The Arabidopsis H3K27me3 demethylase JUMONJI 13 is a temperature and photoperiod dependent flowering repressor

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In plants, flowering time is controlled by environmental signals such as day-length and temperature, which regulate the floral pathway integrators, including FLOWERING LOCUS T (FT), by genetic and epigenetic mechanisms. Here, we identify an H3K27me3 demethylase, JUMONJI 13 (JMJ13), which regulates flowering time in Arabidopsis. Structural characterization of the JMJ13 catalytic domain in complex with its substrate peptide reveals that H3K27me3 is specifically recognized through hydrogen bonding and hydrophobic interactions. Under short-day conditions, the jmj13 mutant flowers early and has increased FT expression at high temperatures, but not at low temperatures. In contrast, jmj13 flowers early in long-day conditions regardless of temperature. Long-day condition and higher temperature induce the expression of JMJ13 and increase accumulation of JMJ13. Together, our data suggest that the H3K27me3 demethylase JMJ13 acts as a temperature- and photoperiod-dependent flowering repressor.
Varying day-length (photoperiod) and ambient temperature are two environmental cues that play central roles in plant development, including the regulation of flowering time. As a facultative long-day (LD) plant, Arabidopsis thaliana flowers earlier under LD conditions compared to short-day (SD) conditions. Higher ambient temperature also promotes flowering and there is considerable crosstalk between the photoperiodic pathway and the ambient temperature pathway. These environmental cues regulate the expression of key flowering integrators such as FLOWERING LOCUS T (FT)\(^1\). In the photoperiodic pathway, FT is regulated by the oscillating expression of the transcription factor CONSTANS (CO), which integrates circadian clock and day-length signals\(^2\,\,^8\). In the ambient temperature pathway, the MADS-box domain proteins Flowering Locus M (FLM) and Short Vegetative Phase (SVP) are involved in regulating FT expression and thus flowering time\(^7\,\,^8\). SVP protein is degraded and its level declines as temperature increases\(^8\). FLM responds to ambient temperature changes by switching between protein isoforms FLM-β and FLM-δ\(^9\,\,^10\). FLM-β forms a repressive complex with SVP to prevent flowering, whereas the dominant negative FLM-δ forms an SVP–FLM-β complex that lacks DNA binding and therefore repressors activity, allowing the activation of flowering\(^11\,\,\,12\). The bHLH transcription factor PHYTOCHROME INTERACTING FACTOR4 (PIF4) binds directly to the FT promoter in a temperature-dependent manner, and strong binding of PIF4 to FT depends on the eviction of H2A.Z nucleosomes induced by high temperature\(^13\,\,\,14\). Furthermore, the MYB transcription factor protein Early Flowering MYB Protein (EFM) acts as a convergence point for temperature and light regulation of flowering\(^15\).

Epigenetic regulators modulate chromatin conformation and composition, thus affecting the expression of key flowering integrators\(^16\). Histone methylation, which has important roles in transcriptional regulation and genome integrity\(^17\), is written by histone methyltransferases and erased by histone demethylases\(^18\). The highly diverse jumonji domain-containing histone demethylases are classified into subfamilies based on their catalytic domain sequence\(^19\,\,\,20\). Demethylases from each subfamily target specific substrates and perform distinct functions. For example, the human Lysine (K)-Specific Demethylase 4 (KDM4), KDM5, and KDM6 subfamilies are H3K9me3/H3K36me3-specific, H3K4me3-specific, and H3K27me3-specific demethylases, respectively\(^20\). In plants, histone demethylases have plant-specific features and different evolutionary relationships compared with their animal counterparts\(^19\). For example, plants do not possess the KDM6 subfamily H3K27me3 demethylases\(^19\). Instead, two known plant H3K27me3 demethylases, EARLY FLOWERING 6 (ELF6)/JUMONJI 11 (JMJ11) and RELATIVE OF EARLY FLOWERING 6 (REF6)/JUMONJI 12 (JMJ12), show sequence similarities to the human H3K9me3/H3K36me3 bi-specific KDM4 subfamily demethylases\(^19\,\,\,20\). The elf6 and ref6 loss-of-function mutants display early and late flowering phenotypes, respectively\(^21\), suggesting that various plant H3K27me3 demethylases influence flowering via different pathways. REF6 genome-wide DNA-binding requires four tandem Cys2His2 zinc fingers and functions to counteract Polycomb-mediated gene silencing\(^22\,\,\,24\).

Here we investigate the role of histone demethylases in plant flowering through the analysis of Arabidopsis JMJ13, which we show possesses H3K27me3 site-specific demethylase activity in vitro and in vivo. We further determined the crystal structure of JMJ13 in peptide-free and H3K27me3 peptide-bound forms. JMJ13 possesses a unique C4HCHC-type zinc finger, and not the previously predicted C5HC2-type zinc finger, despite the two zinc finger types sharing a similar folding topology. The substrate H3K27me3 peptide is specifically recognized by hydrogen bonding and hydrophobic stacking interactions, providing detailed structural insight into the substrate specificity of a plant H3K27me3 demethylase. In addition, we show that JMJ13 plays a role in temperature- and day-length-regulated flowering. Impaired JMJ13 function leads to early flowering in both LD and SD conditions at high temperature, but not in SD conditions at low temperature. Our genetic studies suggest that JMJ13 acts as a flowering repressor, which regulates flowering time in a temperature- and photoperiod-dependent manner.

**Results**

**JMJ13 specifically demethylates H3K27me3.** We previously identified 21 JmJC domain-containing proteins in the Arabidopsis thaliana genome and predicted JMJ13 as one of the 15 potentially active histone demethylases\(^19\). JMJ13 is a homolog of ELF6/JMJ11 and REF6/JMJ12, the two Arabidopsis KDM4 subfamily H3K27me3 demethylases (Supplementary Table 1)\(^23\,\,\,26\). To determine whether JMJ13 is an active demethylase, we performed enzymatic activity assays in vivo using a Nicotiana benthamiana leaf-based assay\(^25\,\,\,27\) (Fig. 1a). In cells where JMJ13-GFP was over-expressed, H3K27me3, but not H3K27me2 and H3K27me1, was markedly reduced (Fig. 1b, c). In contrast, there were no significant differences in the tri-, di- and mono-methylation levels of H3K4, H3K9, or H3K36 sites (Supplementary Fig. 1). The H3K27me3 demethylase activity of JMJ13-GFP was abolished when His293 and Glu295, the two conserved iron-binding amino acids, were replaced by alanine (Fig. 1d, e).

We further performed a MALDI-TOF mass-spectrometry-based in vitro demethylase assay\(^24\). The recombinant expressed JMJ13 catalytic domain (JMJ13CD, residues 90–578, Fig. 2a) displayed unambiguous demethylation activity against H3K27me3 peptides, but not H3K4me3, H3K9me3, or H3K36me3 peptides, confirming that JMJ13 is an H3K27me3 site-specific histone demethylase (Supplementary Fig. 2a–d). Further assays show that JMJ13 has high H3K27me3 demethylase activity, but no significant activity on H3K27me2 and H3K27me1 (Supplementary Fig. 2d–f). Together, the in vivo and in vitro results demonstrate that JMJ13 is predominately an H3K27me3-specific demethylase.

**Crystal structure of JMJ13CD.** To understand the mechanism of H3K27me3-specific demethylation by JMJ13, we performed structural studies. JMJ13 has a central catalytic domain flanked by flexible regions on both N- and C-termini (Fig. 2a). The crystal structure of the JMJ13 catalytic domain (JMJ13CD), including the predicted jumonji, helical, and zinc finger domains, in complex with the co-factor α-ketoglutarate (α-KG), was determined to 2.4 Å resolution (Fig. 2b and Supplementary Table 2). Despite being in different subfamilies, the overall structure of JMJ13CD resembles the previously reported structures of human KDM5A/B/C and Arabidopsis JMJ14, which are composed of two parts: the jumonji domain and the helical plus zinc finger domains\(^24\,\,\,28\,\,\,29\) (Fig. 2b). The jumonji domain adopts a typical α-KG-dependent oxygenase fold with a double-stranded β-helix in the center surrounded by several α-helices (Fig. 2b). A Ni\(^{2+}\) ion, which replaced the endogenous Fe\(^{2+}\) ion during nickel column purifcation, and the co-factor α-KG are located in the active center of the jumonji domain (Fig. 2b). The helical domain consists of four long α-helices forming a helical bundle (Fig. 2b).

JMJ13 and other members of the KDM5 subfamily of histone demethylases are predicted to possess a C5HCHC-type (5 cysteine residues followed by a histidine and 2 additional cysteine residues) zinc finger domain to coordinate two Zn\(^{2+}\) ions. However, our structural analysis showed that the predicted C5HCHC zinc finger of JMJ13 in fact adopts a C4HCHC arrangement, which is composed...
of a CCCH-type zinc finger (Cys500, Cys503, Cys522, and His525) and a CCHC-type zinc finger (Cys514, Cys516, His519, and Cys534) (Fig. 2b). Rather than the predicted Cys507, the His525 contributes to zinc coordination in the C4HCHC-type zinc finger domain in JMJ13. Although the C4HCHC zinc finger domain is embedded in the primary sequence of the helical domain,
structurally it forms an independent domain that interacts with the helical domain (Fig. 2a–b) in a similar manner to the helical and C5HC2 domains of JM1434.

Recognition of the H3K27me3 peptide by JM13. To investigate the substrate recognition and catalytic mechanism of JM13, we determined the crystal structure of JM13CD in complex with the α-KG analog N-oxalylglycine (NOG) and an H3K27me3 peptide, at 2.6 Å resolution (Fig. 2c and Supplementary Table 2). Overall the structure closely resembles the JM13-α-KG complex with a root-mean-square deviation (RMSD) of 0.49 Å (Supplementary Fig. 3a). The peptide can be traced from H3A24 to H3A31 (Supplementary Fig. 3b). The peptide binds in a negatively charged cleft with the residues H3K27me3 to H3P30 located at the bottom of the cleft and the other flanking residues extending out from the cleft (Fig. 2d). The side chain of H3K27me3 inserts into a deep binding pocket within the active site cleft (Fig. 2d). An NOG molecule and a Ni²⁺ ion are deeply buried in the center of the active site (Fig. 2c).

The interactions between JM13 and the H3K27me3 peptide are restricted to the region between H3R26 and H3P30. H3R26 forms salt bridge and hydrogen bonding interactions with Asp236 of JM13 (Fig. 3a). H3S28 forms a side-chain hydrogen bond with Asp296 of JM13 (Fig. 3a). H3P30 positions its side chain prolyl ring such that the plane of the propyl ring is parallel with and stacks on top of the phenyl ring of Phe179 of JM13 (Fig. 3a), resulting in hydrophobic stacking and CH–π interactions. Compared with the JM13 peptide-free structure, the side chain of Phe179 undergoes a significantly rotation to allow stacking with H3P30, indicative of a peptide binding-induced conformational change (Fig. 3b).

The active site of JM13 displays features typical of α-KG-dependent oxygenases observed for other jumonji domain histone demethylases (Fig. 3c)20,30. The trimethyllysine docks into the deep binding pocket with the three methyl groups anchored by an extensive CH–O hydrogen-bonding network (Fig. 3c), which is essential for fixing the conformation of the head group of the bound trimethyllysine. A Ni²⁺ ion and an NOG molecule are coordinated by surrounding residues (Fig. 3c).

To dissect the catalytic mechanism, we performed a structure-based mutagenesis study. To that end, we produced the JM13D236A and JM13D296A proteins, which have alanine replacements of the key residues involved in recognition of H3R26 and H3S28, respectively. These proteins showed significantly reduced demethylase activity (Fig. 3d). Mutation of Phe179 to serine, a small hydrophilic residue, produced a moderate decrease of activity, and replacement by glutamine, a hydrophilic residue with a larger side chain, significantly impaired activity (Fig. 3d), confirming the importance of the hydrophobic interaction between Phe179 and H3P30. Mutations of trimethyllysine or the nickel ion-binding residues completely abolished the activity of JM13, revealing their essential role in catalysis (Fig. 3d).

The H3K27me3 mark resides within the same ARKme3S consensus motif as the H3K9me3 mark, making the two marks chemically difficult to distinguish. In the JM13 structure, the specific stacking interaction between H3P30, at the n + 3 position, and Phe179 selects against the H3K9me3 substrate, which has an H3G12 at the n + 3 position, thereby ensuring substrate specificity.
Structural comparison with other histone demethylases. JMJ13 belongs to the KDM4 subfamily of histone demethylases based on phylogenetic sequence analysis of the catalytic domain. However, JMJ13 possesses a domain architecture that does not resemble the two known KDM4 subfamily H3K27me3 demethylases, REF6 and ELF6, which contain four tandem DNA-binding C2H2 zinc fingers at the end of the C-terminus (Supplementary Table 1). Instead, JMJ13 resembles the two substrate specific KDM5 subfamily histone demethylases, such as human KDM5/JARID1 and Arabidopsis JMJ14. The two proteins have similar domain arrangements and overall structures with an RMSD of 1.99 Å (Fig. 4a)24. However, the zinc coordination topologies differ. The first Zn$^{2+}$ ion occupies an analogous position in both JMJ13 and JMJ14. However, the second zinc ion occupies different sites in the two proteins and each is coordinated differently (Fig. 4f–h), yet the proteins maintain very similar
overall zinc finger domain topologies (Fig. 4f–h). Whether this relationship is indicative of similar function is presently not known.

Our structure-based sequence alignment identified the key residues involved in H3K4me3 recognition by JMJ14 and H3K27me3 recognition by JMJ13. These key residues are not conserved (Supplementary Fig. 4), although generally the JMJ13 and JMJ14 show similar overall structures (Fig. 4e). It suggests that the conserved jumonji-helical-zinc finger cassette may function as a general scaffold for histone demethylases, with specific features evolving within individual demethylases to enable differential regulation and substrate specificity.

**Ectopic JMJ13 expression causes pleiotropic defects.** H3K27me3 functions as a repressive histone mark. To investigate the function of JMJ13, which removes H3K27me3, we generated JMJ13-GFP (JMJ13ox) over-expressing transgenic plants. JMJ13ox plants had pleiotropic phenotypes, with dwarf seedlings, early flowering, and upward curling leaves.
Fig. 4  Comparison of JMJ13 and other histone demethylases.  a  A superposition of the JMJ13-NOG-H3K27me3 complex (in green) and UTX-NOG-H3K27me3 (in orange, PDB code: 3AVR) shows the similar overall structures and peptide-binding sites.  b The structures of the C4HCHC dual zinc finger of JMJ13 (left panel, in green) and the C4 single finger of UTx (right panel, in orange) showing different zinc coordination but similar folding topology.  c The superposition of the JMJ13-NOG-H3K27me3 complex (in green) and the UTX-NOG-H3K27me3 (in orange) shows that the two proteins employ correspondent residues Asp236 of JMJ13 and Asp1089 of UTX to recognize H3R26 in the two complexes, respectively.  In contrast, JMJ13 recognizes H3S52 by a side chain hydrogen bond with Asp296, but UTX lacks this recognition.  d The superposition of the JMJ13-NOG-H3K27me3 complex (in green) and UTX-NOG-H3K27me3 (in orange) shows that JMJ13 recognizes H3P30 by Phe179, whereas UTX recognizes H3P30 by Pro114A at different positions.  e A superposition of JMJ13-NOG-H3K27me3 complex (in green) and JMJ14-NOG-H3K4me3 complex (in magenta, PDB code: 5YKO) shows almost identical overall structures.  f A detailed view of the comparison of the zinc finger domains of JMJ13 (in green) and JMJ14 (in magenta). The two zinc finger domains possess similar overall structures with the first zinc ions occupying the same position but the second zinc ions located at different positions.  g The detailed structures of the zinc coordination of two zinc ions from JMJ13 (upper panel, in green) and the JMJ14 (lower panel, in magenta) show that the first zinc ions have the same coordination but the second ones have different coordination.  h The schematic representation of the sequence motif and coordination of the zinc ions of the JMJ13 C4HCHC domain (upper panel) and JMJ14 C5HC2 domain (lower panel). The spacing residues are denoted as X.

(Supplementary Fig. 5a), all of which increased in severity with increasing JMJ13 expression levels (Supplementary Fig. 5b). The JMJ13-GFP seedlings showed phenotypes similar to H3K27me3 silencing-deficient mutants hpe134 or REF6 over-expression plants, with reduced size of leaf surface cells (Supplementary Fig. 5c, d)25.

Some H3K27me3 target genes, including APETALA1 (AP1), APETALA3 (AP3), PISTILLATA (PI), AGAMOUS (AG), and SEPALLATA3 (SEP3), which are normally expressed in flowers, were ectopically activated in the JMJ13ox seedlings. (Supplementary Fig. 5e). Other genes, for example, FT and SOC1, which are expressed at low levels in 10-day-old seedlings, and the meristem function genes KNOTTED-LIKE FROM ARABIDOPSIS THALIANA 1 (KNAT1) and TOUCH 4 (TOC4), all of which are targets of REF6, were also upregulated in JMJ13ox seedlings (Supplementary Fig. 5e). Consistent with these changes in gene expression, H3K27me3 but not H3K4, H3K9, and H3K6 showed a strong global reduction in the two strong JMJ13ox transgenic lines (Supplementary Fig. 5f).

JMJ13 represses flowering.  We identified jmj13, a T-DNA insertion in the second intron of JMJ13 (GABI116B03, At5g46910) (Supplementary Fig. 6a)35. No JMJ13 transcript was detected in the jmj13 mutant (Supplementary Fig. 6b–c). The plant Polycarya group (PC-G) protein CURLY LEAF (CLF), which functions as an H3K27me3 methyltransferase, is required to repress targets such as AG and SHOOTMERISTEMLESS (STM). CLF, thereby controls flowering time, leaf morphology, and floral organogenesis36,37. The jmj13 clf double mutant phenotype was similar to but weaker than that of clf in terms of flowering time, curling of leaves (Supplementary Fig. 6d–g) and fertility, indicating that jmj13 partially suppresses the clf phenotype. Consistently, the expression levels of some H3K27me3 target genes that were ectopically activated in clf were reduced in the jmj13 clf double mutant (Supplementary Fig. 6h). However, the expression of SEP3 and AG is not reduced in jmj13 clf relative to clf, although the leaf curling and early flowering phenotypes are weakly suppressed in the jmj13 clf double mutant. This could be due to other H3K27me3 target genes, or due to functional redundancy with REF6, ELF6 or other possible H3K27me3 demethylases. Together, these genetic interactions demonstrate that JMJ13 functions as an H3K27me3 demethylase that partially antagonizes the H3K27 methyltransferase CLF in vivo, although further investigation would be required to reveal the details of crosstalk between CLF and JMJ13 and whether the phenotypic suppression is due to direct antagonistic function at particular loci.

We further measured flowering time and scored total leaf number at bolting, in different photoperiod conditions (LD and SD) and under different ambient temperature conditions (low, 16 °C; and high, 28 °C). The jmj13 mutants displayed early flowering in LD conditions38, regardless of low or high temperatures. However, jmj13 plants flowered early at 28 °C, but not at 16 or 22 °C, when grown in SD conditions (Supplementary Fig. 7).

To verify that loss of JMJ13 function is responsible for the early flowering phenotype of jmj13, we transformedjmj13 with a 7.0 kb genomic construct (gJM13) including 2.6 kb upstream and 0.5 kb downstream of the coding region. All the transgenic plants exhibited comparable flowering time to Col plants at 22 °C in LD conditions (Fig. 5a). Compared with the fully rescued pJM13: JMJ13-3A jmj13 lines, lines containing a construct with point mutations in the two conserved iron-binding amino acids, H293A and E295A (pJM13:JMJ13-3A-H293Ajmj13), showed similar flowering time to jmj13 mutants. This suggests that the enzyme activity is necessary for JMJ13 function, together with the jumonji and zinc finger domains (Fig. 5a, b).

In Col plants, LD conditions and elevated ambient temperatures promote flowering. However, when grown in SD conditions, increased temperature could, to some extent, overcome the unfavorable day-length condition to promote flowering. Consistent with these phenotypic observations, the reproductive development of jmj13 mutants was hypersensitive to, and promoted by, both LD conditions and elevated ambient temperatures (Fig. 5b). Collectively, these results suggest that JMJ13 negatively modulates flowering time under LD and in SD under higher temperature.

Crosstalk between JMJ13 and other flowering time regulators. To investigate the mechanism underlying JMJ13’s regulation of flowering in response to temperature, we crossed jmj13 with flm and syp. The flm jmj13 and syp jmj13 double mutants flowered earlier in all conditions, including 22 °C and SD, whereas jmj13 showed a wild-type flowering phenotype in 22 °C and SD (Supplementary Fig. 8a). Over-expression of FLM-β but not FLM-Δ repressed early flowering in jmj13 (Supplementary Fig. 8b–d). These genetic data are consistent with the possibility that FLM and SVP may act downstream or in parallel with JMJ13 in the regulation of flowering.

In the photoperiodic pathway, the circadian clock-regulated GIGANTEA (GI) protein positively regulates the oscillating expression of CO2,5. The transcript level and protein activity of GI are coordinately controlled by the light signaling pathway and the circadian clock. FT expression is activated by CO and repressed by FLM and SVP. Thus, transcriptional regulation of FT is a key output resulting from integration of photoperiodic cues with temperature signals. The flowering time of double mutants of ft jmj13, co jmj13, and gi jmj13 were assessed by counting rosette and cauline leaf numbers in bolting seedlings (Supplementary Fig. 9a–d). We found that all three double mutant lines displayed similar rosette leaf numbers to the ft, co, and gi single mutants (Supplementary Fig. 9a-d), and FT expression was upregulated in
Interestingly, we observed that high temperature and long-day conditions increased the expression of the gene jmj13 (Supplementary Fig. 10a, b and Supplementary data 1), indicating that many genes are regulated directly or indirectly by JMJ13.

Furthermore, we performed RNA-seq for Col and jmj13 grown under 22 °C LD conditions. The expression of SVP was decreased in jmj13 compared with wild type (Supplementary Fig. 10a, b), which is consistent with the genetic data. By contrast, the expression of CO, Late Elongated Hypocotyl (LHY) and Circadian Clock Associated 1 (CCA1) was upregulated in jmj13 (Supplementary Fig. 10a, b and Supplementary data 1), indicating that many genes are regulated directly or indirectly by additional mechanisms, such as circadian rhythm. Interestingly, we observed that high temperature and long-day photoperiod induced JMJ13 expression and increased JMJ13 protein accumulation in 7-day-old seedlings (Supplementary Figs. 10c, d and 11). These results suggested that JMJ13 may affect flowering time through temperature- and photoperiod-mediated pathways.

We further analyzed the H3K27me3 levels in Col and jmj13 by chromatin immunoprecipitation followed by sequencing (ChIP-Seq) in different day-length and temperature conditions. However, we did not observe the over-accumulation of H3K27me3 in jmj13 at FLM/SVP and CO/GI and no significant differences between Col and jmj13 were observed at these loci (Supplementary Fig. 12). This may due to the functional redundancy of ELF6, REF6 or other possible H3K27me3 demethylases. These observations are in consistent with recent report that global H3K27me3 levels were strongly elevated only in the triple elf6 ref6 jmj13 mutant in Arabidopsis

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**Fig. 5** JMJ13 negatively modulates flowering time in a temperature- and photoperiod-dependent manner. a Three complementary lines Comp−1,−2 and −3 (pMJ13::JM13-HAjmj13) showed similar flowering time with wild type in 22 °C LD conditions, but the complementary lines Comp-Mu−1,−2 and −3 with site mutation in the two conserved iron-binding amino acids (pMJ13::JM13mu-HAjmj13), the complementary lines with truncation in JmjC (Comp-ΔJmjC−1,−2,−3 (pMJ13::JM13-ΔJmjC-HAjmj13)) and zinc finger domain Comp-ΔZF−1,−2,−3 (pMJ13::JM13-ΔZF-HAjmj13) remain early flowering. The plant images were created by the authors in this research. b The flowering time of complementary lines (Comp−1,−2,−3; Comp-Mu−1,−2,−3; Comp-ΔJmjC−1,−2,−3; Comp-ΔZF−1,−2,−3) were assessed by counting rosette leaf numbers in bolting seedlings grown under 22 °C, 28 °C LD, and 22 °C, 28 °C SD conditions. Values are means ± SE of indicated number of plants for three independent biological repeats. N number was marked in the bottom of the column. Student’s t test was used to calculate the P value between Col and other lines. ****P value < 0.0001; ***P < 0.001; **P < 0.01; *P < 0.05; Bar = 1 cm. The individual data points are shown as dots. Source data are provided as a Source Data file.
Discussion
In most eukaryotic genomes, a large proportion of chromatin is enriched with H3K27me3. Erasure of these methyl groups is tightly controlled during development, and as part of acclimation to environmental conditions. Flowering is strictly regulated by various epigenetic factors. For example, we previously showed that the H3K4me3 demethylase JM14, and H3K27me3 demethylases REF6 and ELF6 directly regulate flowering integrators FT, FLC, and SOC1.[22,23–27] Recently, it was reported that JM13, REF6, and ELF6, shape the genome-wide distribution of H3K27me3 and control tissue-specific gene activation.[28] Global H3K27me3 levels were only increased in the triple elf6 ref6 jmj13 mutant, conforming that JM13, REF6, and ELF6 are functional redundant. REF6 plays a major role in determining the distribution of H3K27me3, with ELF6 and JM13 apparently supplementing REF6 function[28], which partially explained that there is no over-accumulation of H3K27me3 in jmj13. During flowering, REF6 and ELF6 have distinct functions, as ref6 and elf6 mutants have late and early flowering phenotypes, respectively.[21]. Here, we show that JM13 represses flowering in a temperature- and day-length-dependent manner in Arabidopsis. JM13’s repressive function prevents precocious flowering under unfavorable environmental conditions. Although JM13, ELF6, and REF6 are all H3K27me3 demethylases, they each have different functions in Arabidopsis flowering control, indicating that their differential targeting lead to diverse pathways of flowering regulation. Further work would be needed to determine how JM13 affects flowering time, which JM13 targets that might be phenotypically relevant, and what are the differences and similarities among ELF6, REF6, and JM13. Ambient temperature-dependent, flowering-time control is fine-tuned by multiple pathways in Arabidopsis. A distinct chromatin mark, H3K36me3 has been shown to affect temperature-induced alternative splicing, a major co-transcriptional/post-transcriptional regulatory mechanism to fine-tune ambient temperature-dependent flowering-time in Arabidopsis.[39,40]. Promotion of flowering at higher temperatures has been proposed to counter the unfavorable SD photoperiod.[41] Here we show that another chromatin mark H3K27me3 can also regulate temperature- and photoperiod-dependent flowering regulation. The H3K27me3 demethylase JM13 may negatively modulate temperature-photoperiod-dependent compensation by dampering temperature-driven flowering-promotion in the absence of LD induction. LD photoperiod, and especially high temperature, induces JM13 mRNA expression and JM13 protein accumulation. These results suggest that JM13 plays an important role in temperature- and photoperiod-dependent flowering time regulation.

Methods
Plant materials and growth conditions. Arabidopsis plants used in the study were in the Columbia background. Primers for genotyping jm13 are listed in Supplementary Table 3. Plants were grown in growth chambers under LD conditions (16 h light/8 h dark) or SD conditions (8 h light/16 h dark) at 16 °C, 22 °C and 28 °C, respectively. The rosette leaves number were counted when the seedlings booting. The mutants used in this study are flm-3, ap-32 (SALK_072930), f-1 (CS56), co-2 (CS55), gi-3 (CS51), elf6 (SALK_074694), ref6 (SALK_001018), and cf (SALK_139371). The lbp1 mutant used in this study was fji2-42.

Molecular cloning of JM13. To clone the full-length JM13 cDNA, we amplified a 2364-bp cDNA fragment from a reverse transcribed cDNA pool derived from Col-0 seedlings. The PCR product was cloned into pENTR/D-TOPO (Invitrogen). A Quick Change Site-Directed Mutagenesis Kit (Stratagene) was used to make the point mutations in JM13. The wild-type and mutant JM13 coding sequences were introduced into the pEarley-Gate104 vector. Primers used for cloning are listed in Supplementary Table 3.

Transient expression in Nicotiana benthamiana leaves. The constructs for JM13, or JM13[3293A/3295A] fused with GFP were transformed into Agrobacterium tumefaciens cells (strain EHA105). These cells were then injected into Nicotiana benthamiana leaves, which were harvested for nuclear isolation and immunostaining or immunoprecipitation after 48 h.[42]

In vivo histone demethylation assays. Nuclei transfected with JM13-GFP or JM13[3293A/3295A] GFP mutants were visualized by observing the GFP signal under a fluorescence microscope. Immunolabeling was performed by using histone methylation-specific antibodies (H3K27me3, Millipore 07-449; H3K27me2, Millipore 07-452; H3K27me1, Millipore 07-448; H3K4me3, Millipore 07-473; H3K4me2, Millipore 07-350; H3K4me1, Millipore 07-436; H3K36me3, Millipore 07-442; H3K36me2, Millipore 07-441; H3K36me1, Millipore 07-450; H3K36me3, Abcam ab9050; H3K36me2, Millipore 07-274; H3K36me1, Millipore 07-548). All these antibodies were diluted by 1:100. Alexa Fluor 555 or 488-conjugated goat anti-rabbit (1:300, Invitrogen) were used as secondary antibodies to determine the specific lysine modification site in vivo. After incubation with secondary antibody, nuclei were mounted by one drop of Vectashield Mounting Medium with 4',6-diamidino-2-phenylindole (DAPI) (Vector Laboratories), then photographed under a fluorescent microscope (Olympus BX51). Image (National Institutes of Health) was used for quantification of the immunolabeled nuclei.

Transcript level analysis. Total RNA was extracted using TRIzol Reagent (Invitrogen) for reverse transcription-PCR (RT-PCR) and real-time quantitative PCR (qPCR) analysis. Real-time qPCR analysis was performed using SYBR Green (CWBio, CW0760A). Primers used for transcriptional analysis are listed in Supplementary Table 3.

RNA sequencing analysis. Total RNA was isolated from newly fertilized silique of six week seedlings using a TRIZol kit (Invitrogen). Paired-end sequencing libraries with an average insert size of 400 bp were prepared with a TruSeq RNA Sample Preparation Kit v2 (Illumina) and sequenced on the HiSeq2500 (Illumina). Raw data obtained from Illumina sequencing were processed and filtered using the Illumina pipeline (http://www.illumina.com) to generate Fastq files. Finally, about 8 Gb high-quality 150-bp paired-end reads were generated from each library. FastQC (http://www.bioinformatics.babraham.ac.uk/projects/fastqc/) was initially run to assess the overall quality of all sample reads. Poor quality bases were filtered out using Sickle with parameters “–mode pe; –sanger; –q 20 –Q 50” (https://github.com/najoshi/sickle). The quality filtered reads were aligned to the Arabidopsis reference genome using TopHat version 2.0.9[43] with the parameters “-N 3 –read-edit-dist 3–segment-mismatches 1 -p 20 r =20 g =20-microexon-search =2 b =2-D =2-b –R 3 –no-software-search”, HTSeq software (http://www-huber.embl.de/users/ anders/HTSeq/doc/overview.html) was used to count the number of reads mapped to each of the genes. DESeq was used for differential expression analysis with the “Fisher’s Exact Test” method[44]. The genes showing an absolute value of log2 (fold change; jm13 mutant/WT) ≥ 0.6 and adjusted P value (false discovery rate; FDR) < 0.05 were considered as differentially expressed genes.

ChIP sequencing. A total of 2 g of 7-day-old seedlings sample was powdered in liquid nitrogen using a pestle and mortar and fixed with formaldehyde then immunoprecipitated with anti-H3K27me3 (Millipore 07-449), anti-H3 (Abcam; ab1791) antibodies. After decross-linking, proteinase K, and RNase treatment, the immunoprecipitated DNA was purified by phenol/chloroform extraction for the additional experiments including Illumina single-end sequencing[45]. The purified DNA was ligated with Illumina single-end genome sequencing adapters and then fragments fractioned, PCR amplified and sequenced according to standard protocols (single-end 36 cycles). ChIP-seq reads were aligned to Arabidopsis genome build TAIR10 by Bowtie 2 using default parameters with a local alignment model. Duplicated reads and low-mapping quality reads were identified and removed with SAMtools[4]. Enriched intervals were identified by MACS version 2.1.9 (with default parameters. Density maps of reads for visualization were based on reads of the 200-bp extension of sequencing reads in the 3′-direction after total reads normalization[46].

Western blot assay. Ten-day-old seedlings were ground in liquid nitrogen and the powder was boiled for 5 min in protein sample buffer. The proteins were resolved on a 15% SDS/PAGE gel and transferred onto nitrocellulose membranes (Bioline). After the membranes probed with anti-H3K27me3 (Millipore 07-449); anti-H3K27me2 (Millipore 07-452); anti-H3K27me1 (Millipore 07-448); anti-H3K4me3 (Millipore 07-473); anti-H3K4me2 (Millipore 07-3030); anti-H3K4me1 (Millipore 07-436); anti-H3K36me2 (Millipore 07-441); anti-H3K36me1 (Millipore 07-450); anti-H3K36me3 (Abcam ab1791) or anti-GFP (Roche 11814460001) in TBST (137 mM NaCl, 20 mM Tris·HCl pH 7.6, 0.1% Tween-20). After three washes with TBST, the signals were detected with Immobilon Western Chemiluminescent HRP Substrate (Thermo Fisher Scientific) for GFP antibody.

Protein expression and purification. The catalytic fragment of Arabidopsis JM13 (JM13CD, residues 90–578) was cloned into a pE3F Sumo vector to fuse an N-terminal hexahistidine plus yeast sumo tag. The plasmid was transformed into E. coli strain BL21(DE3) RIL and the transformants were cultured at 37 °C in LB...
medium. When the OD_{590} of cell culture reached 0.7, the protein expression was induced by adding IPTG to a final concentration of 0.2 mM and the cells were cooled to 20 °C. The recombinant expressed protein was purified using a HiTrap column (GE Healthcare). The hexahistidine plus yeast sumo tag was removed by ulp1 protease digestion followed by a second step HiTrap column (GE Healthcare). The target protein was further purified on Heparin and Superdex 200 columns (GE Healthcare). The untagged JMJ13CD easily precipitates in the in vitro activity assay. We further cloned JMJ13CD into a pMal vector (New England Biolabs) to fuse a MBP tag to the target protein. The MBP-tagged JMJ13CD was expressed in E. coli strain BL21(DE3) RIL with IPTG induction and purified using amylose resin (New England Biolabs), Heparin, and Superdex 200 columns (GE Healthcare). All the mutations were generated using a PCR based method and purified using the same protocol as wild-type protein. The chemicals and peptides were purchased from Sigma-Aldrich and GL Biochem Company, respectively.

**Crystalization and structure determination.** The purified JMJ13CD was concentrated to a final concentration of 10 mg ml⁻¹ and mixed with a-2G with a molar ratio of 1:4 at 4 °C for one hour. The crystal screening was carried out using sitting-drop vapor diffusion method at 4 °C. The JMJ13-a-2G complex was crystallized in 0.1 M MES, pH 6.5, 8% dioxane, and 1.6 M ammonium sulfate. To get crystals of the JMJ13–NOG–H3K27me3 complex, we first grew crystals of JMJ13–NOG in the same conditions as JMJ13-a-2G, and then soaked the crystals with 10 mM H3(24–35)K27me3 peptide for 60 h. All the diffraction data were collected at beamline BL19U1 of the National Center for Protein Sciences at Shanghai (NCPSX) at the Shanghai Synchrotron Radiation Facility (SSRF) and processed using the program HKL2000/3000 package.

A summary of the data collection statistics is listed in Supplementary Table 2. The structure of the JMJ13-a-2G complex was determined using the SAD method with the anomalous signal of the zinc peak as implemented in the program Phenix. The model building and structure refinement were conducted using the programs Coot and Phenix, respectively. Throughout the refinement, the geometry of the structure was monitored using the program Molprobity. The structure of the JMJ13–NOG–H3K27me3 complex was solved using the molecular replacement approach. The structure of JMJ13-a-2G complex as the search model was refined and was redefined using the same protocol as used for the JMJ13-a-2G complex. Density modification was applied with the program SOLOMON. A summary of the structure refinement statistics is listed in Supplementary Table 2.

**In vitro histone demethylation assay.** Since the untagged JMJ13CD easily precipitates during the in vitro assay, we used MBP-tagged JMJ13CD in the assay to increase the stability of the protein. The purified MBP-tagged JMJ13CD or its mutants (10 μM) were incubated with the H3(20–34)K27me3 peptide (80 μM) in a reaction buffer with 80 mM Fe(NH4)2(SO4)2, 2 mM ascorbic acid, 1 mM a-KG, 50 mM Tris-HCl pH 7.3, and 150 mM NaCl at 25 °C. The reaction solution was incubated at 25 °C for 2 h and then was quenched by heating to 95 °C for 3 mins. The reaction mixture was further desalted by ZipTip (Millipore). The eluted peptides were spotted on the MALDI plate and cocrystallized with 10 mg ml⁻¹ 4-cyano-4-hydroxyoxamic acid in 60% acetonitrile with 0.1% trifluoroacetic acid (TFA). Then, a 5000 MALDI-TOF/TOF mass spectrometer (ABSciex, Foster City, CA) was used for analyzing the samples. For quantification, peptide mass spectra were recorded in reflector mode from triplicate reactions with each spectrum acquired from 1250 laser shots. The resulting mass spectra were processed by the Data Explorer software.

**Data availability**

X-ray structures have been deposited in the RCSB Protein Data Bank with the accession codes: 6P0D for the JMJ13-a-2G complex and 6P4P for the JMJ13–NOG–H3K27me3 complex. RNA-Seq and ChIP-Seq data have been uploaded to NCBI SRA with accession numbers SRP164434 and SRP174856, respectively. The Source Data underlying Figs. 1, 3, 5, Supplementary Figs. 1, and 5–10 are provided as a Source Data file.

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Author Contributions

S.Z., H.H., H.R., Z.L., Q.Q., W.Q., X.L., X.Chen, X.Cui, S.L., and B.Z. performed the experiments and analyzed the data. S.Z., D.S., X.Cao, and J.D. conceived the study, designed the experiments, and wrote the article.

Additional information

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