bright field, GFP spectrum, and the overlay of both images.
2.5. AsPI Overexpression in Arabidopsis and pi-1 Mutant Complementation

To analyze the function of the isolated AsPI, we overexpressed the gene in the model plant Arabidopsis. We also tested whether AsPI could replace AtPI function in Arabidopsis by overexpression of the Argan gene in the pi-1 mutant background, a loss-of-function mutant allele with a strong floral phenotype [24,34].

We transformed a 35S::AsPI construct into heterozygous pi-1 plants. Twenty-four transformants were recovered in plates containing selection antibiotics and genotyped for the presence of the pi-1 mutation. A total of 12 of the transgenic lines were homozygous for the mutation, 10 were heterozygous, and 2 did not contain the mutation (Supplementary Figure S4). The expression of the transgene was analyzed in 11 lines (Figure 5g), of which 10 were homozygous for the pi-1 mutation and 1 (35S::AsPI #23) that did not contain the mutation.

![Figure 5](image_url)

Figure 5. Overexpression of AsPI in Arabidopsis plants and complementation of pi-1 floral phenotype. (a) Wild-type flower; (b) pi-1 flower; (c) Flower from 35S::AsPI transgenic plants (line #23); (d-f) Flowers from pi-1; 35S::AsPI plants (lines #5, #8 and #16) showing different degrees of complementation. (g) Relative expression of AsPI gene in the Arabidopsis 35S::AsPI transgenic lines. All the transgenic lines are homozygous for the pi-1 mutation except for line #23 that does not contain the mutation. Data were normalized to the expression of the AtTIP41 gene, and the gene expression level in line #11 was set as 1. Each value corresponds to the mean (±SD) of three replicates.

The ectopic expression of AsPI in a wild-type background modified flower anatomy as expected. We observed previously described phenotypes caused by overexpression of the Arabidopsis PI [35] and PI homologs [29,36], including strong separation between sepals, petals, and petals narrowing and sepals edge whitening (Figure 5C). The white areas in sepals are attributed to the partial homeotic transformation of sepals into petals [29,35,36]. Our results indicate that AsPI is able to induce B-function identity when ectopically expressed in Arabidopsis plants. In addition, we analyzed the floral phenotypes of the overexpressing lines in the pi-1 background. The flowers of the pi-1 mutant contain a first outer whorl of sepals, a second whorl of sepal-like organs, and an abnormally large gynoecium in the center of the flower. Stamens are missing, and, occasionally, filamentous structures appear [34,37] (Figure 5 and Table 2).

Among the overexpressing lines, we observed a low level of complementation in two of the lines (pi-1, 35S::AsPI #11 and pi-1, 35S::AsPI #20; Table 2) that correlate with the low expression level of AsPI detected in these plants (Figure 5g). The remaining lines showed full recovery of petals in the second whorl and partial recovery of staminoid structures in the third whorl (Figure 5 and Table 2). In the first whorl, we observed the characteristic white sepal edges that correspond to petaloid tissues, but the more conspicuous phenotype was the full recovery of white petals (Figure 5d–f). In the third whorl, we recorded an increase in the number of organs compared to the pi-1 mutant (Table 2). These organs correspond to filaments (92% of the flowers analyzed/line), filaments ending in stigmatic...
tissue (20%–30% of the flowers analyzed), and occasionally, filaments with anther tissue (10%–20% of the flowers analyzed) (Figure 5d–f and Table 2). In the innermost whorl, we observed many carpel abnormalities: opened, curved or incomplete carpels (Figure 5d–f) caused by the ectopic expression of AsPI in this tissue. In summary, the observed complementation of the pi-1 floral phenotype by AsPI confirms its ability to replace PI function.

### Table 2. Complementation of pi-1 floral phenotype by AsPI overexpression. Average number of organs per whorl and phenotypes observed (n = 10–20).

| Genotype      | 1st Whorl | 2nd Whorl | 3rd Whorl | 4th Whorl |
|---------------|-----------|-----------|-----------|-----------|
| Wild-type     | 4.00 ± 0.00 | 4.00 ± 0.00 | 3.68 ± 0.50 | 2.00 ± 0.00 |
| pi-1 mutant   | 4.00 ± 0.00 | 4.00 ± 0.00 | 0.19 ± 0.40 | 2.76 ± 0.54 |
| pi-1, 35S::AsPI #3 | 4.00 ± 0.00 | 4.00 ± 0.00 | 5.10 ± 0.88 | 2.30 ± 0.67 |
| pi-1, 35S::AsPI #5 | 4.00 ± 0.00 | 4.00 ± 0.00 | 5.00 ± 0.82 | 2.30 ± 0.48 |
| pi-1, 35S::AsPI #7 | 4.00 ± 0.00 | 4.00 ± 0.00 | 4.24 ± 1.60 | 2.12 ± 0.33 |
| pi-1, 35S::AsPI #8 | 4.00 ± 0.00 | 4.00 ± 0.00 | 2.57 ± 1.29 | 2.62 ± 0.59 |
| pi-1, 35S::AsPI #10 | 4.00 ± 0.00 | 4.00 ± 0.00 | 3.60 ± 1.64 | 2.10 ± 0.31 |
| pi-1, 35S::AsPI #11 | 4.00 ± 0.00 | 4.00 ± 0.00 | 1.00 ± 1.00 | 3.00 ± 0.00 |
| pi-1, 35S::AsPI #15 | 4.00 ± 0.00 | 4.00 ± 0.00 | 4.15 ± 1.31 | 2.30 ± 0.47 |
| pi-1, 35S::AsPI #16 | 4.00 ± 0.00 | 4.00 ± 0.00 | 4.60 ± 0.70 | 2.40 ± 0.52 |
| pi-1, 35S::AsPI #20 | 4.00 ± 0.00 | 4.00 ± 0.00 | 0.55 ± 0.83 | 2.45 ± 0.51 |
| pi-1, 35S::AsPI #21 | 4.00 ± 0.00 | 4.00 ± 0.00 | 3.60 ± 1.96 | 2.90 ± 0.99 |

In the wild-type, floral worlds correspond to sepals, petals, stamens, and carpels. Novel phenotypes observed that differ from morphological wild-type organs: a white sepal edges; b sepal-like organs; c petals; d filaments; f filaments fused to carpel; e filaments with stigmatic tissue; f anther tissue; g incomplete, bent or abnormal carpel.

### 3. Discussion

The analyses of the flower disposition and flower number of the argan tree population for Tunisia showed an important variability within and between sites. This could be the consequence of influence at different levels: geographic origin, genotype (tree/location), and genotype x environment interaction. The flowering-fruiting cycle covers a period of 9 to 16 months for the Argan trees in Morocco, from flower opening to fruit maturation [12].

Season was the main source of variation in the number of glomeruli on Argan shoots, and genotype x environment interaction. The flowering-fruiting cycle covers a period of 9 to 16 months for the Argan trees in Morocco, from flower opening to fruit maturation [12]. Season was the main source of variation in the number of glomeruli on Argan shoots, which may explain conflicting reports on flowering times in the literature [36]. Other studies associate the precocity of flowering with the precocity of precipitation [15,38].

Argan populations from sites characterized by a drier and hotter climate (Sfax) showed the lowest frequency of flower per branch and often as scattered single flowers. However, in the areas characterized by higher pluviometry (botanical garden and Korbous), we recorded the highest frequency of flowers. Besides the differences, we observed higher homogeneity among trees from Korbous, Sousse, and Sfax sites than within the botanical garden population. In addition, trees from Sousse and Sfax seem to be more closely related, in agreement with previous analyses [7,11].

In this study, we used a molecular approach to identify regulatory genes involved in floral organ development in Argan. We isolated and functionally characterized a MADS-box gene from Argania spinosa. Phylogenetic analysis showed that AsPI falls within a clade that includes several PI/GLO-like genes. In this group, Arabidopsis PI and Antirrhinum majus GLO (AmGLO) correspond to single genes while the petunia and Medicago counterparts correspond to duplicated genes, PhGLO1/PhGLO2 and MtPI/MtNGL9 in petunia and Medicago, respectively [29,33,39–43]. Gene duplications in the GLO/PI lineage have led to functional divergence and specialization of these MADS-box genes [29,41,44]. B-function MADS-box genes show multiple examples of gene duplication followed by gene loss or paralogs diversification, suggesting that gene duplication has been a crucial factor in shaping flower evolution [45,46]. Our analysis identified a single PI copy in the Argan genome, but we cannot fully exclude the presence of additional gene copies. Further analyses will...
be needed to support or discard the presence of duplicated genes in the PI/GLO-like lineage in this species.

Floral MADS proteins share a common modular conserved structure with four characteristic domains: MADS domain (M), intervening domain (I), keratin-like domain (K), and C-terminal (C) domains [47]. At the C-terminal region, most PI lineage proteins, including AsPI, harbored a highly conserved sequence of approximately 16-amino acid (PI-motif) considered essential for protein function [48]. However, additional research demonstrated that C-terminal motifs of AP3 and PI proteins are not required for floral organ identity specification [41,49,50]. The AsPI transcript, isolated from Argan flowers, encodes for a typical PI protein showing the classical MIKC structure and the presence of the conserved PI-motif. In lily (Lilium longiflorum), two PI genes were characterized, one of them lacking the PI-motif, and it was found that its presence was related to the ability of PI proteins to form homodimers [51].

AP3 and PI genes are present as a single copy in the Arabidopsis genome and are required for the specification of petals and stamens. Although their pattern of expression during flower development are similar, their single ectopic expression causes different phenotypic alterations. Overexpression of PI causes partial first whorl conversion of sepals to petals [37], whereas plants ectopically expressing AP3 exhibit partial conversion of carpels to stamens [52]. Simultaneous ectopic expression of both genes causes vegetative phenotypes (curling of leaves) and early flowering. In these plants, the flowers show stronger homeotic transformation on the first and fourth whorls; sepals tissues are absent on the outermost whorl, and carpelloid structures are rarely found in the innermost whorl [37].

Ectopic expression of AsPI in Arabidopsis wild-type plants phenocopies the effects described for the Arabidopsis PI gene. The flowers showed partial transformation of sepals into petaloid organs. In addition, sepals are widely separated in 35S::AsPI transgenic flowers, a phenotype associated with the ectopic expression in Arabidopsis of several PI orthologs from distant plant species such as Medicago truncatula [29], Catalpa bungei [35], Lilium longiflorum [51], among others. Interestingly, the ectopic expression of TrPI, a PI ortholog from the eudicot species Taihangia rupestris (Rosaceae) in Arabidopsis caused severe modifications in vegetative plant architecture [53]. Besides specifying the identities of petals and stamens, TrPI might function in regulating plant architecture in accordance with its expression in leaves and inflorescence stems [53]. We did not observe defects in vegetative development because of the ectopic expression of AsPI, suggesting a specific role for this gene during Argan flower development.

Previous studies in Arabidopsis showed that the simultaneous overexpression of PI and AP3 in different pi mutant alleles have different degrees of complementation of the mutant phenotype. Ectopic expression of PI and AP3 fully rescues a pi-2 mutant but only partially pi-1 or pi-3 mutant alleles that showed full recovery of petals identity but only mild recovery of stamens in addition to carpel defects [37]. The pi-1 allele was obtained by EMS (ethyl methanesulfonylate) mutagenesis and contained a point mutation that introduces a premature stop codon (TGG to TGA). The mutation results in a 79aa truncated protein and are considered a null allele [24]. We obtained similar results overexpressing AsPI in the pi-1 mutant background, confirming the ability of AsPI to confer B-class organ identity function in the Arabidopsis heterologous system.

The isolation and functional characterization of the AsPI gene is a pioneering study on the understanding of the molecular network controlling flower development in Argania spinosa. Our results support the functional conservation of MADS-box factors regulating the formation of the floral organs in angiosperms. Moreover, these findings provide molecular information for future studies in this species, a non-model plant with an ecologic value and local economic importance.
4. Materials and Methods

4.1. Plant Material

*Argania spinosa* flowers were collected from 60 Argan trees located in four stations in Tunisia: Tunis Botanical Garden, Korbous, Sousse, and Sfax (Table 3). The four locations have differences in altitude, climate, and soil composition [7]. The Argan trees located in Tunis at the botanical garden (Site 1) were introduced decades ago from Morocco. The rest of the sites contained trees from an unknown origin.

Table 3. Localization and general climatic conditions of the sites of the collection of Argan trees used in this study.

| Site Name and Code | Individual Code | Geographic Region | Bioclimatic Stage | Latitude | Longitude | Rainfall (mm/year) |
|--------------------|-----------------|-------------------|-------------------|----------|-----------|-------------------|
| Botanical Garden (BG) | P1A, P2F1, P3A2, P3A6, P3B8, P3C3, P4A8, P5F9, P6F9, P8G10, P9A11, A1JB, A3JB, A6JB, A8JB | North | Upper semi-arid | 36°81′ N 10°16′ E | 400–500 |
| Korbous (A) | A1, A2, A3, A4, A5, A6, A7, A8, A9, A10, A11, A12, A13, A14, A15 | Northeast (Cap Bon) | Sub-humid | 36°81′ N 10°56′ E | 500–700 |
| Sousse (H) | H1, H2, H3, H4, H5, H6, H7, H8, H9, H10, H11, H12, H13, H14, H15 | Central-East (Sahel) | Lower semi-Arid | 35°82′ N 10°64′ E | 300–400 |
| Sfax (S) | S1, S2, S3, S4, S5, S6, S7, S8, S9, S10, S11, S12, S13, S14, S15 | Southeast | Upper arid | 34°72′ N 10°33′ E | 100–200 |

Arabidopsis plants were grown in growth chambers at 21 °C under long-day (16 h light/8 dark) conditions, in a mixture of 1:1:1 soil:perlite:vermiculite. Wild-type *Arabidopsis thaliana* seedling (Landsberg erecta accession) and pistillata-1 (pi-1) mutant [24,48].

4.2. RNA Extraction and First-Strand cDNA Synthesis

Total RNA was isolated from Argan flower buds using Ribozol RNA extraction reagent (VWR Life Science, Solon, OH, USA) following the manufacturer’s instructions. After the extraction, we treated the samples with DNAase (Turbo DNA-free kit; Invitrogen, Waltham, MA, USA) to eliminate possible contamination with traces of genomic DNA. First-strand cDNA synthesis was performed by reverse transcription using 1 µg of total RNA and with the system Primer Script RT reagent kit (TaKaRa, Shiga, Japan) using a mix of poly-dT and random primers.

4.3. Cloning of AsPISTILLATA

A partial sequence of the *AsPI* gene was isolated from first-strand cDNA obtained from flower buds and using degenerated oligos PI-DEG For and PI-DEG Rev. The PCR product (321 bp) was purified and inserted into the pGEM-T Easy vector (Promega, Madison, WI, USA) for sequencing. The cloned partial cDNA sequence was used to design gene-specific primers to amplify the 5′ and 3′ end of cDNA using the SMARTer RACE 5′/3′ Kit (TaKaRa Bio, Shiga, Japan) according to the manufacturer’s instruction. The gene-specific primers were AsPI GSP2 for 3′ RACE and AsPI GSP7 for 5′ RACE. The sequence of the primers used is listed in Supplementary Table S1.

4.4. Phylogenetic Analyses

The evolutionary history was inferred by using the maximum likelihood method and Tamura-Nei model [54]. The tree with the highest log likelihood (−13,496.01) is shown. Initial tree(s) for the heuristic search were obtained automatically by applying Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated using the Tamura-Nei model and then selecting the topology with a superior log likelihood value. A discrete Gamma distribution was used to model evolutionary rate differences among sites (5 categories (+G, parameter = 1.8084)). The rate variation model allowed for some sites to be evolutionarily invariable ([+I], 9.65% sites). The tree is drawn to scale, with
branch lengths measured in the number of substitutions per site. The analysis involved 26 DNA sequences obtained from GenBank and the PI-like sequence that we isolated from A. spinosa. Codon positions included were 1st + 2nd + 3rd + Noncoding. There were a total of 803 positions in the final data set. Evolutionary analyses were conducted in MEGA X [55]. Sequences used in this analysis are listed in Supplementary Table S2.

4.5. Expression Analyses

For expression analyses, leaves and flower buds were collected. Flower buds were dissected under a stereomicroscope (Nikon, Tokyo, Japan) to separate floral worlds (sepals, petals, stamens/staminoids, and pistil) before total RNA extraction. First-strand cDNA synthesis was performed as previously described. The cDNA was diluted 10-fold with RNase-free water for qRT-PCR. Quantitative RT-PCR (qRT-PCR) was carried out with cDNA and Pyro Taq EvaGreen qPCR Mix (Cultek) using the QuantStudio 3 Real-Time PCR System (Applied Biosystems, Waltham, MA, USA). In a single experiment, each sample was assayed in triplicate (technical replicates).

For the expression analyses in Argan, the ycf2 gene (GenBank: JQ290464) was chosen as the internal control. For expression analyses in Arabidopsis, we used the TIP4 gene (At4g34270) as the internal control. The amplification efficiency of each set of primers was tested before the expression studies. Relative expression levels were calculated using the endogenous control genes as normalizers with the $2^{-\Delta\Delta Ct}$ method [56].

4.6. Subcellular Localization of AsPI Protein by Transient Expression in Nicotiana benthamiana

The complete AsPI cDNA was amplified using AsPI ATG For and AsPI Rev primers (Primers are listed in Supplementary Table S1) and cloned by recombination into the pEarleyGate 101 vector [57] to obtain an in-frame fusion to the fluorescent YFP protein. The final construct was confirmed by sequencing. Constructs containing AsPI-YFP fusion proteins were transformed into Agrobacterium tumefaciens C58/pMP90.

Overnight cultures of Agrobacterium containing the construct were diluted in infiltration buffer and used to infiltrate 4-week-old Nicotiana benthamiana leaves [58]. After 48 h of the infiltration, the localization of the fluorescent protein was determined on leave disks by confocal scanning microscopy (LSM 780, Zeiss, Jena, Germany) analyses.

4.7. Arabidopsis Transformation for AsPI Overexpression

For the overexpression of the AsPI protein, the coding sequence of the gene was cloned by recombination into de pEarleyGate100 vector [57] that placed the cDNA under the cauliflower mosaic virus 35S promoter. The final construct was confirmed by sequencing. Arabidopsis plants heterozygous for the pi-1 mutation were transformed by floral dipping according to standard procedures [59] after electrophorating the generated plasmid (35S::AsPI) into Agrobacterium strain C58/pMP90. Transformants were selected on phosphinotricin (PPT; Duchefa Biochemie, Haarlem, The Netherlands).

4.8. Histological Studies of Argan Flowers

For histological analyses, flower buds at different developmental stages were taken and fixed in FAE solution (3.7% formaldehyde, 5% acetic acid, 50% EtOH) overnight at 4 °C. The next day, FAE solution was removed and samples dehydrated in increasing concentrations of ethanol (70%, 80%, 90%, 95%, and 100%), cleared with Histo-Clear (VWR Life Science, Solon, OH, USA), and embedded in paraffin. Microtome sections of 8 µm were obtained and stained with 0.05% toluidine blue [60]. Stained slides were examined under a bright field using a Leica DM5000 light microscope equipped with a digital camera.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/plants10081665/s1, Figure S1. Alignment of partial cDNA sequences (360 nucleotides) of PISTILLATA genes from selected woody species and from the mode legume Medicago truncatula. Figure S2. PISTILLATA gene sequence. Figure S3. Protein alignment of AsPI and functional characterized PI/GLO-related proteins from different species. Figure S4: Genotyping of transgenic plants
transformed with 35S::AsPI for the presence of the pi-1 mutation. Table S1: Primers used in this work. Table S2: Accession numbers of the AP3/DEF- and PI/GLO-related MADS-box genes used for phylogenetic analysis.

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**References**

1. O’Brien, T.P.; Feder, N.; McCully, M.E. Polychromatic staining of plant cell walls by toluidine blue O. *Protoplasma* **1964**, *59*, 368–373. [CrossRef]

2. Guillaume, D.; Pioch, D.; Charrouf, Z. Argan [Argania spinosa (L.) Skeels] Oil; Springer: Berlin/Heidelberg, Germany, 2019; ISBN 9783030124724.

3. Kechebar, M.S.A.; Karoune, S.; Belhamra, M.; Rahmoune, C. Etude structurale des peuplements d’arganier (*Argania spinosa*) en Algerie. *J. Algérien Régions Arid.* **2013**, *46*, 54–.

4. Adib, S.E.L.; Slim, S.; Jeddi, F.B. Etude de la dynamique de la colonisation mycorhizienne de deux variéités d’arganier en Tunisie. *J. New Sci.* **2015**, *17*, 603–614.

5. Emberger, L. A Propos de La Distribution Géographique de l’arganier. *Bull. Soc. Sci. Nat. Phys. Maroc* **1924**, *4*, 151–153.

6. Metougui, M.L.; Mokhtari, M.; Maughan, P.F.; Jellen, E.N.; Benlhhabib, O. Morphological Variability, Heritability and Correlation Studies within an Argan Tree Population (*Argania spinosa* (L.) Skeels) Preserved in situ. *Int. J. Agric. For.* **2017**, *7*, 42–51. [CrossRef]

7. Mapelli, F.; Riva, V.; Vergani, L.; Choukrallah, R.; Borin, S. Unveiling the Microbiota Diversity of the Xerophyte *Argania spinosa* L. Skeels Root System and Residuesphere. *Microb. Ecol.* **2020**, *80*, 822–836. [CrossRef] [PubMed]

8. Louati, M.; Ucarli, C.; Arikan, B.; Ghada, B.; Hannachi, A.S.; Turgut-Kara, N.; Kara, T. Genetic, Morphological, and Biochemical Diversity of Argan Tree (*Argania spinosa* L.) as Case Study. *J. Food Sci. Technol.* **2013**, *12*, 747–754. [CrossRef]

9. Moukrim, S.; Lahssini, S.; Rhazi, M.; Alaloui, H.M.; Benabou, A.; Wahby, I.; El Madili, M.; Arahou, M.; Rhazi, L. Climate change impacts on potential distribution of multipurpose agro-forestry species: *Argania spinosa* (L.) Skeels as case study. *Agrofor. Syst.* **2018**, *93*, 1209–1219. [CrossRef]

10. Charrouf, Z.; Guillaume, D. Argan oil: Occurrence, composition and impact on human health. *Eur. J. Lipid Sci. Technol.* **2008**, *110*, 632–636. [CrossRef]

11. El Kharrassi, Y.; Maata, N.; Mazri, M.A.; El Kamouni, S.; Talbi, M.; El Kebjaj, R.; Moustaid, K.; Essamadi, A.K.; Andreoletti, P.; El Mzouri, E.H.; et al. Chemical and phytochemical characterizations of argan oil (*Argania spinosa* L. skeels), olive oil (Olea europaea L. cv. Moroccan picholine), cactus pear (*Opuntia megacantha* salmy-dyck) seed oil and cactus cladode essential oil. *J. Food Mass. Charact.* **2017**, *12*, 747–754. [CrossRef]

12. Louati, M.; Khouja, A.; Ben Abdelkrim, A.; Hannachi, A.S.; Baraket, G. Adaptation of *Argania spinosa* L. in Northern Tunisia: Soil analysis and morphological traits variability. *Sci. Hortic.* **2019**, *255*, 220–230. [CrossRef]

13. Benlahbil, S.; Zahidi, A.; Bani-Aameur, F.; El Mousadik, A. Duration of Blossoming, Longevity of Argan Flower and Assessment of Pollen Fertility by Staining. *Int. J. Agric. For.* **2015**, *5*, 291–304. [CrossRef]

14. Zahidi, A.; Bani-Aameur, F.; El Mousadik, A. Morphological variability of the fruiting branches in *Argania spinosa*: Effects of seasonal variations, locality and genotype. *J. Hortic. For.* **2013**, *5*, 168–182. [CrossRef]

15. Bani-Aameur, F. Phenological phases of *Argania spinosa* (L. Skeels) flower. *For. Genet.* **2000**, *7*, 329–334.

16. Bani-Aameur, F. *Argania spinosa* (L.) Skeels flowering phenology. *Genet. Resour. Crop. Evol.* **2002**, *49*, 11–19. [CrossRef]

17. Yan, W.; Chen, D.; Kaufmann, K. Molecular mechanisms of floral organ specification by MADS domain proteins. *Curr. Opin. Plant Biol.* **2016**, *29*, 154–162. [CrossRef]

18. Ali, Z.; Raza, Q.; Atif, R.M.; Aslam, U.; Ajmal, M.; Chung, G. Genetic and Molecular Control of Floral Organ Identity in Cereals. *Int. J. Mol. Sci.* **2019**, *20*, 2743. [CrossRef] [PubMed]
Plants 2021, 10, 1665

19. Coen, E.S.; Meyerowitz, E.M. The war of the whorls: Genetic interactions controlling flower development. *Nat. Cell Biol.* 1991, 335, 31–37. [CrossRef]

20. Favaro, R.; Pinyopich, A.; Battaglia, R.; Kooiker, M.; Borghi, L.; Ditta, G.; Yanofsky, M.F.; Kater, M.; Colombo, L. MADS-Box Protein Complexes Control Carpel and Ovule Development in Arabidopsis. *Plant Cell* 2003, 15, 2603–2611. [CrossRef] [PubMed]

21. Angenent, G.C.; Franken, J.; Busscher, M.; Van Dijken, A.; Van Went, J.L.; Dons, H.J.; Van Tunen, A.J. A novel class of MADS box genes is involved in ovule development in petunia. *Plant Cell* 1995, 7, 1569–1582. [CrossRef]

22. Pelaz, S.; Ditta, G.S.; Baumann, E.; Wisman, E.; Yanofsky, M.F. B and C floral organ identity functions require *SEPALATA* MADS-box genes. *Nat. Cell Biol.* 2000, 405, 200–203. [CrossRef] [PubMed]

23. Ditta, G.; Pinyopich, A.; Robles, P.; Pelaz, S.; Yanofsky, M.F. The *SEP4* Gene of Arabidopsis Italianna Functions in Floral Organ and Meristem Identity. *Curr. Biol.* 2004, 14, 1935–1940. [CrossRef]

24. Bowman, J.; Smyth, D.; Meyerowitz, E.M. Genes directing flower development in Arabidopsis. *Plant Cell* 1989, 1, 37–52. [CrossRef] [PubMed]

25. Goto, K.; Meyerowitz, E.M. Function and regulation of the Arabidopsis floral homeotic gene *PISTILLATA*. *Genes Dev.* 1994, 8, 1548–1560. [CrossRef]

26. Wuest, S.; O’Maioleidigh, D.; Rae, L.; Kwasniewska, K.; Raganelli, A.; Hanczaryk, K.; Lohan, A.; Loftus, B.; Graciet, E.; Wellmer, F. Molecular basis for the specification of floral organs by *APETALA3* and *PISTILLATA*. *Proc. Natl. Acad. Sci. USA* 2012, 109, 13452–13457. [CrossRef]

27. Theissen, G.; Becker, A.; Di Rosa, A.; Kanno, A.; Winter, K.-U.; Saedler, H. A short history of MADS-box genes in plants. *Plant Mol. Biol.* 2000, 42, 115–149. [CrossRef]

28. Stellari, G.M.; Jaramillo, M.A.; Kramer, E.M. Evolution of the *APETALA3* and *PISTILLATA* Lineages of MADS-Box–Containing Genes in the Basal Angiosperms. *Mol. Biol. Evol.* 2004, 21, 506–519. [CrossRef] [PubMed]

29. de Oliveira, R.R.; Cesarino, I.; Mazzaferra, P.; Dornelas, M.C. Flower development in *Coffea arabica L.*: New insights into MADS-box genes. *Plant Reprod.* 2014, 27, 79–94. [CrossRef]

30. Roque, E.; Fares, M.A.; Yenush, L.; Rochina, M.C.; Wen, J.; Mysore, K.; Gómez-Mena, C.; Beltrán, J.P.; Cañas, L.A. Evolution by gene duplication of *Medicago truncatula* PISTILLATA-like transcription factors. *J. Exp. Bot.* 2016, 67, 1805–1817. [CrossRef]

31. Zhang, W.; Xiang, Q.-Y.; Thomas, D.T.; Wiegmann, B.; Frohlich, M.W.; Solits, U.E. Molecular evolution of *PISTILLATA*-like genes in the dogwood genus *Cornus* (Cornaceae). *Mol. Phylog. Evol.* 2008, 47, 175–195. [CrossRef]

32. Zhang, B.; Su, X.; Zhou, X. A MADS-box gene of *Populus deltoides* expressed during flower development and in vegetative organs. *Tree Physiol.* 2008, 28, 929–934. [CrossRef]

33. Ge, J.; Zhao, D.; Han, C.; Wang, J.; Hao, Z.; Tao, J. Cloning and expression of floral organ development-related genes in herbaceous peony (*Paeonia lactiflora* Pall.). *Mol. Biol. Rep.* 2014, 41, 6493–6503. [CrossRef] [PubMed]

34. Kramer, E.M.; Dorit, R.L.; Irish, V.F. Molecular Evolution of Genes Controlling Petal and Stamens Development: Duplication and Divergence Within the *APETALA3* and *PISTILLATA* MADS-Box Gene Lineages. *Genetics* 1998, 149, 765–783. [CrossRef] [PubMed]

35. Krizek, B.; Meyerowitz, E. The Arabidopsis homeotic genes *APETALA3* and *PISTILLATA* are sufficient to provide the B class organ identity function. *Development* 1996, 122, 11–22. [CrossRef]

36. Jing, D.; Xia, Y.; Chen, F.; Wang, Z.; Zhang, S.; Wang, J. Ectopic expression of a *Catalpa bungei* (Bignoniaceae) *PISTILLATA* homologue rescues the petal and stamen identities in Arabidopsis *pi-1* mutant. *Plant Sci.* 2015, 231, 40–51. [CrossRef]

37. Bowman, J.L.; Smyth, D.R.; Meyerowitz, E.M. Genetic interactions among floral homeotic genes of Arabidopsis. *Development* 1991, 112, 1–20. [CrossRef]

38. Ferradossi, A.; Bani-Aameur, F.; Dupuis, P. Station climate, phenology and fruiting of Argan tree (*Argania spinosa* (L.) Skeels). *Actes Inst. Agron. Veto* 2016, 17, 59–60.

39. Rathcke, B.; Lacey, E.P. Phenological Patterns of Terrestrial Plants. *Annu. Rev. Ecol. Syst.* 1985, 16, 179–214. [CrossRef]

40. Angenent, G.C.; Busscher, M.; Franken, J.; Mol, J.N.; Van Tunen, A.J. Differential expression of two MADS box genes in wild-type and mutant petunia flowers. *Plant Cell* 1992, 4, 983–993. [CrossRef]

41. Angenent, G.C.; Franken, J.; Busscher, M.; Colombo, L.; Tunen, A.J. Petal and stamen formation in petunia is regulated by the homeotic gene *fbp1*. *Plant, J.* 1993, 4, 101–112. [CrossRef]

42. Benilch, R.; Roque, E.; Ferrández, C.; Cosson, V.; Caballero, T.; Pennmetsa, R.V.; Beltrán, J.P.; Cañas, L.; Ratep, P.; Madueño, F. Analysis of B function in legumes: *PISTILLATA* proteins do not require the PI motif for floral organ development in *Medicago truncatula*. *Plant J.* 2009, 60, 102–111. [CrossRef]

43. Kramer, E.M.; Irish, V.F. Evolution of the Petal and Stamens Developmental Programs: Evidence from Comparative Studies of the Lower Eudicots and Basal Angiosperms. *Int. J. Plant Sci.* 2000, 161, S29–S40. [CrossRef]

44. Tröbner, W.; Ramírez, L.; Mottet, P.; Hue, I.; Huijser, P.; Lönnig, W.E.; Saedler, H.; Schwarz-Sommer, Z. GLOBOSA: A homeotic gene which interacts with DEFICIENS in the control of *Antirrhinum* floral organogenesis. *EMBO J.* 1992, 11, 4693–4704. [CrossRef]

45. Vandenbussche, M.; Zethof, J.; Royaert, S.; Weterings, K.; Gerats, T. The Duplicated B-Class Heterodimer Model: Whorl-Specific Effects and Complex Genetic Interactions in *Petunia hybrida* Flower Development. *Plant Cell* 2004, 16, 741–754. [CrossRef]

46. Airoldi, C.; Davies, B. Gene Duplication and the Evolution of Plant MADS-box Transcription Factors. *J. Genet. Genom.* 2012, 39, 157–165. [CrossRef] [PubMed]
47. Alvarez-Buylla, E.R.; Pelaz, S.; Liljegren, S.J.; Gold, S.E.; Burgess, C.; Ditta, G.S.; de Pouspana, L.R.; Martínez-Castilla, L.; Yanofsky, M.F. An ancestral MADS-box gene duplication occurred before the divergence of plants and animals. *Proc. Natl. Acad. Sci. USA* 2000, 97, 5328–5333. [CrossRef] [PubMed]

48. Theißen, G. Development of floral organ identity: Stories from the MADS house. *Curr. Opin. Plant Biol.* 2001, 4, 75–85. [CrossRef]

49. Lamb, R.S.; Irish, V.F. Functional divergence within the APETALA3/PISTILLATA floral homeotic gene lineages. *Proc. Natl. Acad. Sci. USA* 2003, 100, 6558–6563. [CrossRef]

50. Berbel, A.; Navarro, C.; Ferrándiz, C.; Cañas, L.; Beltrán, J.-P.; Madueño, F. Functional Conservation of PISTILLATA Activity in a Pea Homolog Lacking the PI Motif. *Plant Physiol.* 2005, 139, 174–185. [CrossRef]

51. Piwarzyk, E.; Yang, Y.; Jack, T. Conserved C-Terminal Motifs of the Arabidopsis Proteins APETALA3 and PISTILLATA Are Dispensable for Floral Organ Identity Function. *Plant Physiol.* 2007, 145, 1495–1505. [CrossRef] [PubMed]

52. Chen, M.-K.; Hsieh, W.-P.; Yang, C.-H. Functional analysis reveals the possible role of the C-terminal sequences and PI motif in the function of lily (*Lilium longiflorum*) PISTILLATA (PI) orthologues. *J. Exp. Bot.* 2011, 63, 941–961. [CrossRef]

53. Jack, T.; Fox, G.L.; Meyerowitz, E.M. Arabidopsis homeotic gene APETALA3 ectopic expression: Transcriptional and posttranscriptional regulation determine floral organ identity. *Cell* 1994, 76, 703–716. [CrossRef]

54. Lü, S.; Fan, Y.; Liu, L.; Liu, S.; Zhang, W.; Meng, Z. Ectopic expression of TrPI, a *Taihangia rupestris* (Rosaceae) PI ortholog, causes modifications of vegetative architecture in Arabidopsis. *J. Plant Physiol.* 2010, 167, 1613–1621. [CrossRef]

55. Tamura, K.; Nei, M. Estimation of the number of nucleotide substitutions in the control region of mitochondrial DNA in humans and chimpanzees. *Mol. Biol. Evol.* 1993, 10, 512–526. [CrossRef]

56. Kumar, S.; Stecher, G.; Li, M.; Knyaz, C.; Tamura, K. MEGA X: Molecular Evolutionary Genetics Analysis across Computing Platforms. *Mol. Biol. Evol.* 2018, 35, 1547–1549. [CrossRef] [PubMed]

57. Livak, K.J.; Schmittgen, T.D. Analysis of relative gene expression data using real-time quantitative PCR and the 2\(^{-\Delta\Delta Ct}\) Method. *Methods* 2001, 25, 402–408. [CrossRef]

58. Earley, K.W.; Haag, J.R.; Pontes, O.; Opper, K.; Juehne, T.; Song, K.; Pikaard, C. Gateway-compatible vectors for plant functional genomics and proteomics. *Plant J.* 2006, 45, 616–629. [CrossRef] [PubMed]

59. Sparkes, I.A.; Runions, J.; Kearns, A.; Hawes, C. Rapid, transient expression of fluorescent fusion proteins in tobacco plants and generation of stably transformed plants. *Nat. Protoc.* 2006, 1, 2019–2025. [CrossRef] [PubMed]

60. Clough, S.J.; Bent, A. Floral dip: A simplified method for *Agrobacterium*-mediated transformation of *Arabidopsis thaliana*. *Plant J.* 1998, 16, 735–743. [CrossRef]