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Paramutation-like features of multiple natural epialleles in tomato

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Abstract

Background: Freakish and rare or the tip of the iceberg? Both phrases have been used to refer to paramutation, an epigenetic drive that contravenes Mendel's first law of segregation. Although its underlying mechanisms are beginning to unravel, its understanding relies only on a few examples that may involve transgenes or artificially generated epialleles.

Results: By using DNA methylation of introgression lines as an indication of past paramutation, we reveal that the paramutation-like properties of the *H06* locus in hybrids of *Solanum lycopersicum* and a range of tomato relatives and cultivars depend on the timing of sRNA production and conform to an RNA-directed mechanism. In addition, by scanning the methylomes of tomato introgression lines for shared regions of differential methylation that are absent in the *S. lycopersicum* parent, we identify thousands of candidate regions for paramutation-like behaviour. The methylation patterns for a subset of these regions segregate with non Mendelian ratios, consistent with secondary paramutation-like interactions to variable extents depending on the locus.

Conclusion: Together these results demonstrate that paramutation-like epigenetic interactions are common for natural epialleles in tomato, but vary in timing and penetrance.

Keywords: Epigenetics, Paramutation, DNA methylation, Small RNAs, RNA-directed DNA methylation, Hybrids, Introgression lines

Background

Paramutation is an epigenetic process in plants (including pea, maize, tomato [1]) and animals (worm [2], fruit fly [3], mouse [4]) that is associated with gene silencing. It is unlike other epigenetic mechanisms, however, in that it involves transfer of the silent state from an allele with epigenetic modification to its active homologue. This paramutated allele then becomes silenced and it acquires the ability to silence other active alleles in subsequent generations so that inheritance patterns are non Mendelian [1, 5].

The best characterised examples of paramutation are from maize, at the *b1*, *r1*, and *pl1* loci. Genetic screens have implicated NRPD1 (*rmr6* [6]), the major subunit of Pol IV; the RNA-dependent RNA polymerase RDR2 (*mop1* [7]) and NRPD2a (*mop2* /*rmr7* mutants [8, 9]) the shared subunit of Pol IV and Pol V. These proteins are all required for RNA-directed DNA methylation (RdDM) in

which DNA methyltransferases are guided to the target sequence in the genome by base pairing of small RNAs (sRNAs). RdDM is also associated with paramutation of the *SULFUREA* locus in tomato [10, 11] and trans-chromosomal DNA methylation in *Arabidopsis thaliana* hybrids [12, 13]. Based on these findings the dominant model of paramutation implicates RdDM in the establishment and/or maintenance of the epigenetic mark.

Our interest in paramutation follows from an earlier study of sRNA in tomato lines in which homozygous regions were introgressed from *Solanum pennellii* into *Solanum lycopersicum* (cv. M82) [14]. The resulting introgression lines (ILs) each carry many loci at which sRNAs are more abundant than in either parent: they are transgressive [15]. To explain these findings we invoked epigenetic mechanisms because, in some instances, there was hypermethylation of the genomic DNA corresponding to the sRNA locus.

In this present study we focus initially on one locus, *H06*, at which there is transgressive sRNA and DNA hypermethylation in multiple introgression lines [15]. We were prompted to consider the involvement of a paramutation-like process at *H06* because presence of

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this epiallele in multiple ILs was not consistent with Mendelian inheritance. The ILs are produced by recurrent backcrossing of the F1 hybrid to the *Solanum lycopersicum* cv. M82 parent so that Mendelian features of the hybrid genome would co-segregate with specific regions of introgressed DNA. At *H06*, however, the segregation must have been independent of any introgressed regions. We could rule out that the anomalous behaviour of *H06* was due to a spontaneous epigenetic change because we could reproducibly recapitulate the transgressive sRNA and DNA hypermethylation at this locus in F2 progeny of the *S. lycopersicum* × *S. pennellii* cross [15].

An alternative explanation of the *H06* epiallele invoked non-Mendelian inheritance due to a hybrid-induced ‘paramutation’ that, once triggered, would be inherited in the recurrent backcross progeny independently of any one region in the *S. pennellii* genome. The genetic and molecular tests presented here are fully consistent with that hypothesis and they indicate further that the timing of *H06* paramutation correlates with the production of sRNAs from the paramutagenic epiallele. We also identify other methylated DNA epialleles in the ILs with paramutation-like properties: they are absent from the parental lines, present in multiple ILs independently of a specific introgressed region from *S. pennellii* and they transfer their epigenetic mark to a non-methylated allele following backcrossing to M82. Based on the characterisation of these loci we propose that multiple paramutation-like events occur in the progeny of a *S. lycopersicum* cv. M82 × *S. pennellii* cross. These events illustrate how epigenetic effects, of which paramutation is an extreme example, may be induced by hybridisation of divergent genomes. Further characterisation of this epigenetic spectrum will reveal the defining features of paramutation loci and the extent to which hybrid-induced changes to the epigenome can influence transgressive segregation in crop plants and natural evolution.

Results

Paramutation at the *H06* locus in multiple lines

The *H06* locus [15] is in the euchromatin of chromosome 8, nine kilobases upstream of two genes in divergent orientation (Fig. 1a). It is unmethylated and lacks small RNAs in M82 but, in IL8–3, it is methylated in all three contexts (CG, CHG and CHH, where H is any nucleotide but G) and produces abundant 24-nt sRNAs (Fig. 1a–c). We refer to this as the *H06*^{IL} epiallele. The homologous locus in *Solanum lycopersicum* cv. Micro-Tom and *Solanum pimpinellifolium* has an epigenetic profile similar to *H06*^{IL} both in terms of DNA methylation and sRNA production (Fig. 1b and c). In *Solanum pennellii*, however, this locus was distinct from the other species in that there was full methylation at CG and CHG but no sRNAs and only partial CHH methylation

in the corresponding region (Fig. 1b and c). This finding is different from our previous report in which the *S. pennellii* locus was described as hypomethylated at all three C contexts [15] (in the aerial part of two-week-old seedlings, whereas here we use only the leaf of a two-week-old seedling). The present data are, however, consistent with the previous finding in that the RdDM features of *H06* - mCHH and sRNA - are low or absent in *S. pennellii* seedlings.

To find out whether the *H06*^{IL} has properties consistent with paramutation, we crossed M82 × IL8–3 and monitored the DNA methylation in F1 (M82 × IL8–3), F2 and BC1 (M82 × F1) generations. The DNA was extracted from the leaves of 15-day-old plants and it was assayed with McrBC digestion. The first aim of these tests was to establish whether the epigenetic mark could be heritably transferred from *H06*^{IL} to *H06*^{M82} to create an *H06*^{IL'} epiallele. The second aim was to find out whether *H06*^{IL'} could mediate secondary paramutation and transfer its epigenetic mark to *H06*^{M82}.

Fourteen of the fifteen F1 plants had highly methylated *H06* DNA in this assay (Fig. 2a) indicating that *H06*^{M82} had been converted to *H06*^{IL'}. The level of methylation in the 30 F2 plants derived from highly methylated F1s was always high, showing that *H06*^{IL'} and the newly established *H06*^{IL'} are normally stable, thus fulfilling the first criterion of paramutation that the epigenetic mark could be transferred from *H06*^{IL} to *H06*^{M82} to form a stable *H06*^{IL'} epiallele. There was, however, instability of *H06*^{IL} in a single F1 plant and its F2 progeny in which the *H06* DNA was completely unmethylated (Fig. 2a).

The secondary paramutation criterion was tested in the M82 × F1 backcrossed (BC) progeny. If *H06*^{IL} but not *H06*^{IL'} is paramutagenic then half of the BC1 plants would have fully methylated *H06* DNA and half would have at most 50% methylated DNA. In contrast, if both epialleles were paramutagenic, there would be more than half of the plants with full methylation of *H06* DNA. The data (Fig. 2a) are consistent with secondary paramutation because there was an excess of highly methylated plants (18 highly methylated plants, 8 half-methylated or lower, *p*-value = 0.038, one-sided binomial test). It is likely, however, that some of the *H06*^{IL} or *H06*^{IL'} alleles in the BC1 were unstable and reverted to *H06*^{M82} because five of these BC plants had less than 50% methylated DNA at *H06*.

To investigate the paramutagenic properties of the *H06* allele in Micro-Tom, *S. pimpinellifolium* and *S. pennellii* we made a further series of crosses with M82 and analysed the F1 progeny. With Micro-Tom and *S. pimpinellifolium* the results were as with IL8–3: all of the *H06* alleles were highly methylated in the F1, irrespective of the direction of the cross (Fig. 2b). In contrast, in M82 × *S. pennellii*, about half of the *H06* alleles were

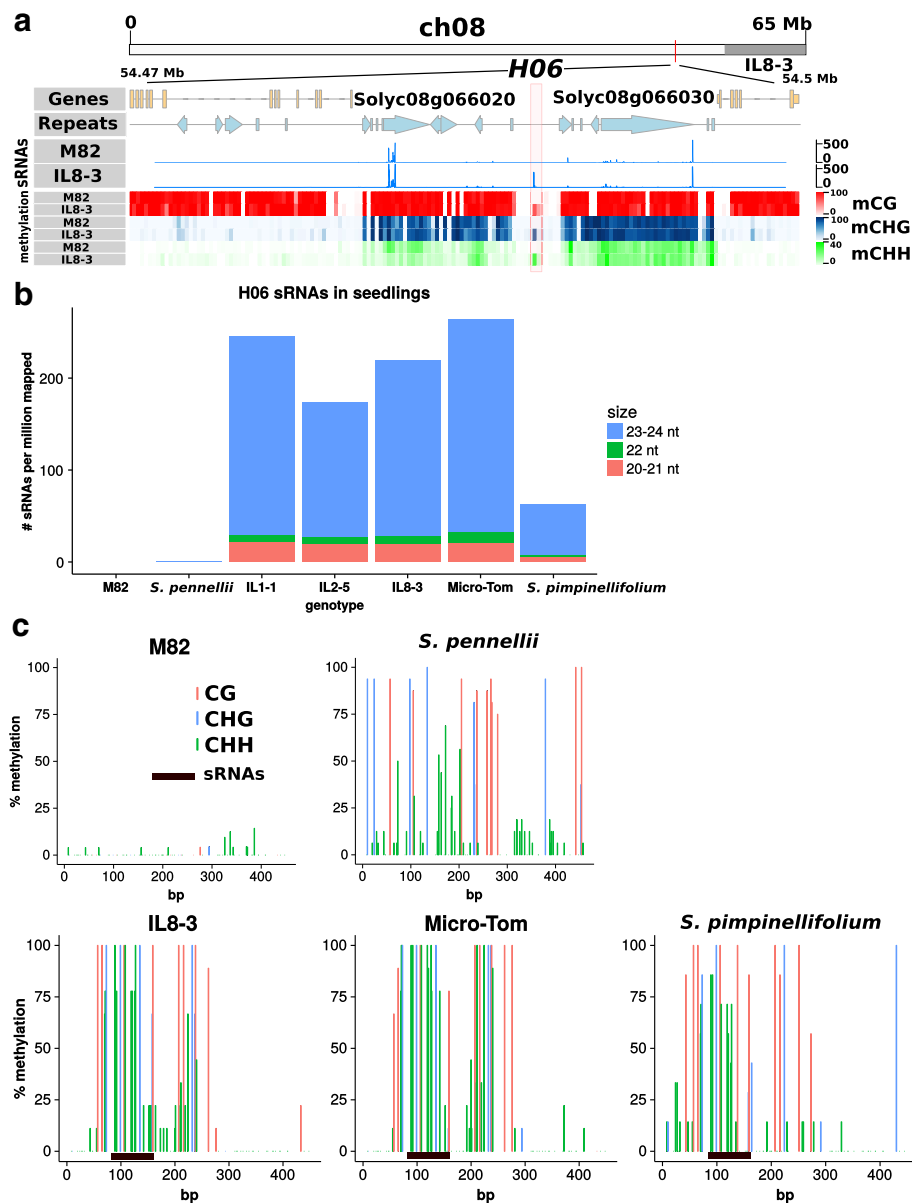


Fig. 1 *H06* epialleles. **a** Genomic location of *H06*, sRNAs in M82 and IL8-3 whole seedlings and methylation in seedling leaves. The *H06*^{IL} epiallele in IL8-3 was methylated in the CG, CHG and CHH contexts and the source of abundant sRNAs. Neither DNA methylation nor sRNAs were detected in the *H06*^{M82} epiallele. **b** High sRNA production at *H06* in seedlings of three introgression lines, the Micro-Tom cultivar and the wild relative *S. pimpinellifolium*. **c** *H06* methylation in leaves of M82, *S. pennellii*, IL8-3, Micro-Tom and *S. pimpinellifolium* determined by Sanger bisulfite sequencing (at least 10 independent clones per genotype). Primer sequences are given in Additional file 10, and *H06* sequences in Additional file 11

methylated in the 15-day leaves (Fig. 2c). This level increased in flowers and, in pollen and 15 d leaves of F2 plants, it was close to 100% (Fig. 2c). From these data we conclude that our various *Solanum* genotypes carried three distinct epialleles at *H06*: *H06*^{M82}, the *S. pennellii* allele referred to as *H06*^{pen} and *H06*^{IL}. The *H06*^{M82} allele had neither methylation nor sRNAs. *H06*^{pen} had DNA methylation but without the sRNA characteristics of RdDM in leaves of seedlings. This allele triggered an

epigenetic change to *H06*^{IL} but only after reproductive development in the F1. The third epiallele - *H06*^{IL} - was present in Micro-Tom and *S. pimpinellifolium* in addition to IL8-3, had both abundant sRNAs and DNA methylation in leaves of seedlings, and it could trigger a paramutation-like change to *H06*^{M82} without any evident lag. The *H06*^{IL} alleles had the general characteristics of *H06*^{IL} but with incomplete penetrance of its effect on *H06*^{M82}.

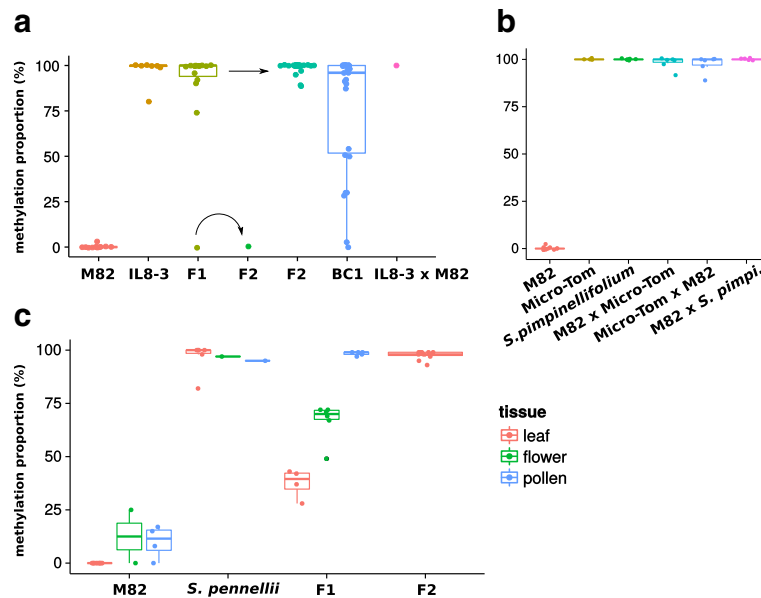


Fig. 2 *H06* paramutation assessed by McrBC. McrBC digests methylated DNA, so that comparing amplification of McrBC-treated and non-treated DNA by qPCR reveals the proportion of molecules that were methylated. **a** *H06* McrBC in M82 × IL8-3 seedling leaves. Unmethylated *H06*^{M82} ($n = 12$), crossed with the methylated *H06*^{IL} from IL8-3 ($n = 7$), became methylated in most F1s (14 out of $n = 15$), and remained methylated in the F2 ($n = 30$). Back-crossing the F1 with M82 demonstrated weak paramutagenicity (BC1, $n = 26$, p -value = 0.03776, one-sided binomial test). Paramutation also occurred with the IL as the female parent (IL8-3 × M82, $n = 1$). **b** *H06* McrBC in M82 × Micro-Tom and *S. pimpinellifolium*. F1 seedling leaves are fully methylated ($n = 6$ for each cross), like Micro-Tom and *S. pimpinellifolium* themselves ($n = 4$ and 5 respectively). **c** *H06* McrBC in M82 × *S. pennellii*. The F1 vegetative tissue was half-methylated ($n = 4$) whereas the F1 pollen and the F2 leaves were fully methylated ($n = 5$ and 11 respectively)

H06 paramutation timing correlates with sRNA production

To further investigate the interaction of *H06*^{M82} with *H06*^{pen}, we used allele-specific Sanger bisulfite sequencing of *H06* DNA from leaves and pollen of M82, *S. pennellii* and their F1. Consistent with the McrBC results, the allelic methylation of the F1 in the leaves mirrored that of the parents: low methylation of the M82 allele and high methylation of the *S. pennellii* allele (Fig. 3a) at CG and CHG. In pollen, however, the M82 allele of the F1 became hypermethylated in CG, CHG and CHH contexts (Fig. 3b), whereas pollen in the M82 parent is hypomethylated.

This gain of methylation by the M82 allele in the F1 correlated with an increase in sRNA production at *H06* (Fig. 3c). The *H06* sRNA levels were highest in flowers of *S. pennellii* and the F1, and very low in M82 leaves, flowers and pollen. These levels were, however, much lower than in IL8-3: in the flowers of *S. pennellii* there were 1–2 reads per million mapped whereas in IL8-3 seedlings there were more than 150.

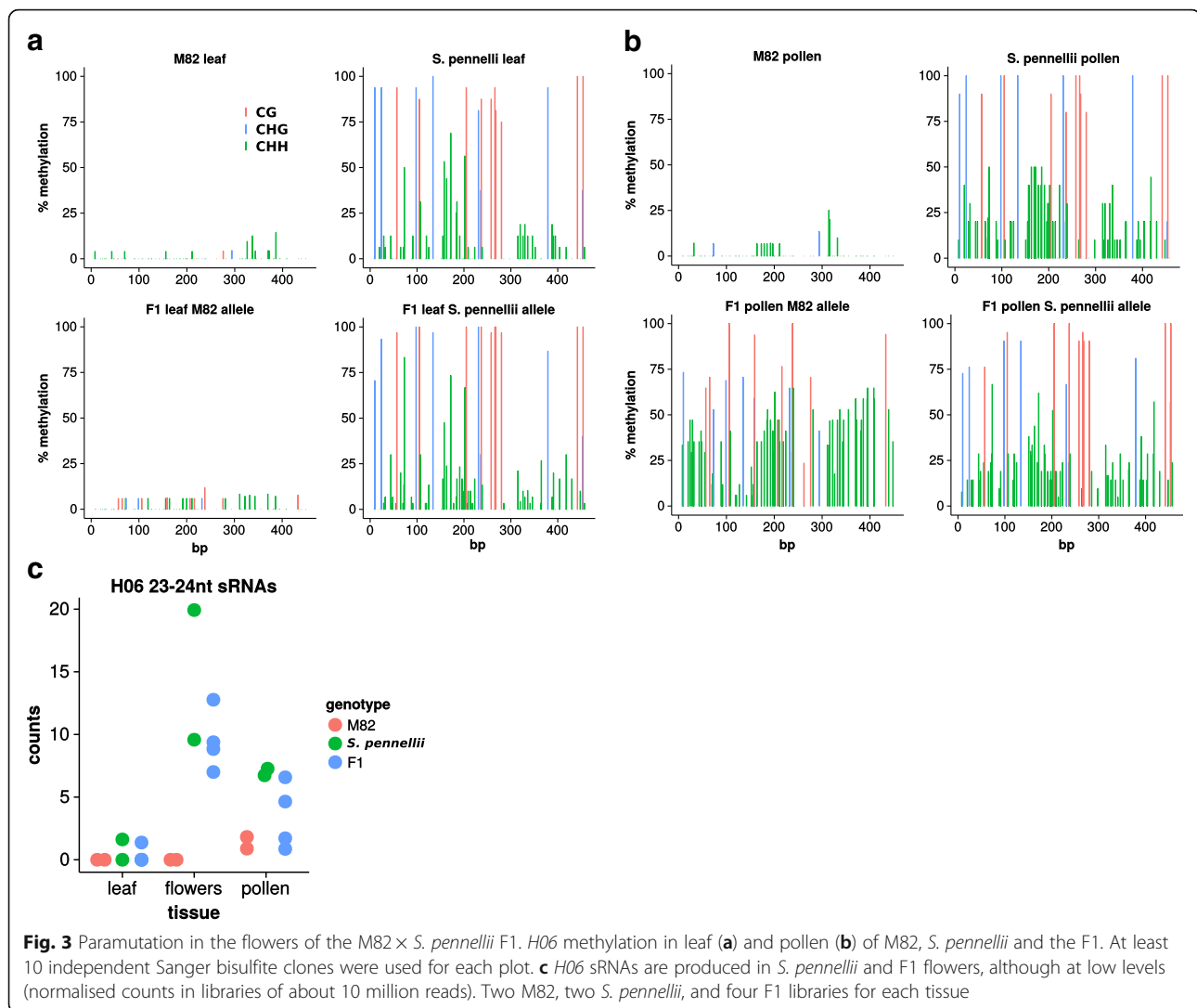
From these data there is a clear correlation of sRNA with the paramutation-like properties of the various *H06* loci. *H06*^{M82} lacks both sRNA and DNA methylation and is paramutable; *H06*^{pen} with low levels of sRNAs triggers a delayed paramutation-like process and *H06*^{IL}, at which the sRNA levels are high, mediates a rapid transfer of the epigenetic marks to *H06*^{M82}. This correlation of paramutation

and sRNA implicates RdDM in the process. Furthermore the changes in both mCHH and sRNA in pollen and flowers suggest that the reproductive phase may be a key developmental stage in the transfer of an epigenetic mark between alleles of *H06*.

De novo methylated *H06* recapitulates paramutation

In principle the association of RdDM with the paramutation-like properties of *H06* could be either a cause or consequence of the transition from *H06*^{M82} to *H06*^{IL}. To test the possibility of a causal role of RNA we used virus-induced gene silencing (VIGS) in which an RNA virus is modified to carry a small host genomic sequence insert. Such RNA viruses may direct DNA methylation of the corresponding genomic DNA of the infected plant [16] and we have previously used this approach to recapitulate the silencing of the *sulf* locus in tomato [11].

To test this system at *H06*, we infected unmethylated M82 plants with a Tobacco Rattle Virus (TRV) containing a 500-bp *H06* sequence (TRV-*H06*, Fig. 4a). Out of 11 successfully infected plants (V_0 generation) spread across three replicate experiments, only one plant gave rise to methylated progeny (V_1 generation) (Fig. 4b), with 30% of the plants having high methylation (20/59). None of 6 the control plants infected with unmodified TRV produced methylated progeny (56 tested V_1 plants) and,



with many tens of M82 plants tested over a period of 3 years, we have never observed spontaneous methylation of *H06* DNA. It is likely therefore that the new epiallele (*H06*^{VIGS}) was triggered by the TRV-*H06* VIGS.

This new epiallele (*H06*^{VIGS}, Fig. 4c) was distinct from *H06*^{pen} and *H06*^{IL} (Fig. 1c) in that the hypermethylation was in a restricted region of 200 bp rather than 500 bp. The *H06*^{VIGS} epialleles were stable in the V₂ generation (Fig. 4b). We tested whether this new epiallele was paramutagenic by backcrossing V₁ plants to M82: if *H06*^{VIGS} is inherited as a standard Mendelian locus without paramutation, the BC₁ plants would have no more than 50% of methylated *H06* DNA in an McrBC assay; if however *H06*^{VIGS} is paramutagenic, some backcrossed plants would exhibit methylation above 50%. Of 21 BC₁ plants, 4 had substantially more than 50% of methylated *H06* DNA (Fig. 4b), and we conclude

that *H06*^{VIGS} has weak paramutagenic activity. There were also many plants with less than 50% of methylated *H06* in these BC₁ plants that is due, presumably, to instability of *H06*^{VIGS} in the heterozygous condition. The ability to epigenetically modify and confer paramutation properties to *H06* by VIGS indicates that sRNAs could be causal in paramutation. However the low efficiency of VIGS in methylating *H06* precludes definitive conclusions about the mechanism of paramutation.

Genome-wide 'paramutation' in introgression lines?

Is *H06* paramutation an isolated example, or could there be other similar loci in *Solanum* genomes? To address this question we screened the DNA methylome of M82 and three different introgression lines, IL1–1, IL2–5 and IL8–3, in duplicates (summarised in Additional file 1). We reasoned that paramutation loci would be differentially methylated compared to M82 in multiple

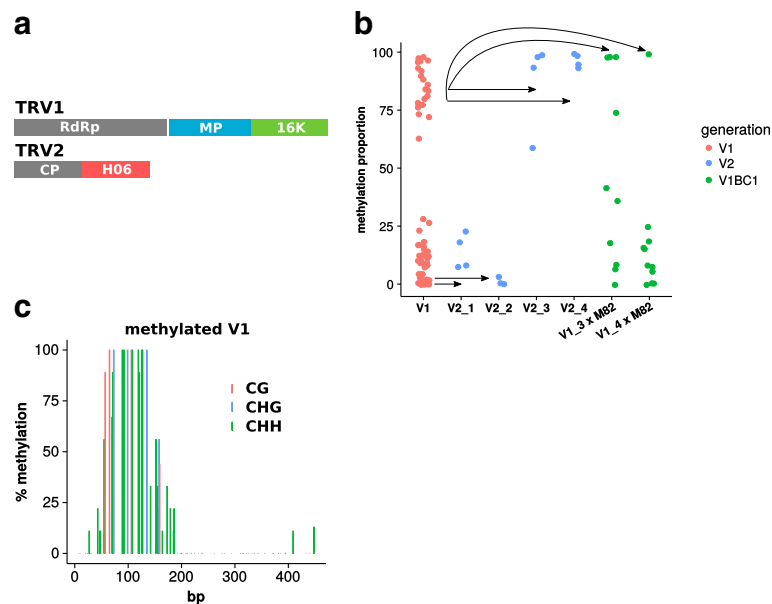


Fig. 4 Virus-induced paramutation. **a** Tobacco Rattle Virus modified to contain a 500-bp *H06* fragment. **b** DNA methylation in the progeny of the V_0 infected plant that gave methylated offspring, assessed by McrBC ($n = 59$ V_1 plants). V_1 methylation patterns were stable in the V_2 generation ($n = 4, 3, 4$ and 4 for V_{2_1} to V_{2_4}), and exhibited weak paramutagenicity in backcrosses to M82 (V_1BC_1 , $n = 10$ for both $V_{1_3} \times M82$ and $V_{1_4} \times M82$). **c** *H06* methylation of a V_1 plant assessed by Sanger bisulfite sequencing (9 independent clones)

introgression lines. To identify DMRs with this characteristic we compared methylation counts in CG, CHG and CHH of each introgression lines to M82 in 300-bp sliding windows. Each introgression line had 10,000–30,000 DMRs depending on context (CG, CHG and CHH) as compared to M82 (Fig. 5). Only a subset of these DMRs were shared in all three ILs: around 4600 CG hypermethylated DMRs, 1800 CG hypomethylated DMRs, 3600 CHG hypermethylated DMRs, 1000 CHG hypomethylated DMRs, 262 CHH hypermethylated DMRs and 900 CHH hypomethylated DMRs (Fig. 5 and Additional file 2). Of these DMRs there were 25, including *H06*, that were hypermethylated in all three contexts (Additional file 3). There were also 10 hypomethylated DMRs in all three contexts (Additional file 4).

The RdDM model of paramutation suggests that there would be 24-nt sRNAs at the paramutated loci, as is the case for *H06*. Using seedling sRNA libraries from [15], we identified 134 loci upregulated for these sRNAs in the ILs, and 68 down-regulated (Fig. 6a and b). Of the 134 upregulated sRNA loci, 9 overlapped hypermethylated DMRs and 3 overlapped hypomethylated DMRs. In addition to *H06*, one other locus (ch12:62124801–62,125,100, referred to as Hyper1) had increased sRNAs and hypermethylation in all three contexts in the introgression lines (Additional file 5: Figure S1). This locus and the other hypermethylated DMRs that overlap differential sRNAs are

prime candidates for paramutation that would follow the same pattern as *H06*.

DMRs can be associated with differences in gene expression, so we performed differential expression analysis between M82 and the three ILs under scrutiny with data from [17]. In this data set for seedlings grown in sun and shade, we found 124 genes that were upregulated in ILs compared to M82, and 108 that were downregulated (Fig. 6c). The reported log₂ fold changes of the differentially expressed genes were modest, in part because a large proportion of differentially expressed genes had low average expression (and hence low counts, leading to a shrinkage of the log₂ fold change estimates, Fig. 6c and Additional file 6: Figure S2). There was no significant enrichment in biological process, molecular function or cellular compartments of gene ontology terms associated with the differentially expressed genes (upregulated, downregulated or the full list). 10 and 5 of the upregulated genes were overlapped (over their 2 kb promoter or transcribed sequence) by shared hypermethylated and hypomethylated DMRs, respectively, while 4 of the down-regulated genes overlapped hypermethylated DMRs and 7 hypomethylated ones. These genes may be of particular interest to investigate transcriptional and physiological effects of paramutation following a cross between M82 and *S. pennellii*. Although this data set encompasses two environmental conditions (sun and shade), DMRs may affect the transcription of more genes in a tissue- or development-specific manner.

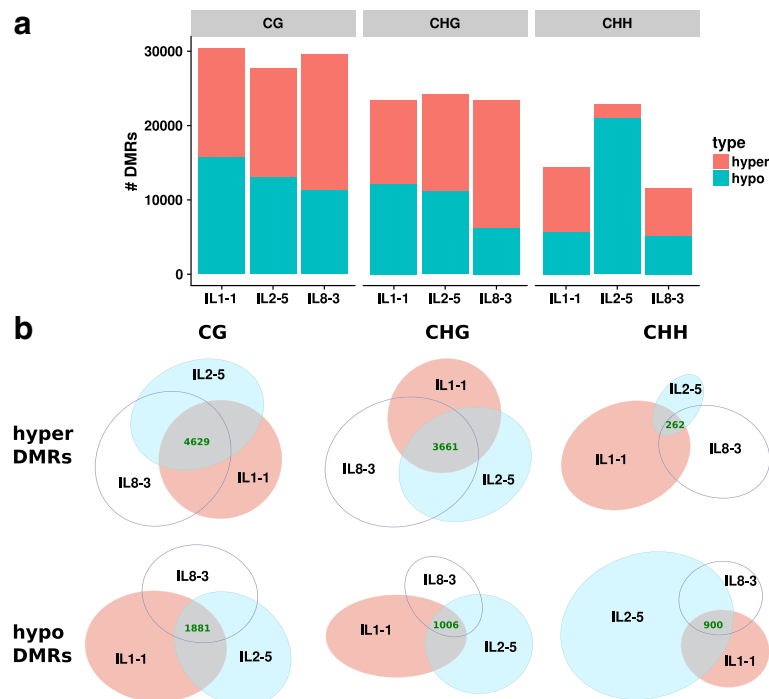


Fig. 5 Introgression lines DMRs. **a** Number of DMRs in each introgression line compared to M82 for CG, CHG and CHH contexts. Two plants per genotype. **b** DMR overlap across introgression lines. Hypermethylated DMRs have higher methylation in the introgression lines compared to M82. Area-proportional Venn diagrams

Paramutation-like activity of DMRs

Hypermethylated DMRs

To evaluate the paramutagenic properties of the hypermethylated DMRs, we selected 9 regions based on hypermethylation in multiple cytosine contexts in the three ILs, amenability to McrBC assays, presence of differential sRNAs, and proximity to differentially expressed genes (Hyper1–9, Table 1, Additional file 5: Figure S1). We tested their inheritance in the progeny of M82 × IL8–3 by McrBC-qPCR. Mendelian segregation would give 50% methylation in the F1, and in the F2 either 100% (1/4

progeny), 50% (1/2 progeny) or 0% (1/4 progeny) methylation. We tested these DMRs in 10 different F1s and in 33 F2s from three different F1 plants. Any evidence for more than 50% methylation of the F1 alleles or more than 3/4 F2 progeny with 50% or higher methylation would indicate non-Mendelian inheritance and evidence for a paramutation-like mechanism.

According to these criteria, there was evidence of a paramutation-like process at four loci (Hyper1–4, Fig. 7a and Table 1). For three of those (Hyper1–3), some F1s had more than 50% of methylated alleles. At the fourth

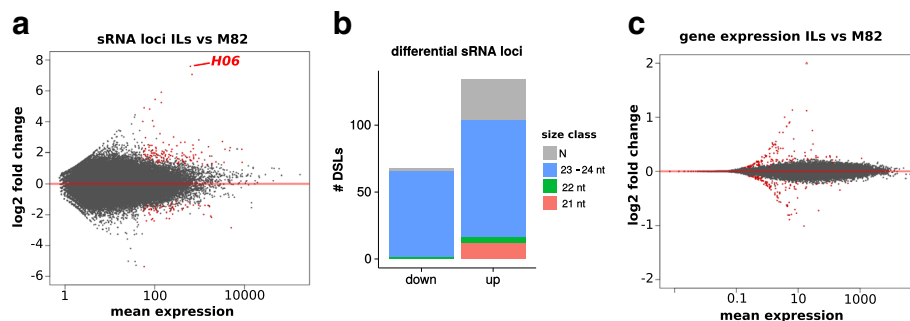


Fig. 6 Differential sRNA and gene expression in the introgression lines. **a** Differential analysis of sRNAs between M82 (two plants) and IL1–1/IL2–5/IL8–3 seedlings (one plant per genotype). Data from [15]. **b** Classification of differential sRNA loci by ShortStack. **c** Differential gene expression between M82 ($n = 40$) and IL1–1/IL2–5/IL8–3 seedlings ($n = 30$). Data from [17]. sRNA loci and gene that are differentially expressed ($p\text{-adj} < 0.05$) are coloured in red

Table 1 Summary of paramutation candidates and their validation

DMR	chr	Start	End	Type	Differential sRNAs	DE nearest gene	Evidence for heritable methylation change
Hyper1	ch12	62,124,801	62,125,100	hyperCG/CHG/CHH	yes	no	strong, incomplete penetrance
Hyper2	ch02	48,226,401	48,226,800	hyperCG/CHG/CHH	yes	no	strong, incomplete penetrance
Hyper3	ch09	61,960,201	61,961,700	hyperCG/CHG	no	yes	weaker
Hyper4	ch04	26,469,001	26,469,500	hyperCG/CHG	no	yes	weaker
Hyper5	ch09	61,962,201	61,962,500	hyperCG/CHG/CHH	no	yes	no
Hyper6	ch08	59,748,601	59,748,900	hyperCG	yes	no	no
Hyper7	ch04	58,749,201	58,749,500	hyperCG/CHG/CHH	yes	no	inconclusive
Hyper8	ch12	39,437,001	39,437,700	hyperCG/CHG	no	yes	inconclusive
Hyper9	ch04	26,561,001	26,561,500	hyperCG/CHG	no	yes	no, high variability
Hypo1	ch07	12,086,201	12,088,700	hypoCG/CHG	no	no	strong, incomplete penetrance
Hypo2	ch06	31,383,601	31,384,100	hypoCG/CHG	no	no	weaker
Hypo3	ch06	47,125,401	47,125,700	hypoCG/CHH	yes	yes	weaker
Hypo4	ch03	1,025,801	1,026,100	hypoCG/CHG	no	no	no
Hypo5	ch09	67,640,601	67,641,100	hypoCG/CHG	no	no	no
Hypo6	ch05	22,508,201	22,508,700	hypoCG/CHG	no	no	no, high variability
Hypo7	ch12	34,481,401	34,482,700	hypoCG/CHG/CHH	no	no	no, high variability
Hypo8	ch03	1,023,801	1,024,100	hypoCG/CHG	no	no	no, high variability

locus (Hyper4, Fig. 7a) the F1 progeny had approximately 50% of methylated alleles consistent with Mendelian inheritance but more than 3/4 of the F2 progeny had more than 50% methylated alleles (only 3/33 F2s had low methylation, p -value = 0.021, one-sided binomial test). This pattern is compatible with a partial gain of methylation by the M82 allele later in the development of the F1 (as was the case for *H06* in M82 × *S. pennellii*). Of note, Hyper1 corresponds to a region of highly transgressive sRNAs (log₂ fold change of 7) that are 24-nt in length (Additional file 5: Figure S1). It is therefore very similar in behaviour to *H06*. As for Hyper3, it is located in the immediate promoter of the *Solyc09g064640* gene, which is expressed at lower levels in the introgression lines compared to M82 (Additional file 6: Figure S2).

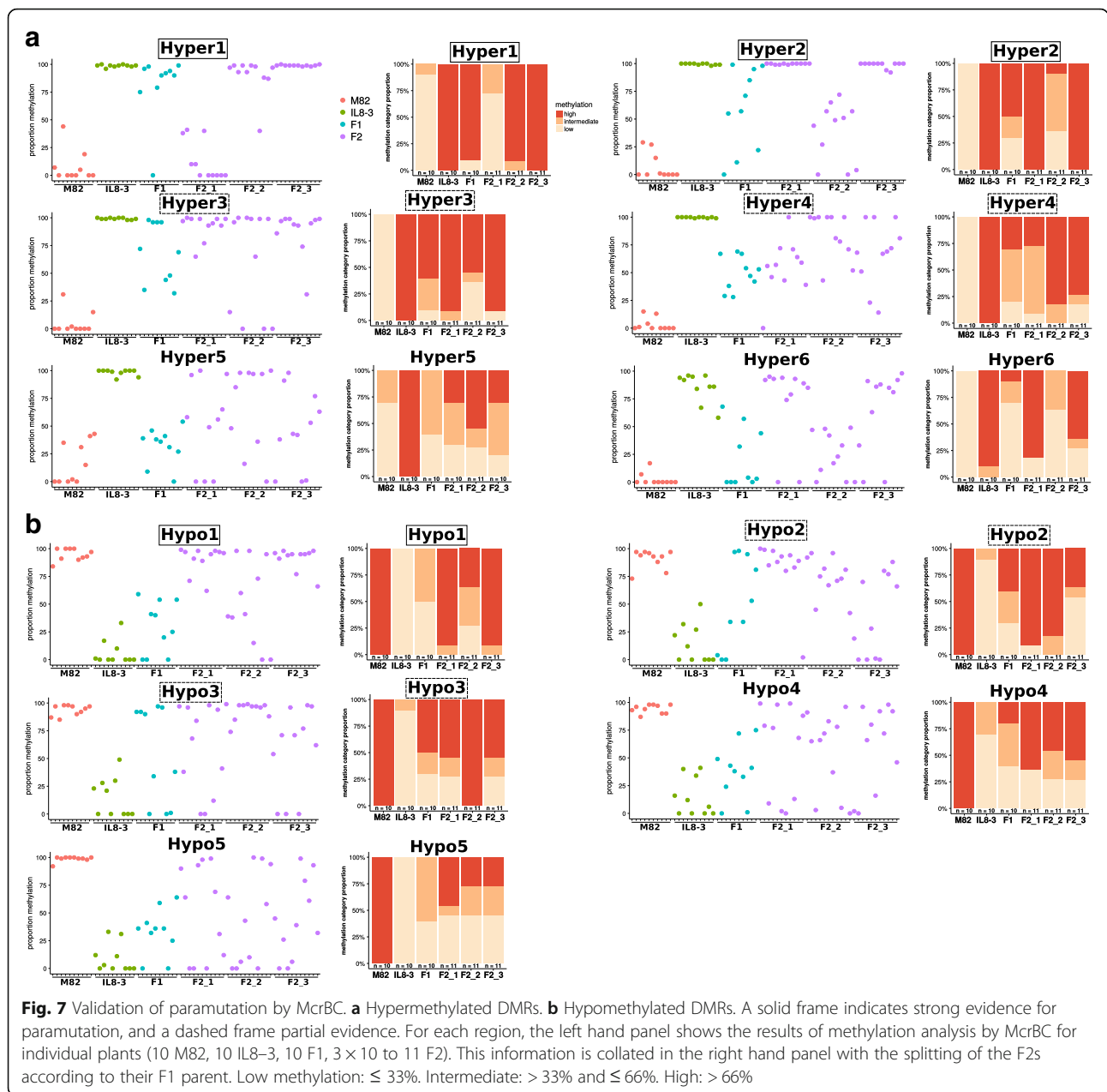
It is clear, however, that the DNA methylation marks at these loci are not completely stable. At Hyper1 and Hyper2, for example, there were F1 s with very little methylation, suggesting that there must have been loss of methylation even from the IL8–3 allele. It is likely therefore that there are two oppositely acting mechanisms influencing the methylation status of these loci: a paramutation-like event leading to de novo establishment of DNA methylation and a second process leading to removal of DNA methylation. The net effect of these processes is likely to account for the extent to which the pattern of DNA methylation deviates from Mendelian ratios in these F1 and F2 plants. However once established in the F1, methylation may be fairly stable: F2 progeny from two out of three F1s were consistently highly methylated for Hyper1 and Hyper2.

In addition to these four loci (Hyper1–4, Fig. 7a) we also investigated inheritance at five other loci that were hypermethylated in the IL1–1, IL2–5 and IL8–3 (Additional file 7: Figure S3 and Table 1). At Hyper5 and Hyper6 there was no conclusive evidence for paramutation-like behaviour (Fig. 7a) and we conclude either that these are conventional Mendelian loci or that the instability of the methylation mark in the progeny of M82 × IL8–3 offset the effects of the paramutation-like mechanism. At Hyper7 and Hyper8, high methylation in some M82 plants as assessed by McrBC does not allow us to conclude that the lack of F2s with low methylation reflects non-Mendelian segregation (Additional file 7: Figure S3). Methylation in IL8–3 at Hyper9 was variable, and there was no evidence for paramutation-like interactions (Additional file 7: Figure S3).

Hypomethylated DMRs

We also tested inheritance of 8 hypomethylated DMRs in IL1–1, IL2–5 and IL8–3 (Hypo1–8, Fig. 7b and Additional file 7: Figure S3). For one locus (Hypo1), there was a high proportion of the hypermethylated allele in the F2 (from two F1s in particular) although in the F1 the trend was towards medium-to-low methylation. These proportions indicate that methylation of this locus is subject to instability but that the hypermethylated M82 epiallele has some paramutation-like activity.

Similarly at Hypo2 and Hypo3, there was evidence for contrasting dynamics of methylation: 4 and 5 (out of 10) F1s were highly methylated, while 3 in each case had very low methylation (Fig. 7b). Overall the proportions of lowly



methylated F2s did not significantly depart from Mendelian ratios, but the F1 of origin had clear effects, with progenies of some F1s homogeneously hypermethylated. These results argue for paramutation-like activity as well as instability of the hypermethylated M82 epialleles.

At Hypo4 and Hypo5 there was no evidence for non-Mendelian segregation (Fig. 7b), while for the remaining three loci (Hyper6-8, Additional file 7: Figure S3), McrBC estimates of methylation in the parents were too variable to interpret the methylation levels of their progeny.

In summary, of the 17 DMRs tested in detail, 7 showed at least partial evidence of a heritable gain of methylation upon crossing (Table 1). Because the methylation

differences were observed in three independent introgression lines, it is unlikely that they are due to a trans effect from the introgressed *S. pennellii* region and a more likely explanation is that there is an epigenetic effect that may be partially related to paramutation-like mechanisms when methylation is gained in the M82 × IL8-3 F1s and F2s.

Discussion

Origins of DNA methylation differences: Paramutation-like mechanisms versus spontaneous epigenetic variation

The main aim of this study was to explore the epigenetic changes associated with wide cross hybridisation of

M82 × *S. pennellii* through the characterisation of *H06* and other DMRs in at least three different ILs relative to the parental lines. In principle these shared DMRs, as with DMRs in hybrid progeny of maize [18], rice [19] and *Arabidopsis thaliana* [12, 20], could be due to spontaneous epimutation [21, 22] or genetic differences in seedstocks (e.g. a transposon insertion) [23, 24] as well as to genetic or epigenetic interactions between the parental genomes. For many of the DMRs shared by the three ILs under study, we cannot say which of these explanations is likely to apply. For *H06*, however, we could reproduce its epimutation whenever we made the M82 × *S. pennellii* cross and paramutation in backcrosses of IL8–3 with M82 occurred at a high frequency, suggesting that the epigenetic state of the ILs at *H06* is due to the hybridisation process. To explain hybrid-induced paramutation it could be that there are genetic or epigenetic differences between the parental genomes. A genetic mechanism might apply if the two genomes differ by insertion or deletion mutation at the affected locus. In such a scenario it could be, by analogy with epigenetic marks in *Neurospora crassa* that the transition is triggered by unpaired DNA during meiosis of the F1 [25]. The difference between the plant and fungal systems in the properties of the epigenetic mark: in *N. crassa* it would be a standard heritable mark with Mendelian inheritance whereas in tomato it would be a heritable mark that can transmit between alleles.

This requirement for unpaired DNA would account for the delayed onset of the paramutation-like process of *H06* until the meiosis in the reproductive phase of the F1 (Fig. 3). An alternative epigenetic explanation could invoke sRNAs from one of the interacting alleles that, in the parental line, are at too low an abundance to trigger paramutation. If the second allele has characteristics that favour production of secondary sRNA, for example repeats [5], then the formation of the hybrid would trigger a high level of sRNA and RdDM. The well established feedback loops of the RdDM process [26] would ensure that, once established at one allele, the amplified RdDM is stable and has the epigenetic drive characteristic of paramutation in subsequent generations.

These hypotheses are not mutually exclusive because both RdDM and meiotic silencing of unpaired DNA are dependent on an RNA-dependent RNA polymerase [27, 25]. The involvement of RdDM is supported by several lines of evidence that link small RNAs and in particular 24-nt sRNAs with paramutation-like events in tomato in addition to the findings from other species [11, 5, 12, 13]. At *H06*, for example, the timing of the paramutation-like process correlates with the timing of sRNA production (Fig. 3) and in VIGS the viral sRNAs produced could mediate (although inefficiently) the stable methylation of *H06* and the onset of paramutation-like

properties (Fig. 4). In addition to *H06*, out of four loci where the hypermethylated epiallele was associated with upregulated sRNAs, three showed evidence of paramutation-like behaviour (Hyper1, Hyper2 and Hypo3, Fig. 7 and Table 1).

There are, however, epialleles with paramutation-like characteristics (Hyper3, Hyper4, Hypo1 and Hypo2, Fig. 7) that are not associated with differential small RNAs. It remains possible that RNA-independent mechanisms are involved although some of these examples may be due to sRNA production and gain of methylation in tissues or developmental stages that are not well represented in the samples used for this analysis. Conversely, not all epialleles with differential sRNA have distinct epigenetic marks or, even if they do, are paramutagenic. Clearly paramutation requires additional, as yet unknown, factors beyond those associated with heritable epigenetic marks.

Stability of epialleles and penetrance of paramutation-like effects

For one locus with two epialleles (unmethylated and methylated), there are four possible states in a non-mosaic diploid organism: ‘unmethylated/unmethylated’, ‘unmethylated/methylated’, ‘methylated/unmethylated’, ‘methylated/methylated’ (Additional file 8: Figure S4). Local positive feedback loops in the maintenance of methylation [26, 27] or non-methylation [28] contribute to the stability of each state, while spontaneous epimutations and epiallele interactions in epi-heterozygotes trigger state transitions and distortions in the expected Mendelian ratios (Additional file 8: Figure S4).

Epialleles associated with paramutation in maize can be highly stable as with the *b1* locus, or unstable and revert to the active state, as with the *pl1* locus [29]. Similarly, with the paramutation-like loci in tomato there are varying degrees of stability. For some loci there were F1 plants without methylation (*H06*, Hyper1, Hyper2, Hypo1, Hypo2, Hypo3, Fig. 7), indicating a degree of instability of the methylated epiallele. Currently there are no clear mechanisms to explain the loss of methylation in an epi-heterozygote, apart from a dilution by a factor two of the sRNAs that could weaken a RdDM positive feedback loop acting in trans. The penetrance of the paramutation-like interaction (transition to the ‘methylated/methylated’ state) also varied: while 9/10 F1s had high methylation at Hyper1, only 5/10 were highly methylated at Hyper2 (Fig. 7). It is therefore crucial to garner information about stability and paramutagenicity of a large number of epialleles in order to identify their determinants. However we note that a binary depiction of the epigenetic state (e.g. methylated/unmethylated) is not complete: while *H06*^{IL} is very stably methylated, the paramutated *H06*^{IL} can revert when backcrossed to M82 (Fig. 2).

Frequency of paramutation in tomato hybrids

Paramutation with perfect efficiency will be fixed very rapidly in a population and will therefore be difficult to detect. Using a wide cross such as *Solanum lycopersicum* cv. M82 × *Solanum pennellii* LA0716 increases the likelihood of detecting highly penetrant paramutation events. We based our genome-wide search for paramutation on the characteristics of *H06* paramutation, which is associated with a gain of methylation in all sequence contexts and abundant sRNAs in multiple ILs at the seedling stage. We only found one other locus, Hyper1, that shared all of these characteristics, and this locus also displayed paramutation-like properties. However segregation analyses (Fig. 7) revealed that loci without differential sRNAs or without hypermethylation in all contexts could also engage in paramutation-like behaviours. Unexpectedly, loci that were hypomethylated in the ILs could also gain methylation upon crossing to M82.

These findings suggest that paramutation-like interactions are in fact common; occurring at hundreds of loci in a M82 × IL8–3 cross. However fully penetrant paramutation is rare: primary paramutation at *H06* was the most penetrant, but it displayed weak secondary paramutation (Fig. 2); and the penetrance of epigenetic changes and their inheritance in the seven validated loci (from seventeen tested) was incomplete. Therefore we conclude that full penetrance is rare and the IL crossing scheme may be too stringent (three consecutive backcrosses) to identify many traces of paramutation from the initial M82 × *S. pennellii* cross. Further studies aiming at assessing the prevalence of paramutation will need to be powered to detect incomplete penetrance.

Effects on gene expression

DNA methylation is primarily associated with the silencing of repeated sequences. Consistently earlier examples of paramutation are often linked with transposable elements [5]. Thus, as a mechanism of identity-based silencing, paramutation is thought to contribute to silencing multiple copies of transposable elements throughout the genome, and differential methylation may only occasionally also affect gene expression. We found no evidence for differential expression of the genes surrounding *H06* in the seedling expression data set. However *H06* is a target of the RIN (RIPENING INHIBITOR) transcription factor [30] and its surrounding genes are most highly expressed in the ripening fruit [31] (Additional file 9: Figure S5), so its differential methylation may have an effect in this particular tissue. For three of the validated DMRs that engaged in paramutation-like interactions (Hyper3, Hyper4 and Hypo3), the neighbouring genes were differentially expressed in the introgression lines compared to M82 (Additional file 6: Figure S2B).

Generating novel epialleles can be a source of phenotypic diversity [32, 33]. If the epialleles are stable enough and their epigenetic state can be efficiently controlled, epigenetic drives via paramutation may be used to improve crops.

Conclusions

In this study we show that interactions between natural tomato epialleles are relatively common, can lead to heritable differences in methylation, and occur mostly in one direction (gain of methylation) while stochastic loss of methylation may act as a counterbalance. There were however variations between pairs of epialleles in timing, penetrance and heritability of the gain of methylation. We propose that interactions between epigenomes should be viewed as a continuum, of which fully penetrant and fully heritable paramutation is an extreme manifestation. Taking into account the partial penetrance and stochasticity of epigenetic interactions will be crucial to properly investigate transgenerational inheritance.

Methods

Plant material

Tomato (*Solanum lycopersicum*) cv. M82, *S. lycopersicum* Micro-Tom, *Solanum pimpinellifolium* and *Solanum pennellii* plants were raised from seeds in compost (Levington M3) and maintained in a growth room at 23 °C with 16/8 h light/dark periods with 60% relative humidity, at a light intensity of 150 $\mu\text{mol photons m}^{-2}\cdot\text{s}^{-1}$. Unless otherwise indicated, *S. pennellii* refers to accession LA0716 that was used to generate the introgression lines [14].

DNA extraction

DNA from leaf (2-week-old seedlings), mature flowers or pollen was extracted with the Puregene kit (QIAGEN) following manufacturer's instructions.

Genotyping

IL8–3 was genotyped based on the TG510 marker: the sequence was amplified with primers TG510 fw and TG510 reverse, and digested with AluI. The M82 sequence is cleaved in a 133-bp and a 245-bp fragment, while the introgressed sequence from *S. pennellii* remains intact (378 bp). The *H06* sequence from M82 was differentiated from *S. pennellii* and *S. pimpinellifolium* by digesting *H06* amplicons with HpaI. The *S. pennellii* and *S. pimpinellifolium* are digested into two fragments, 260 and 210 bp in length, while the M82 sequence remains uncut. Even brightness of the *H06* genotyping bands for F1 s on an agarose gel indicate that McrBC-qPCR results should not be distorted by differences in DNA amplification efficiency between alleles.

RNA extraction

Total RNA samples were prepared from 100 mg of tissue using TRIzol (LifeTechnologies).

VIGS

H06 genomic insert was cloned into the binary TRV RNA2 vector using the KpnI and XhoI restriction sites of the multiple cloning site as described previously [16, 34]. Cotyledons of tomato seedlings were agro-infiltrated 10 d after sowing with a 1:1 mixture of *Agrobacterium tumefaciens* (strain GV3101:pMP90 + pSOUP) carrying TRV RNA1 and RNA2 at OD₆₀₀ = 1.5. 11 plants were successfully infected with TRV-H06 and 6 with wild-type TRV.

Viral load quantification

Quantification of viral load was performed with Precision One-Step qRT-PCR (Primerdesign) on 15 ng RNA per well and normalised with *TIP41*.

sRNA-Seq

sRNAs from leaf (2-week-old seedlings), flower and pollen of M82 (two individual plants), *S. pennellii* LA0716 (two plants) and their F1 (four plants) were cloned from 10 µg total RNA using the Illumina TruSeq Small RNA cloning kit and libraries were indexed during the PCR step (12 cycles) according to the manufacturer's protocol. Gel size-selected, pooled libraries were sequenced on a HiSeq 2000 50SE. Padubiri Shivaprasad prepared Micro-Tom (one plant) and *S. pimpinellifolium* (one plant) sRNA libraries from the aerial part of two-week-old seedlings according to [15]. Sequences were trimmed and filtered with Trim Galore! (with the adapter parameter '-a TGGAATTCTCGGGTGCCAAGG'). Mapping to *H06* (*S. pennellii* and M82 sequences) was performed with Bowtie 1.1.1 [35] without mismatches (options '-v 0 -a').

sRNA counts on *H06* in Fig. 1 were obtained by mapping sRNA libraries from [15] to Heinz genome SL2.50 with Bowtie 1.1.1 without mismatches (options '-v 0 -a'), and extracting the counts for the interval "SL2.50ch08:54487325-54,487,842". We found that there was a mistake in the labelling of the libraries deposited in [15]: the libraries labelled IL1-1, IL2-5, IL8-1-1, IL8-1-D, IL8-1-3, IL8-2, IL8-2-1, IL8-3 and IL8-3-1 actually correspond to IL8-3-1, IL8-3, IL8-2-1, IL8-2, IL8-1-5, IL8-1-D, IL8-1-1, IL2-5 and IL1-1. For convenience the correctly labelled sRNA libraries for the introgression lines used in this study (IL1-1, IL2-5 and IL8-3) are included in the GEO data set.

To find differential sRNA loci between M82 and ILs, sRNA libraries from IL1-1, IL2-5, IL8-3 and two M82 seedlings (from [15]) were mapped and clustered on Heinz genome SL2.50 using ShortStack v3.3.3 [36] with

default parameters. sRNA counts on the defined loci were analysed with DESeq2 v1.8.1 [37].

Gene expression analysis

Transcript counts for two-week-old IL1-1, IL2-5, IL8-3 and M82 seedlings from [17] were analysed with DESeq2 [37] with the design factors 'condition' (sun or shade), 'experimental batch', and 'genotype' (either IL or M82). In each of the five experimental replications, there were 4 M82 and 1 IL1-1, 1 IL2-5, and 1 IL8-3 in each treatment (sun or shade), totalling 40 M82 and 30 IL libraries on which to run the differential expression analysis. Genes were considered differentially expressed between genotypes when the adjusted *p*-value was for the 'genotype' factor was < 0.05.

Gene ontology enrichment analysis

Differentially expressed gene IDs were tested for gene ontology term enrichment against the full set of *S. lycopersicum* genes with the PANTHER Overrepresentation Test (Released 20,171,205) (<http://geneontology.org/>, GO Ontology database Released 2017-12-27, Fisher's exact test with FDR multiple test correction).

Analysis of DNA methylation

McrBC

Analysis of methylation by *McrBC* was performed as previously described in [16].

Sanger bisulfite

For Sanger bisulfite sequencing, 450 ng of DNA was bisulfite-converted with EZ DNA Methylation-Gold Kit (Zymo Research) and amplified with primers specific to the region and the Kapa Uracil+ HotStart DNA polymerase (Kapa Bioscience). Amplification products were size selected on a 1.5% agarose gel and gel extracted with the QIAquick gel extraction kit (QIAGEN), A-tailed following the same protocol as for the library preparation (below), and cloned into pGEM-T easy (Promega). Sequences aligned with MUSCLE were then analysed with CyMATE [38] and PCR-duplicates were removed. Each figure represents at least nine independent clones.

MethylC-Seq

Bisulfite library preparation was performed in duplicates for each genotype with a custom protocol similar to ref. [39]. 1.2 µg DNA was sonicated on a Covaris E220 to a target size of 400 bp and purified on XP beads (Ampure, ratio 1.8). DNA was end-repaired and A-tailed using T4 DNA polymerase and Klenow Fragment (NEB) and purified again using XP beads (ratio 1.8×). Methylated Illumina Y-shaped adapters for paired-end sequencing were ligated using Quick-Stick Ligase (BioLigne). 450 ng of purified (ratio 1.8×), adapter-ligated DNA was bisulfite-converted using

EZ DNA Methylation–Gold Kit (Zymo Research) according to manufacturer’s instructions. DNA was barcoded using 12 cycles of PCR amplification with KAPA HiFi HotStart Uracil+ Ready Mix (Kapabiosystems) with PE1.0 and custom index primers (courtesy of the Sanger Institute). Duplicate libraries for M82, IL1–1, IL2–5 and IL8–3 (leaves of two-week-old seedlings) were sequenced in paired-end mode. Reads were trimmed and filtered with Trim Galore! v0.4.2 (default parameters), then mapped on Heinz genome SL2.50 using Bismark v0.17.0 [40] (first in paired-end mode with options ‘–score-min L,0,-0.2 -p 4 –reorder –ignore quals –no-mixed –no-discordant –unmapped; then unmapped read1 was mapped in single-end mode with the same quality parameter ‘-N 1’). Reads were deduplicated with ‘bismark deduplicate’ and methylation calls were extracted using Bismark ‘methylation extractor’ (with options ‘-r2 2’ for paired-end reads). Based on Bismark’s cytosine report, methylated and unmethylated counts for cytosines of both strands were pooled into 300-bp bins sliding by 200 bp and separated by context (CG, CHG, and CHH). Bins with fewer than 10 counts or with coverage exceeding the 99 th percentile in one or more libraries were excluded from the analysis. We also excluded the introgressed regions: ch01 1–86 Mb, ch02 44–54 Mb and ch08 61 Mb – 65,866,657 bp (end of chromosome). Each introgression line was then compared to M82 by fitting a logistic regression on the methylated and unmethylated counts (R glm with ‘family = binomial’). The *p*-values were subjected to a Benjamini-Hochberg correction to control the false discovery rate at 5%. Differentially methylated bins were further filtered by imposing thresholds on the absolute difference in methylation (average of replicates) between M82 and the introgression line: a difference of 25% in the CG context, 20% in CHG and 10% in CHH. DMRs were constructed by merging overlapping differentially methylated bins. We included the region Hyper2 into the McrBC validation analysis despite it being inside the introgressed region in IL2–5, because it was hypermethylated in CG, CHG and CHH contexts and produced abundant sRNAs in IL1–1 and IL8–3, thus making it a likely candidate for paramutation. Chloroplast DNA was used as a control for bisulfite conversion efficiency, and sequencing statistics are collected in Additional file 1.

Oligonucleotides

Please refer to Additional file 10.

Additional files

Additional file 1: Summary of MethylC-seq data. Tsv format. (TXT 358 bytes)

Additional file 2: DMRs shared between introgression lines. Tsv format. (TXT 453 kb)

Additional file 3: DMRs shared between introgression lines and hypermethylated in all cytosine contexts. Average proportion of methylated cytosines between two replicates, for each context and genotype. Tsv format. (TXT 4 kb)

Additional file 4: DMRs shared between introgression lines and hypomethylated in all cytosine contexts. Average proportion of methylated cytosines between two replicates, for each context and genotype. Tsv format. (TXT 1 kb)

Additional file 5: Figure S1. Genomic position, small RNAs and DNA methylation of the IL DMRs Hyper1–9 and Hypo1–8. The selected DMR is highlighted in red, the plotted region includes 2 kb upstream and downstream. Genes: ITAG2.4 gene models. Repeats: RepeatMasker annotation. Hypomethylated regions shared between the three introgression lines are annotated in blue, and hypermethylated regions in red. sRNAs: coverage of sRNAs in seedlings. Methylation: percentage of methylated cytosines in 100-bp regions in each context (two replicates per genotype). (PDF 853 kb)

Additional file 6: Figure S2. A) Distribution of zero-count data for the genes differentially expressed between M82 and ILs (70 samples: 40 M82 and 30 ILs, data from [17]). B) Differential expression of genes whose promoters overlap shared IL DMRs selected for segregation analysis. Hypermethylation of Solyc09g064640’s promoter (Hyper3 and Hyper5) is associated with decreased expression in introgression lines. Solyc09g064640 codes for a protein related to the family with sequence similarity 65 member A (FAM65A). Solyc04g025030, encoding a MADS-box transcription factor similar to Agamous-like MADS-box protein AGL62, is associated with Hyper4, and Solyc06g075840, encoding a putative zinc-binding protein of the Yippee family, with Hypo3. Normalised counts in seedlings (40 M82 and 30 ILs, data from [17]). (PDF 64 kb)

Additional file 7: Figure S3. Segregation of DNA methylation patterns by McrBC. These regions show variable methylation in the parental controls M82 and IL8–3. (A) Hypermethylated DMRs Hyper7–9. (B) Hypomethylated DMRs Hypo6–8. For each region, the left hand panel shows the results of methylation analysis by McrBC for individual plants. This information is collated in the right hand panel with the splitting of the F2 s according to their F1 parent. Low methylation: ≤ 33%. Intermediate: > 33% and ≤ 66%. High: > 66%. (PDF 135 kb)

Additional file 8: Figure S4. Schematic of possible epiallelic transitions for one locus in a diploid state with unmethylated (white) and methylated (black) epialleles. There are four possible epigenetic configurations. Recurrent (curved) arrows represent the propensity to conserve the current state, from the action of local positive feedback loops maintaining methylation/unmethylation. Transition (straight) arrows between states represent spontaneous epimutations and, when starting from epigenetically heterozygous states, paramutation-like interactions. The epigenetic state of a genome may be seen as the result of such a stochastic process at each cell/organism generation. (PDF 2 kb)

Additional file 9: Figure S5. Potential role of *H06* in fruit ripening. Expression profiles across tissues of the two genes immediately downstream of *H06*, Solyc08g066020, encoding a Serine C-palmitoyltransferase like protein (A) and Solyc08g066030, encoding an unknown protein (B). Their expression peaks in mature green fruit. Data from [31] visualised on the eFP Browser [41] at http://bar.utoronto.ca/efp_tomato/. Copyright permissions for the re-use of the tomato drawings kindly granted by Prof. Provart. (C) Demethylation of the 3’ end of *H06* during fruit ripening correlates with the binding of the RIN transcription factor at this locus. RIN binding, demethylation and ripening are compromised in the *rin* (ripening inhibitor) and *cnr* (colorless non-ripening) mutants. Data from [30]. (PDF 971 kb)

Additional file 10: Oligonucleotides used in this study. Tsv format. (TXT 1 kb)

Additional file 11: *H06* sequences for *S. lycopersicum* cv. M82, *S. pimpinellifolium* and *S. pennellii* LA0716. Fasta format. (FASTA 1 kb)

Abbreviations

DMR: Differentially methylated region; IL: Introgression line; RdDM: RNA-directed DNA methylation; sRNA: Small RNA; VIGS: Virus-induced gene silencing

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Availability of data and materials

The datasets generated and analysed during the current study are available in the GEO repository under accession GSE97247 (<https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE97247>).

Authors' contributions

QG and DCB designed the research and wrote the manuscript, QG performed the research. All authors read and approved the final version of the manuscript.

Ethics approval and consent to participate

S. lycopersicum cv. M82, *S. lycopersicum* cv. Micro-Tom, *S. pimpinellifolium*, and *S. pennellii* LA0716 seeds were kindly provided by Tamas Dalmay. Seeds for the introgression lines were kindly provided by Dani Zamir. All plants were grown, propagated, crossed and collected in growth rooms at the Plant Growth Facility of the Department of Plant Sciences. Plant work was done under licence from the Department of Food and Rural Affairs (DEFRA) n° 50,973/198463 and VIGS experiments were conducted under DEFRA Plant Health licence 50,973/235589.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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