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CONSERVATION AND VARIATION OF DNA METHYLATION IN LACTUCA SATIVA AND LACTUCA SERRIOLA

A Dissertation Presented

by

TRUDI A. BAKER

Submitted to the Office of Graduate Studies, University of Massachusetts Boston, in partial fulfillment of the requirements for the degree of

DOCTOR OF PHILOSOPHY

December 2017

Biology Program

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CONSERVATION AND VARIATION OF DNA METHYLATION IN

LACTUCA SATIVA AND LACTUCA SERRIOLA

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TRUDI A. BAKER

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ABSTRACT

CONSERVATION AND VARIATION OF DNA METHYLATION IN LACTUCA SATIVA AND LACTUCA SERRIOLA

December 2017

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Directed by Professor Richard V. Kesseli

Molecular techniques for guiding plant breeding have successfully used wild progenitors of domestic crops as sources of genetic variants conveying desirable traits. However, epigenetic variation, in particular DNA methylation, is a significant source of phenotypic variation and epigenetic effects of plant domestication are poorly understood. Described herein are the first single-base pair resolution methylomes of the highly valued crop iceberg lettuce (*Lactuca sativa* cv. Salinas) and its close relative, and ubiquitous weed, *L. serriola*. This work suggests several roles for acquisition and inheritance of methylation in the evolution of *Lactuca* spp. in response to stress. The *Lactuca* spp. have conserved patterns of methylation around genomic regions associated with biotic stress response and conserved changes in average methylation levels in genic and intergenic regions under nutrient deprived conditions. The genotypes also have important differences in both methylation levels and variability in both control and nutrient deprived conditions. Additionally, there are suggestions that abiotic stress associated methylation may be transmitted between generations with fidelity. Together these findings suggest an additional source and mechanism of genomic variation which may be isolated and adapted for improvement of crops.

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Sincere thanks to Richard V. Kesseli for an extraordinary opportunity; to Bonnie A. Gulick, Jonathan D. Baker, Susan M. Huse for endless faith and encouragement; to Stuart H. Morey, Amy Avery, Michie Yasuda for generously sharing your expertise; and to David Weisman for introducing me to Python, which substantially improved the course of my career.

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Amplified fragment length polymorphism	AFLP
Argonaute 4	AGO4
Adenosine tri-phosphate	ATP
Base pair	bp
Chromomethylase 3	CMT3
Demeter	DME
Demeter-like protein 2	DML2
Demeter-like protein 3	DML3
Deoxyribonucleic acid	DNA
Dicer-like 3	DCL3
Differentially methylated cytosine	DMC
Differentially variable cytosine	DVC
Domains Rearranged DNA Methyltransferase 2	DRM2
False discovery rate	FDR
Flagellin-sensitive 2	FLS2
First self-ed generation	S1
Homologous recombination frequency	HRF
Individuals grown for two generations without fertilizer	NN
Individuals grown in nutrient deprived conditions whose parents were	
grown in control conditions	CN
Individuals grown for two generations in control conditions	CC
Individuals grown in control conditions whose parents were grown in	
nutrient deprived conditions	NC
Leucine-rich repeat	LRR
Long terminal repeat	LTR
Nucleotide	nt
Messenger ribonucleic acid	mRNA
Methylation sensitive amplified polymorphism	MSAP
Methylcytosine	mC
Methyltransferase 1	MET1
Mutation accumulator	MA
Mutator-like transposable element related locus	MULE-F19G14
Reduced representation bisulfite sequencing	RRBS
Repressor of silencing 1	ROS1
Restriction fragment length polymorphism	RFLP
Ribonucleic acid	RNA
RNA-dependent RNA polymerase 2	RDR2
RNA-directed DNA methylation	RdDM
RNA polymerase V	Pol V
RNA polymerase IV	Pol IV
Sawadee Homeodomain Homologue 1	SHH1
Single nucleotide polymorphism	SNP

LIST OF ABBREVIATIONS

Suppressor of variegation 3-9 homolog protein 2	SUVH2
Suppressor of variegation 3-9 homolog protein 9	SUVH9
Transcription start site	TSS
Transcription end site	TES
Variable In Methylation	VIM
Whole genome bisulfite sequencing	WGBS

CHAPTER 1

INTRODUCTION

Epigenetic diversity and its importance

To convey desirable traits such as increased yield or disease-resistance into domestic crops, plant breeders have traditionally used wild relatives of crops as sources of genetic variants conveying desirable traits [1–3]. This has been particularly effective in the many high value agricultural crops closely related to hardy weeds [2,3]. Improvement methods are traditionally targeted toward identification of discrete alleles to improve crops by selective breeding or transgenics [4,5]. However, recent studies have shown significant phenotypic variation associated with epigenetic diversity [6–9].

Epigenetics is broadly defined as "the study of mitotically and/or meiotically heritable changes in gene function that cannot be explained by changes in DNA sequence" [10]. Epigenetic modifications can include chemical modifications of nucleotides including methylation of cytosine producing 5-methylcytosine, the "fifth base" [11]. Epigenetic modifications also include chemical modification of proteins closely associated with genomic DNA, most notably the post-translational modification of histone proteins, as well as mitotically heritable protein-DNA associations [12]. Even conformation changes of prion proteins and cytoplasmic inheritance, transmission of

1

plastids, endosymbionts, viruses, and small RNAs through mitosis or meiosis, are sometimes broadly considered epigenetic [13–16]. Indeed, many of these concepts are interrelated. For example, DNA methylation in the CHG (H is any nucleotide but G) context positively reinforces the histone modification Histone3 Lysine9 [17]. The genome of the symbiotic plant root endophyte *Mesorhizobium loti* undergoes adenine methylation in the process of symbiosis, and these modifications are required for the efficient formation of nodules on the plants' roots [18]. Even the stalwart of traditional agricultural breeding programs, heterosis or hybrid vigor, has an epigenetic component [19,20].

Key epigenetic marks and mechanisms

DNA methylation will be the focus of this chapter and the following chapters. Though DNA methylation is one of many epigenetic mechanisms, it is highly prevalent, central to the interaction of other epigenetic mechanisms, and has unique characteristics in plant genomes. In plants, considerable methylation is found in each of the three possible sequence contexts for methylcytosine: CG, CHG and CHH, where H is any nucleotide except G. In plants, as in most eukaryotes, methylation is most frequently found in the CG context [21]. In differentiated human fetal fibroblasts more than 99.98% of methylation is found in the CG context [22], whereas in Arabidopsis immature floral tissue only a slight majority (55%) of methylcytosines are found in the CG context [23]. An additional differentiating characteristic of mammalian and plant epigenomes is the degree to which acquired DNA methylation is reset between generations. Methylation in mammalian genomes goes through two round of erasure during embryogenesis, whereas methylation in plant genomes, particularly in the CG and CHG contexts, are maintained through embryogenesis [24].

Methylation in different plant genomes share some common characteristics. Methylation within the CG context is more frequent than CHG or CHH within gene bodies [23,25–30]. In *Arabidopsis thaliana*, when the MET1-3 methyltransferases were knocked out, CHG methylation in euchromatic regions increased significantly and CHG methylation within gene bodies was enriched, taking on a similar profile to that of CG in wild-type plants [23]. Given the interrelation of the methylation and siRNA pathways, it is not surprising that Lister et al. (2008) found 85% of genomic regions with small RNA sequence identity contained at least one methylated cytosine; those methylation sites comprised 39% of all methylated sites [23].

Methylation polymorphisms and plant phenotypes

Methylation polymorphisms can be introduced stochastically due to a lack of fidelity of DNA methylation maintenance or by presence of a genetic variant such as a repetitive element insertion, or as a targeted response to environmental stimuli. Schmitz et al. (2011) estimated the rate of methylation polymorphism per CG to be 100,000 times greater than the rate of per nucleotide sequence polymorphism for the mutation accumulator (MA) lines of *A. thaliana*. This rate was based upon 30 generations derived from common ancestry. However, sites of methylation polymorphism between MA lines were not evenly distributed through the genome rather they were concentrated in certain

genomic locations [31,32]. This concentration of methylation polymorphism was also seen between members of geographically disparate wild populations of *A. thaliana* having diverged over a century ago [33]. This suggests that the fidelity of DNA methylation maintenance in plants may be highly variable across the genome, as has been shown in mouse embryonic stem cells [34].

Methylation affects plant phenotypes important for reproduction and fitness, including flower morphology [35], flowering time [36–38], sex determination [39], herbivore and pathogen resistance, [40–42] and agronomically important traits such as heterosis [19]. In general, abiotic stress associated methylation results have not been consistently mitotically or meiotically transmitted and, when detected, the direction of methylation change has been, in some cases, inconsistent across species and conditions. Treatment of Zea mays seedlings with intermittent heat, cold and ultraviolet stresses, for instance, did not result in condition-specific methylation patterns in adult plants [43], nor did methylation changes in rice correlate with salt treatment or the salt tolerance of the variety [44,45]. A recent study found both salt sensitive and resistant varieties of rice were globally hypomethylated in salt treatments [45]. In contrast, salt stressed A. thaliana were globally hypermethylated [46]. The duration of the stress and the length of time between stress treatments and tissue sampling may be important, but largely unconsidered, variables when comparing different stress methylation studies; methylation changes can be induced within a few hours of stress treatment [45,47] and a large proportion of induced changes may revert with time [48].

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Methylation can affect phenotype through many different mechanisms including modulating gene expression, transposable element mobility, and alternative splicing patterns. The relationship between methylation levels and gene expression levels differs by gene region. Methylation levels within the promoter regions are traditionally thought of as negatively correlated with gene expression levels through decreased binding affinity of transcription factors for methylated DNA [49] and reduced access of transcription factors and binding sites due to methylation-induced compact chromatin structure [50]. An example of negative correlation between promoter methylation and gene expression, is the expression of key genes involved in ethylene-induced ripening in tomatoes which requires active demethylation of their promoter regions by DNA glycosylases SIDML1/2 (orthologs of ROS1 in Arabidopsis) [27,51]. Interestingly, ROS1 itself is an important counter example to the generally inverse relationship between promoter methylation and gene expression. The ROS1 promoter is the target of both RdDM and active demethylation by ROS1 and transcription of the ROS1 gene is directly and positively correlated with the methylation levels in its promoter [52]. Thus the ROS1 promoter acts as a self-regulating rheostat, maintaining balance in its own methylation levels as well as ROS1 targets such as transposable element proximal genes [52]. Contrary to promoter regions, moderate methylation levels within gene bodies are associated with highly expressed genes, while high and low levels of methylation are associated with low expression levels [53]. Though methylation levels in both promoter and gene bodies are associated with expression, only a small percentage of differentially expressed genes in Arabidopsis and rice are associated with differential methylation [54], and only ~20% of

maize genes with on-off expression differences between inbred maize lines were associated with differentially ethylated regions [55].

Gene body methylation has also been associated with the suppression of transposable element insertion [56] and alternative splicing [57]. Mutator transposons insert preferentially into unmethylated regions [56,58], preferentially into genes, and more specifically into regions depleted in CG, but not CHG or CHH methylation [58]. The high average levels of CG methylation found in the gene bodies of most angiosperms could be an adaptive defense to transposable element insertion. Gene body methylation is also associated with prevalence of isoforms. Regulski (2013) analyzed sites of alternative splicing and found a bias in acceptor sites toward lower levels of CHG methylation, while levels of CHH methylation did not appreciably affect splicing efficiency. For honey-bee genes that are alternatively spliced, skipped exons are significantly hypomethylated relative to included exons, though in both cases exons have higher levels of methylation than flanking introns [57].

Transposable elements generally have higher average levels of methylation than intergenic regions [25,59], and methylation can serve to suppress their transcription and mobility [21]. During gametogenesis, passive demethylation and active DNA demethylation by DEMETER are associated with active transcription of transposons in the vegetative and central cells [21]. These transcripts travel to their respective egg or sperm cells and reinforce transcriptional silencing and RdDM of transposons [60]. In somatic tissue, hypomethylation in loss of function methyltransferase mutants in Arabidopsis resulted in a significant increase in transposon and pseudogene transcription relative to wild-type [23]. A variety of biotic and abiotic stresses are associated with hypomethylation [45,61,62] and treatment of Arabidopsis with the plant stress associated phytohormone salicylic acid resulted in hypomethylation of transposable elements and increased transcription of those elements [40]. However, not all releases of transposable elements are associated with removal of methylation. For example the release of silencing of Mutator-like transposable element related locus (MULE-F19G14) with temperature shifts occurs despite the maintenance of high levels of methylation and repressive histone modifications (H3K9/K27) [63]. And in an additional example, the binding affinity of the Tam3 transposase to its binding site in the sub-terminal repeat region of Tam3 is impaired by DNA methylation *in vitro*, but lack of methylation *in vivo* is not sufficient to induce transposition [64].

The presence and methylation status of transposable elements can affect the expression of proximal genes and potentially alter the organisms' fitness [40,65–67]. For example, genes which are up are upregulated in stress conditions in maize are enriched near certain families of transposable elements [68]. In Arabidopsis, genes which are downregulated in loss-of-function demethylase mutants with increased susceptibility to *Fusarium oxysporum* infection are enriched with transposable element regions in their promoters [69]. Methylation at neighboring transposable elements may have a positive or negative correlation to expression levels of nearby transposable genes [40,70].

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Transgenerational methylation effects

Specific environmental stresses to a parent can result in identifiable differences in their offspring's DNA methylation levels, their expression of stress-related genes, and their competitive ability in the stress environment [46,71]. In considering the evolutionary consequences of inheritance of acquired methylation, two possible scenarios can be considered. Stress associated DNA methylation could be directed towards the stress that is encountered, as has been suggested in mammalian nutritional studies. Several studies have found that offspring of parents with altered nutritional states (starvation, high fat, low-protein diets) had altered DNA methylation of metabolism related genes [71–73]. And in plants, members of the RdDM pathway play a role in transgenerational priming – the phenomenon where exposure to stress can make the individual or its offspring better poised to respond to future incidents of the stress [42]. Alternately, inheritance of stress associated DNA methylation could be beneficial, not by changing the mean level of methylation at a target, but rather by introducing stochasticity. Modeling supports the hypothesis that stochastic variation in mC would be advantageous in a disturbed environment [74]. Additionally, experimental data in dandelion shows increased epigenetic variation within individuals in stress treatments relative to unstressed groups [8]. In Arabidopsis, groups of genetically identical, but epigenetically diverse individuals, were more resistant to pathogen challenge and competition than less epigenetically diverse groups [75].

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Epigenetic effects of domestication of *Lactuca* **species**

Long histories of artificial selection have resulted in significant phenotypic divergence of domestic plants from wild relatives and decreased genetic diversity among selected domestic varieties. The epigenetic consequences of such prolonged selection are unknown. This dissertation seeks to provide a deeper understanding of the directional vs. stochastic hypotheses through a comparative analysis of the methylomes of domestic lettuce *Lactuca sativa* and its closely related wild and weedy relative *L. serriola*. *L. sativa* and *L. serriola* are particularly well suited for this study as they are self-fertile, populations tend to be highly homozygous. In the absence of genetic diversity, epigenetic diversity may be even more important. Additionally, *L. sativa* and *L. serriola* have very different tolerances for stress. *L. sativa* is commercially produced in a narrow range of environmental conditions and production is nutrient intensive, using more nitrogen fertilizer per acre than corn and most other vegetables [76]. *L. sativa* is also susceptible to many pathogens to which its wild relative *L. serriola* is substantially resistant [77].

The domestication history is well documented. Domestication of *L. sativa* has been traced to the Middle Eastern region encompassing modern day Iraq, Turkey, Syria, Lebanon, Israel, and the Egyptian river valley, referred to as the Fertile Crescent [78,79]. Significant cultivation of *L. sativa* is documented through the Grecian (450 B.C.) and Roman (1 A.D.) empires [78]. The first report of a heading type lettuce, *L. sativa* var. *capitate*, dates to a 1543 herbal book of German horticulturalist Leonhart Fuchs [78]. Though Linneaus (1757) classified *L. sativa* and *L. serriola* as distinct species, the accuracy of this taxonomic distinction and the exact relationship of *L. sativa* to *L*. *serriola* and other related species has been contentious. As early as 1851, Bischoff, Boissier, Hooker and Fiori contended that *L. sativa* and *L. serriola* were conspecific, differing only in degree of domestication. The relationships of these taxa were resolved with the application of molecular markers. Kesseli et al. 1991 showed that *L. serriola* alone was progenitor of *L. sativa* and that none of the 143 RFLP loci examined had diagnostic alleles that separated these taxa [80]. This was further confirmed with AFLP data [81]. Interestingly however, the major morphological groups of lettuce each appear to have a monophyletic origin suggesting that they arose from independent lineages of *L. serriola*, an idea first proposed by Sturtevant in 1886 [82]. The first report of *L. serriola* in the United States was in L.H. Pammel's 1863 report on distribution of weeds [78]. *L. serriola* is a particularly ubiquitous weed, found on all continents except for Antarctica, and as a common weed found throughout the lower 48 states [83]. *L. serriola* is a hardy weed commonly found beside highways and in other human disturbed environments.

The experiments described in the following chapters suggest several roles for acquisition and inheritance of methylation in the evolution of *Lactuca* spp. in response to stress. The *Lactuca* spp. have conserved patterns of methylation around genomic regions associated with biotic stress response and conserved, stress-induced changes in average methylation levels in genic and intergenic regions. These experiments also offer insights into the role of variability in methylation in the genotypes' differing response to stress conditions. Additionally, there are suggestions that abiotic stress associated methylation may be transmitted between generations with fidelity. Together these findings suggest an additional source and mechanism of genomic variation which may be isolated and

adapted for improvement of crops. Chapter 2 describes the first whole genome bisulfite sequencing of *Lactuca* species, and examines the differences in methylation patterns between *L. sativa* and *L. serriola* and other plant species through the lens of domestication. Chapter 3 introduces a novel method of characterizing whole genome patterns using reduced representation bisulfite sequencing. Chapter 4 looks at the impact of stress on the acquisition and the transmission of DNA methylation. Finally, Chapter 5 summarizes significant differences between the methylomes of these closely related species and suggests roles for DNA methylation in adaptation to environmental disruption.

CHAPTER 2

CONSERVATION AND VARIATION IN DNA METHYLATION IN *LACTUCA* SATIVA AND L. SERRIOLA

Introduction

To convey desirable traits such as increased yield or disease-resistance into domestic crops, plant breeders have traditionally used wild relatives of crops as sources of genetic variants conveying desirable traits [1–3]. This has been particularly effective in the many high value agricultural crops closely related to hardy weeds [2,3]. Improvement methods are traditionally targeted toward identification of discrete alleles to improve crops by selective breeding or transgenics [4,5]. However, recent studies have shown significant phenotypic variation associated with epigenetic diversity [6–9].

Methylation affects plant phenotypes important for reproduction and fitness, including flower morphology [35], flowering time [36–38], sex determination [39], herbivore and pathogen resistance, [40–42] and agronomically important traits such as heterosis [19]. In general, abiotic stress associated methylation results have not been consistently mitotically or meiotically transmitted and, when detected, the direction of methylation change has been, in some cases, inconsistent across species and conditions. Treatment of *Zea mays* seedlings with intermittent heat, cold and ultraviolet stresses, for instance, did not result in condition-specific methylation patterns in adult plants [43], nor did methylation changes in rice correlate with salt treatment or the salt tolerance of the variety [44,45]. A recent study found both salt sensitive and resistant varieties of rice were globally hypomethylated in salt treatments [45]. In contrast, salt stressed *A. thaliana* were globally hypermethylated [46]. The duration of the stress and the length of time between stress treatments and tissue sampling may be important, but largely unconsidered, variables when comparing different stress methylation studies; methylation changes can be induced within a few hours of stress treatment [45,47] and a large proportion of induced changes may revert with time [48].

Specific environmental stresses to a parent can result in identifiable differences in their offspring's DNA methylation levels, their expression of stress-related genes, and their competitive ability in the stress environment [46,71]. In considering the evolutionary consequences of inheritance of acquired methylation, two possible scenarios can be considered. Stress associated DNA methylation could be directed towards the stress that is encountered, as has been suggested in mammalian nutritional studies. Several studies have found that offspring of parents with altered nutritional states (starvation, high fat, low-protein diets) had altered DNA methylation of metabolism related genes [71–73]. And in plants, members of the RNA directed DNA Methylation (RdDM) pathway play a role in transgenerational priming – the phenomenon where exposure to stress can make the individual or its offspring better poised to respond to future incidents of the stress [42]. Alternately, inheritance of stress associated DNA

methylation could be beneficial, not by changing the mean level of methylation at a target, but rather by introducing stochasticity. Modeling supports the hypothesis that stochastic variation in mC would be advantageous in a disturbed environment [74]. Additionally, experimental data in dandelion shows increased epigenetic variation within individuals in stress treatments relative to unstressed groups [8]. In Arabidopsis, groups of genetically identical, but epigenetically diverse individuals, were more resistant to pathogen challenge and competition than less epigenetically diverse groups [75].

Many biotic stresses are associated with global or loci specific hypomethylation. Global loss of methylation is associated with increased resistance to infection by the bacterial pathogen *Pseudomonas syringae* in Arabidopsis [40] and upregulation of stress response genes in transgenic tobacco (*Nicotiana tabacum* cv. Xanthi) [47]. Treatment of rice with the methyltransferase inhibitor 5-azadeoxycytidine induced global hypomethylation and resistance to infection by *Xanthomonas* [84]. In addition, DNA glycosylase loss of function mutants have been shown to be more susceptible to fungal and bacterial pathogens and showed increased methylation and decreased expression of stress response genes [41,69]. Plants have evolved proteins, encoded by resistance genes, which recognize effector proteins of pathogens resulting in elicitor-triggered immunity [85,86].

The most prevalent class of resistance genes in plants are NBS-LRR proteins, which contain a leucine-rich repeat (LRR) domain [87]. Boyko et al. (2007) found significant hypomethylation and increases in homologous recombination frequency (HRF) in LRR domain containing genes in tobacco plants challenged with tobacco mosaic virus [62]. A key protein in the recognition of pathogen infection is the plant pattern-recognition receptor FLAGELLIN-SENSITIVE 2, it recognizes bacterial flagellin-derived peptide 22 (flg22). Flg22 triggers active demethylation by DNA glycosylase ROS1 and upregulation of some long terminal repeat (LTR) containing transposable elements and some LRR containing resistance genes [41]. In the absence of pathogen pressure, ROS1 constitutively demethylates these transposable elements in balance with constitutive transcriptional gene silencing. Pathogen pressure in a loss of function ROS1 mutant resulted in aberrant methylation in the CHH context of ROS1target LTR transposable elements [41].

Fungal and bacterial pathogens cause significant losses in the production of lettuce (*L. sativa*), the most consumed vegetable in the United States whose annual production is valued at approximately \$2 billion [76]. A close relative, and ubiquitous weed, *L. serriola* shows enhanced resistance to many of these pathogens. Early molecular work showed that *L. serriola* is the sole progenitor of *L. sativa*, supporting the contention that fully cross fertile *L. sativa* and *L. serriola* are conspecific [80,81]. Breeding efforts to enhance pathogen resistance in *L. sativa* include introduction of resistance genes from *L. serriola* [77,88–92]. In this paper we explore differences in the methylomes of *L. sativa* cv. *Salinas*, one of the mostly widely used elite cultivars in the breeding of modern crisphead lettuce varieties [93], and *L. serriola* (UC96US23), a pervasive and hardy weed, in the context of domestication and pathogen resistance phenotypes.

Methods

Samples and Extraction

Lactuca sativa cv. Salinas and *L. serriola* (UC96US23) seeds were obtained from Richard Michelmore's lab at the University of California Davis and the Compositae Genome Project (http://compgenomics.ucdavis.edu/). To reduce variation due to the maternal effect of different growing conditions for the different sources of seeds used in this study, two "progenitor" generations were planted (procedure described below). To avoid individual specific maternal effects, offspring from different self-fertilized parent plants were used as biological replicates. Seeds of each genotype were sterilized according to the following procedure: 1 mL 20% bleach solution and one drop Tween 20 were added to 25 seeds in a 2 mL microcentrifuge tube and gently agitated for 5 minutes. After a quick spin, detergent solution was decanted and 1 mL autoclaved, deionized water added and tubes gently agitated for 5 minutes.

This process was repeated for a total of 10 rinses. The seeds were refrigerated overnight at 4°C. Seeds were planted in commercial potting soil (Fafard Growing Mix 2: 70% Canadian sphagnum peat, 30% perlite and vermiculite) that had been autoclaved (25 minutes wet cycle) in two consecutive days preceding planting. The autoclaved soil was thoroughly moistened with autoclaved deionized water prior to filling half-gallon nursery pots. Sterilized seeds were then planted 2 seeds per container, at approximately 6 mm depth, covered with aluminum foil, then refrigerated at 4°C for 5 days.

Plants were randomly assigned positions within a 72 square grid in a Coviron® PGW36 Plant Growth Chamber at the University of Massachusetts Boston. Standard growth conditions were 16 hours of 800 µmol/m²/s intensity light at 23°C and 8 hours dark at 18°C. For the first two weeks in the growth chamber plants were watered 6 days per week with autoclaved deionized water. Thereafter plants were watered 2 times per week with unamended autoclaved deionized water and once with autoclaved deionized water supplemented with Peter's 20-20-20 all-purpose fertilizer at a concentration of 120 parts per million (N).

Tissue was collected from four biological replicates of *L. sativa* and *L. serriola*. In order to minimize variation between samples due to developmental differences, leaf tissue was collected when the first individual flowers of the secondary inflorescence are visible but still closed [28]. Samples were collected at a consistent time of day, between 1 and 2 hours prior to daybreak, to minimize variation in stress-related transcriptomes [94,95]. Two, 13 mm diameter leaf discs were placed in sterile containers and immediately immersed in liquid nitrogen.

Whole genome bisulfite library preparation and sequencing

DNA extractions were performed using MoBio's PowerPlant Pro DNA extraction kit with the following modifications: 40 μ l of Phenolic Separation Solution was added to 410 μ l of Solution PD1, 50 μ l Solution PD2 and 3 μ l RNaseA samples were added, then incubated at 65°C for 10 mins. Samples were further purified using MoBio's PowerClean Pro DNA Clean-up kit accord to manufacturer's instructions.

For each sample 1.4 µg DNA was fragmented using the Covaris S220 system (Covaris, Woburn, MA) in microTUBE AFA Fiber tubes (Covaris, Cat. No. 520045) to between 100 and 500 bp using the following instrument parameters: 80 s with a duty factor of 10%, a peak incident power of 175W, a temperature of 6°C and 200 cycles per burst. The sonicated DNA was purified using Qiagen DNeasy MinElute columns according to manufacturer's instructions. End-repair and ligation were performed using NEBNext® UltraTM DNA Library Prep Kit for Illumina® and NEBNext® Multiplex Oligos for Illumina[®] (Methylated Adaptors) according to manufacturer's instructions (New England Biolabs, Ipswich, MA). Ligation products were purified using a 1:1 ratio of Agencourt AMPure XP beads (Beckman Coulter Genomics, Danvers, MA) to product, and eluted in 30 µl of 1 x TE buffer. Libraries were size selected with a 1.5% Blue Pippin agarose gel cassette (Sage Science, Beverly, MA) for fragment sizes between 250 and 600 bp to collect fragments with a minimum 180 bp insert size plus the additional length of the ligated adapters. Isolated products were purified using a 1:1 ratio of Agencourt AMPure XP beads to product and eluted in 10 µl nuclease-free water. Bisulfite conversion was performed using NEB's Epimark Bisulfite Conversion kit according to manufacturer's instructions with an additional 5:1 bead clean-up.

Each WGBS library was PCR amplified in three separate 50 µl reactions, each containing 6.65 µl of bisulfite sample, 1 µl 10 mM dNTPs, 0.75 µl NEBNext universal PCR primer, 0.75 µl NEBNext index primer, 10 µl 5X EpiMark hot start Taq reaction buffer, 0.25 µl NEB EpiMark hot start Taq DNA polymerase and 30.6 µl ultrapure water. PCR conditions were 95°C for 30 seconds, followed by 13 cycles of 95°C for 15 seconds, 61° C for 30 seconds, and 68° C for 30 seconds, a final extension at 68° C for 5 minutes and hold at 4°C. Following PCR, the triplicate samples were pooled prior to bead purification. PCR products were purified using a 0.79:1 ratio of Agencourt AMPure XP beads to product, immediately followed by a subsequent bead purification using a 1:1 ratio and eluted in 20 µl 10 mM Tris, pH 8.0. Paired-end sequencing (2x100) was performed on a HiSeq 2000 at the University of Massachusetts Boston, Center for Personalized Cancer Therapy Genomics Core.

Raw reads were converted from bcl to fastq format using bcl2fastq (v. 1.8.4). Reads were trimmed using Trimmomatic (v 0.32) [96]. TruSeq3-PE adapter sequences were used as a reference for adapter trimming, allowing 2 nt mismatches with the adapter sequences, palindrome clip threshold of 30, simple clip threshold of 10. Overlapping paired end reads were merged using leeHom with the alignment of each library to the chloroplast genome serving as a prior [97]. The priors were estimated for each library based on paired end alignment of trimmed (but not merged) reads to the lettuce chloroplast genome.

The rate of bisulfite non-conversion was estimated by aligning paired reads to the bisulfite converted and bowtie2 indexed lettuce chloroplast genome (Accession number: NC_007578.1 by Bismark (v 0.13.1) [98]. Alignments were carried out in a two-step process. First, paired reads were aligned, with the –un option selected. Second, the reads which did not produce a valid, paired alignment were aligned in single read mode. Duplicate reads were marked and removed using Picard Tools' MarkDuplicates (v 1.96) [99]. The resulting bam files were then converted to the sam format using Picard Tools'

SamFormatConverter. Bismark's methylation_extractor script was run with the – bed_graph option which generates a 1-based report ("coverage" file) with the counts of the methylated and unmethylated reads detected at each position and summarizes these results over the entire genome. The coverage files were used as input to Bismark's coverage2cytosine which generates a text file summarizing the counts of methylated and unmethylated reads at each position in the genome regardless of whether any reads covered that position. All samples have apparent chloroplast methylation rates less than 5%.

Genome preparation, read alignment, and preliminary methylation counts

Access to the genome assemblies of *L. sativa* (v6) and *L. serriola* (v6) were generously provided by the Compositae Genome Project (http://lgr.genomecenter.ucdavis.edu; S. Reyes-Chin Wo, A. Kozik, D. Lavelle, and R.W. Michelmore, unpublished data). Sequences were bisulfite converted using Bismark's bismark_genome_preparation.

Trimmed reads were aligned to the bisulfite converted and indexed genome using Bismark [98] and bowtie2 [100]. Alignments were generated in the two step process described above, except the leeHom merged reads were also aligned as single reads in the second step. The resulting genome alignment files for each biological replicate were combined by replicate, and duplicate reads were marked and removed using Picard Tools' MarkDuplicates [99].

Bismark's methylation_extractor script was run with the –bed_graph option which generates a 1-based report ("coverage" file) with the counts of the methylated and unmethylated reads detected at each position and summarizes these results over the entire genome. The coverage files were used as input to Bismark's coverage2cytosine which generates a text file summarizing the counts of methylated and unmethylated reads at each position in the genome regardless of whether any reads covered that position.

Comparison of methylation between L. sativa and L. serriola by genomic feature

To calculate summary statistics for percent methylation by context over coding and repetitive regions, we summed the number of methylated reads and total reads of the biological replicates aligned to their respective genome sequences, selected only positions which were covered by five reads in all replicates of a genotype, and calculated the percent methylation by combining all replicates. These results were saved in the bed format and bedtools intersect [101] was used to define positions based on feature and feature proximity. Gene features were limited to those predicted loci also having transcriptional support and filtered to only the primary transcript per locus to avoid double counting. Repeat features were identified using RepeatMasker v4 [102].

Average methylation levels over protein coding genes and surrounding up- and downstream regions were calculated based on approximately 37,000 predicted genes in these genomes. Average levels were calculated over 100 bp bins in regions 10,000 bp upstream of the transcription start site (TSS), and 10,000 bp downstream of the transcription end site (TES) for all protein coding genes. Protein coding genes greater than 1000 nt in length were divided into 100 bins, and summary statistics computed for each bin position.
To detect differential methylation between *L. sativa* and *L. serriola*, reads from *L. serriola* were aligned to the *L. sativa* (v6) genome as described above. Only genome positions with at least three reads in each of the four *L. sativa* and *L. serriola* replicates and positions with non-zero variance in the proportion of methylated reads were retained for further analysis. Reads were analyzed in R using MethylSig [103]. Local information was included in the estimation of variance but not local methylation level. The local dispersion level was calculated across 8-9 orders of magnitude and repeated for tiled regions of: 1 bp, 10 bp, 100 bp, 1000 bp for each context. The smallest dispersion window that maximized detection was 1 Mb, corresponding to a window of +/- 1.17 cM. The differentially methylated cytosines (DMCs) with q-value <0.05 and a methylation difference >= 20% were considered significant. The predicted protein coding genes and repetitive features which overlapped with these DMC were identified using bedtools intersect.

Detection of differentially variable methylation between L. sativa and L. serriola

We utilized the iEVORA algorithm to test the null hypothesis of equal variances between biological replicates of *L. sativa* and *L. serriola* in the proportion of methylation at cytosines covered by at least ten reads using a q-value threshold of 0.001 [104,105]. The predicted protein coding genes and repetitive features which overlapped with these DMC were identified using bedtools intersect.

Gene ontology and KEGG annotations obtained from the Compositae Genome Project were used to perform gene ontology analysis of genes in differentially methylated and differentially variable regions. Hypergeometric testing using the R package phyper with false discovery rate (FDR) correction (p < 0.05) was performed to detect gene ontology terms over-represented in the entire set of genes containing one or more DMC, and for the sets located within 1 kb or 2 kb of an annotated repetitive element.

Results

Whole genome bisulfite sequencing of L. sativa and L. serriola

We performed whole genome bisulfite sequencing of four *L. sativa* and four *L. serriola* individuals, obtaining an average of 52 million high quality, deduplicated reads per individual. In *L. sativa*, the average methylation percent was 84.2% in the CG context, 70.3% in the CHG context, and 12.6% in the CHH context (Figure 1A). In *L. serriola*, average methylation levels were slightly lower in all sequence contexts; 77.4%, 64.5% and 10% respectively (Figure 1B). These methylation levels were comparable to other plant genomes of similar size (Figure 2A).

Among plant species the proportion of methylation at cytosines in the CG and CHG positions are significantly and positively correlated with genome size (CG: $R^2=0.57$, p-value=0.0046 and CHG: $R^2=0.78$, p-value=0.0002), whereas the relationship between methylation levels in the CHH context and genome size is weak and not statistically significant ($R^2=0.15$ and p-value=0.2149; Figure 3).

Both *L. sativa* and *L. serriola* have the characteristic bi-modal distribution of methylation levels with the vast majority (99.03% CG, 91.77% CHG, 90.14% CHH) of cytosine positions less than 20% or more than 60% methylated. In all sequence contexts the coefficient of variation for percent methylation at a position is inversely related to the

average percent methylation (Figure 4). These results are consistent with the observation in wild and domestic rice [54] suggesting an evolutionary role in conservation of highly methylated positions particularly those inhibiting the spread of transposable elements [60].

Previous work in other plants species has shown significant correlation of methylation levels for up to 5 kb [25]. We found the spatial autocorrelation of methylation in all contexts was highly consistent between all *L. sativa* and *L. serriola* replicates (Figs 5 and 6), though the degree of correlation differed significantly between the CG/CHG and the CHH context. The average correlation of methylation levels between cytosines separated by up to 50 kb is 0.82 in the CG context, 0.70 in the CHG context and 0.22 in the CHH context. The correlation between positions in CG or CHG contexts decreases after 50 kb, whereas the correlation does not vary significantly with genomic distance in the CHH context.

Methylation levels in both *Lactuca* ssp. took on familiar patterns in the regions up- and downstream of protein coding genes patterns where the relatively high levels of methylation in the CG and CHG contexts decrease sharply in the regions preceding the transcription start site, increase over the gene body, decrease towards the 3' end of the transcribed region, then increase in the downstream region. These patterns are very similar to those previously shown in *A. thaliana* [23,25] and other *Brassicaceae* [30], *Oryza. sativa* [26,54], *Populus trichocarpa* [26], *Manihot esculenta* [106], *Glycine max* [107], *Solanum lycopersicum* [27], and *Zea maize* [28] (Figure 6). *Lactuca* also shows increases in CHH methylation within a few hundred nucleotides upstream of the transcription start site and downstream of the poly-adenylation signal similar to previous reports in *Zea maize* [28] (Figure 7).

The average methylation percentage over 1,063 of the resistance genes in *L. sativa* [108] showed strikingly different patterns of methylation compared to the trends observed when considering all protein coding genes. In both *L. sativa* and *L. serriola*, the regions from 100 to 400 bp upstream and downstream of resistance genes were substantially more methylated in the CHH context compared to the average for other genes (Figure 8 A and B). The methylation percentage over the resistance genes themselves were low in all contexts with significant spikes at the 3' end of the genes (Figure 8 C).

Identifying differentially methylated cytosines between L. sativa and L. serriola

To identify cytosines with significant differences in mean methylation level between *L. sativa* and *L. serriola*, we aligned reads from biological replicates of *L. sativa* and *L. serriola* to the *L. sativa* genome sequence. We filtered the aligned positions and considered only positions covered by at least three reads in all replicates of both genotypes and having non-zero variance in the proportion of methylated reads between genotypes. There were 293,264 such cytosines in the CG, 257,984 in CHG, and 1,392,713 in CHH contexts. We tested for differential methylation using a beta binomial model across biological replicates [103]. Cytosine sites for which *L. sativa* and *L. serriola* had significant differences in methylation levels (q < 0.05) and also differed by at least 20% in mean methylation levels were considered for further analysis. There were 5,344 differentially methylated cytosines (DMCs) in the CG, 3,909 in the CHG, and 3,306 in the CHH contexts. Of these positions, 1,064 (7.81%) were associated with known sequence polymorphism between *L. sativa* and *L. serriola* and were excluded from further analyses of DMCs.

The mean level of methylation at DMCs was higher for *L. sativa* in all three contexts; 70% vs. 42% in the CG context, 67% vs. 34% in the CHG context, and 50% vs. 36% in the CHH context. In pairwise comparisons of these DMCs, *L. sativa* had the higher methylation level in 72%, 76% and 62% of positions in the CG, CHG and CHH contexts, respectively. These cytosines mostly were found in repetitive or unannotated regions (Figure 9 A-C). Most DMC within annotated repetitive elements are found in LTR retrotransposons (Figure 9 D-F).

We located 740 DMCs in 318 genes; 357, 282, and 101 DMCs in the CG, CHG and CHH contexts respectively. The majority (67.84%) of DMCs within genes had higher methylation levels in *L. sativa* than *L. serriola*. Genes with the highest frequency of DMC in their gene bodies were annotated with terms including hydroxylase activity, monooxygenase activity, electron carrier activity, transmembrane receptor activity, metal ion transport, and protein kinase activity (Table 1). The gene ontology terms including transmembrane receptor activity, intrinsic to membrane, serine-type endopeptidase activity, quinone binding, and oxidoreductase activity were enriched in DMC containing genes relative to their occurrence in the entire set of genes in the genome (Table 2).

As methylation in upstream regions of resistance genes in close proximity to repetitive elements has been associated with pathogen resistance [40,69], we looked at

DMCs in genes and regions 1 kb regions upstream that were also within 1 kb of a predicted repetitive element. Approximately half of all DMCs in genes, and 18% of DMCs in regions 1 kb upstream of genes, are located within 1 kb a predicted repetitive element. Of DMCs in genes located within 1 kb of a predicted repetitive element, 67% were more highly methylated in L. sativa; 83% of DMCs in upstream regions within 1 kb of a predicted repetitive element had a higher percent methylation in L. sativa. Methylation levels at DMC were significantly higher in L. sativa (p < 0.05) in the CG and CHG contexts, but did not significantly differ in the CHH context. Though the average methylation levels at DMC differed between L. sativa and L. serriola, within L. sativa or within L. serriola the level of methylation in upstream or gene regions was not significantly different between features located within 1 kb of a repetitive element and those greater than 1 kb from a repetitive element. Though methylation levels at DMC within upstream regions were significantly higher in L. sativa than L. serriola, methylation levels were not affected by proximity to repetitive regions (Figure 11 C and D). Gene ontology terms enriched in both genes and upstream regions within 1 kb of a predicted repetitive element included serine-type endopeptidase activity, isomerase activity, endonuclease activity, plastid, and carbon fixation, (Table 3 & Table 5).

Identifying differentially variable methylated cytosines between L. sativa and L. serriola

We utilized the iEVORA algorithm to test the null hypothesis of equal variances between biological replicates of *L. sativa* and *L. serriola* in the proportion of methylation at cytosines covered by at least ten reads using a q-value (FDR) threshold of 0.001 [104,109]. Very few positions (1.11%) co-localized with known sequence polymorphism and these were excluded from further analysis. In all sequence contexts the variability of methylation was greater in *L. sativa*: 55% of differentially variable methylated cytosines (DVCs) were more variable in *L. sativa* in the CG context, 77% in the CHG context and 92% in the CHH context. There were 378 DVCs in the CG context, 139 in the CHG context and 1,180 in the CHH context. Like DMCs, DVCs were mostly found in repetitive regions and unannotated regions, though only 3.71% of all DVCs were also DMCs and 0.5% of all DMCs also DVCs. Most DVCs within annotated repetitive elements were found in LTR retrotransposons (Figure 12 D-F). DVCs in all three sequence contexts were found within predicted protein coding genes (Figure 12 A-C). Genes containing DVCs were enriched for gene ontology terms structural constituent of ribosome, cytochrome-c oxidase activity, iron ion binding, ribosome, and mitochondrial electron transport cytochrome-c to oxygen (Table 7).

To investigate the spatial correlation of DMCs, DVCs, sequence polymorphisms, annotated genes and annotated repetitive elements, we divided each chromosome into 100,000 equally sized regions and counted the features starting within that region. The regions ranged from 2,271 to 4,369 bp in length depending on the chromosome. The number of DMC within a region was not strongly or significantly correlated with the number of SNPs, protein coding genes, or repetitive elements in that region (Figure 13A). The frequencies of DMCs in the three contexts are weakly, but significantly, correlated with each other (tau=0.18-0.22, p-value«0.01, representative values for chromosome 1). Similarly, the frequencies of DVCs in the three contexts are weakly, but significantly,

correlated (tau=0.48-0.49, p-value « 0.01, representative values for chromosome 1). The frequencies of DVCs in the three contexts are also weakly, but significantly, correlated with DMC in the regions (tau=0.17-0.3, p-value«0.01, representative values for chromosome 1) (Figure 13 B, values includes all chromosomes).

Gene ontology analysis of conserved methylation

We defined positions of highly conserved methylation as positions where the variability of percent methylation between biological replicates was in the lowest 25% for that *Lactuca* spp. and the average methylation level of *L. sativa* and *L. serriola* differed by less than 20%. In all sequence contexts, most sites of conserved methylation were located in regions without an annotated protein coding gene or repetitive element (Figure 14). The majority of positions with highly conserved methylation levels had low levels of methylation (Figure 16), and were found in non-repetitive intergenic regions (Figure 15). There were 249,770 cytosines with highly conserved methylation states, 74% of which had average methylation levels of less than 20%, 19.2% had average methylation levels greater than 60%. The majority of positions with conserved low levels of methylation were in the CHH context (80%), with 10.3% and 9.8% in the CG and CHG contexts respectively. In contrast, 51.9% of positions with conserved high levels of methylation were found in the CG context, 34% in the CHG context and 14.1% in the CHH context. The median levels of methylation at conserved sites were very high in all contexts within annotated genes and repetitive elements, and very low in genomic regions not known to contain genes or repetitive regions (Figure 15). There were 1,388 positions which were

100% methylated in all biological replicates of both *L. sativa* and *L. serriola* with an average read coverage of 24 reads and minimum read coverage of 11 reads. Forty-seven of these positions were found in seven genes, one of which was FLAGELLIN-SENSITIVE 2 a main sensor of bacterial infection; the remaining six genes are of unknown function (Table 8).

Discussion

Methylome characterization

When comparing the global methylation percentages in each sequence context of *L. sativa* and *L. serriola* to those of other genomes, a general positive relationship between percent global methylation and genome size is apparent, as has been detected previously using HPLC [110]. Similar to a recent report [111], we found that the relationship between methylation levels and genome size is sequence context dependent. There is a strong positive correlation between genome size and the total proportion methylation in the CG and CHG contexts, but that CHH methylation is only weakly correlated with genome size. Similarly, methylation levels at CG and CHG sites are highly correlated with methylation levels at sites within the same context that are separated by up to 50,000 bp, while methylation levels at neighboring CHH sites are less strongly correlated and that correlation does not vary significantly with distance. Both the correlation of methylation with genome size and spatial autocorrelation differences between the CG/CHG and the CHH contexts follow logically from the different mechanisms that maintain DNA methylation. Methylation in the CHH context must be

maintained by continual *de novo* RdDM targeted via small 21/24 nt RNA, where as methylation in the CG and CHG contexts is stably maintained through DNA replication by highly processive DNA methyltransferases MET1 and CMT3, [28].

DNA methylation has been proposed to act as a means of introducing stochasticity and evolutionary advantage to organisms in highly variable environments [74]. The overwhelming relative variability of CHH methylation in L. sativa, highlights the potential importance of differences in RdDM to significant phenotypic differences between L. sativa and L. serriola. We identified 1,697 cytosines having significantly different variances in the methylation between L. sativa biological replicates and L. serriola biological replicates. The majority of these positions were found in the RdDM driven CHH context (Figure 17). The proportion of DVCs which were more variable in the domestic L. sativa was much greater in the CHH context (92%) than in the CG (55%) or CHG (77%) contexts (Figure 18). The balance between RdDM and active demethylation by ROS1 may be finely turned for quick activation of pathogen defense response [41,112], an important phenotypic difference between L. sativa and L. serriola. Additionally, differences in the fidelity or kinetics of methylation or demethylation in the CHH context may contribute to differences in pathogen response and resistance. In Arabidopsis, loss of function mutants in the RdDM pathway harbored lower titers of bacteria than wild type plants [41]. If the RdDM pathway had a similar affect in *Lactuca* spp. this could contribute to the differences in pathogen tolerance between L. serriola relative to L. sativa.

The increased variation in methylation in domesticated lettuce is consistent with the increased methylation diversity seen in domesticated, relative to wild, soybeans [113]. These findings are particularly interesting in relation to the recent study by Latzel et al. that found populations of epigenetically diverse plants were more competitive as measured by increased plant density[75]. Increased planting density is a desirable trait selected for in agricultural production to increase crop yield, and the increased variability seen in *L. sativa* may be an adaptation to domestication.

In a pairwise comparison of sites in L. sativa and L. serriola, we found 13,623 differentially methylated cytosines (DMCs), most of which (92.19%) were not associated with known sequence polymorphisms between the species. Previous studies comparing methylation between related plant species have found the majority of methylation variation was associated with sequence diversity [29,30,54,114], though cases of pure epialleles, methylation variants independent of sequence, have been identified in maize [115], soybean [116], and *Arabidopsis* [31,117]. We found that the distribution of known SNPs was not significantly correlated with the distribution of DMCs in *L. sativa* and *L.* serriola and that the majority of DMCs between L. sativa and L. serriola did not colocalize with SNPs. Like Rambani et al. (2015) we found that the majority of DMC in protein coding genes were in the CG or CHG contexts (Figure 8 A and B) [118]. As with DVC, we found that the majority of DMCs were located in annotated repetitive regions (Figure 9 A, B and C). Becker et al. (2011) found fewer DMC among related Arabidopsis lines in regions near transposable elements, regions that were also enriched in small interfering RNAs [32]. However, we did not see an appreciable difference in the relative

frequency of DMC in protein coding genes or their upstream regions based on their proximity to annotated repetitive elements. Becker's strains were derived from a common ancestor 30 generations prior, whereas *L. sativa* and *L. serriola* are separated by centuries of selective breeding. It is possible the relative abundance of DMC and DVC in these regions in *Lactuca* species reflects an altered balance between RdDM and active demethylation between the species, and that the relatively short period of divergence between Becker's strains has not appreciably altered the balance between RdDM and active demethylation at these regions.

The majority of DMCs between *L. sativa* and *L. serriola* were more highly methylated in the domestic variety. Eichten et al. (2013) also found a positive association between location of differential methylation and repetitive elements, however they found that 81% of differentially methylated regions were more highly methylated in wild progenitor teosinte than in domesticated relative maize [114]. General categories of genes associated with domestication of crops, including transcription factors, enzymes and transporter proteins [119], were enriched among *Lactuca* genes containing a DMC.

In *Lactuca* both DMC and DVC are most prevalent in repetitive elements and could be associated with the high production of "sports" or rare unusual phenotypes in some varieties of *L. sativa* [120]. Though these sports appear to be generated randomly the variants, and their phenotypes, can be inherited similar to the stochastic generation [121,122] and inheritance with fidelity of differential methylation observed in offspring of regenerated plants [121]. Genomic stress of clonal propagation and tissue culture of plants are known to produce somatic variants or sports which are associated with

differential DNA methylation [121,122] and activity of transposable elements [123]. Though not grown clonally, the long history of inbreeding and selection to derive, in particular the heading varieties of, *L. sativa* from *L. serriola* could be thought of as a genomic stress.

It is tempting to hypothesize that the relative hypomethylation of DMCs in *L. serriola* may be associated with *L. serriola*'s superior performance in stressful and disturbed environments as has been suggested for other plant species. Resistance to biotic and non-biotic stresses have been associated with global [40,45,84] and loci specific [69] hypomethylation. Previous work in *A. thaliana* has shown that global loss of methylation is associated with increased resistance to infection by the bacterial pathogen *Pseudomonas syringae* [40]. Further, treatment of rice with the methyltransferase inhibitor 5-azadeoxycytidine induced global hypomethylation and resistance to infection by *Xanthomonas* [84]. In addition to the relative hypomethylation of *L. serriola* at DMC in general, all of the DMC within resistance genes (n=9) or within 1 kb upstream (n=10) or downstream (n=3) of resistance genes had higher percent methylation in *L. sativa* than *L. serriola* suggesting a particular role for methylation in and around these genes.

An interesting direction for future research would be to investigate active demethylation around resistance genes in *L. sativa* and *L. serriola*. The *L. sativa* and *L. serriola* genomes contain possible homologs to *A. thaliana* DNA glycosylases DEMETER and ROS1, but we did not find likely homologs to DML2 and DML3. In *A. thaliana* ROS1 is expressed in vegetative tissues while DEMETER is expressed in the endosperm and central cell during gametogenesis [124,125]. In *A. thaliana*, the region

between a Helitron transposon and ROS1's 5' UTR contains a DNA methylation monitoring sequence that is targeted by both RNA-directed DNA methylation and active de-methylation by ROS1 [126]. ROS1 expression is increased when methylated and decreased when de-methylated [52]. The active demethylation by ROS1 around genes in close proximity to repetitive elements has been associated with resistance to fungal and bacterial pathogens in A. thaliana. A triple demethylase mutant, deficient in all three known A. thaliana DNA glycosylases, was more susceptible to fungal infection and showed increased methylation and decreased expression of stress response genes with promoters in close proximity to transposons [69]. Yu et al. (2013) showed that growth of bacterial pathogen Pseudomonas syringae pv. tomato strain DC3000 was enhanced in ROS1 loss of function mutant, but not in loss of function mutants for the other DNA glycosylases, Demeter-like 2 (dml2) and Demeter-like 3 (dml3) [41]. Yu's work also showed that sRNAs accumulated in the region upstream of select resistance genes in Arabidopsis; regions where we see a spike in methylation in the CHH context in both species of Lactuca. Though the L. sativa and L. serriola methylation profiles around resistance genes are not significantly different on average from each other, analysis of differential rates of methylation and demethylation of resistance genes and flanking regions could highlight differences in pathogen responsiveness which could complement traditional gene based breeding approaches.

Our findings highlight significant differences in the methylomes of *L. sativa* and *L. serriola* that suggest future work in investigating epigenetic underpinnings of these closely related organisms' differing ability to adapt to disturbed environments.

Figures

Figure 1. Genome-wide levels of methylation are highly reproducible between biological replicates of *L. sativa* and *L. serriola*. Genome-wide levels of methylation in the CG, CHG and CHHs contexts are highly consistent among biological replicates of *L. sativa* (A) and biological replicates of *L. serriola* (B).



Figure 2. Genome-wide levels of methylation in plant species. Average methylation levels of *L. sativa* and *L. serriola* are high when compared to other plant methylomes especially in the CHH context (Figure 2A). Plants are listed from left to right in order of increasing genome size from *Arabidopsis thaliana* (135 MB) to *Zea mays* (2.5 GB), and literature reported genome sizes of each plant are plotted in Figure 2B. *Arabidopsis thaliana* [127], *Oryza sativa japonica* [128], *Oryza rufipogon* and *Oryza nivara* [129], *Oryza sativa indica* [130], *Brassica oleracea* [131], *Solanum lycopersicum* [132], *Glycine max* [133], *Zea maize* [134].



Figure 3. Correlation between genome size and genome-wide levels of methylation. There is a strong positive correlation between genome size and genome-wide levels of methylation in the CG (A) and CHG (B) sequence contexts for the 11 plant species shown in Figure 2. Methylation levels in the CHH sequence context (C) tend to be much lower and are not significantly correlated with genome size. Genome sizes are represented as C-values, the amount in picograms (pg) of DNA in a single copy of the genome.



Figure 4. Inverse relationship of variation and average methylation levels. In all sequence contexts the coefficient of variance ("CV") of a position across biological replicates is inversely related to the percent methylation at that position in both *L. sativa* (A-C) and *L. serriola* (D-F) for cytosines covered by more than 10 reads in all samples.



Figure 5. Spatial autocorrelation of methylation over long genomic distances. Methylation levels of cytosines in the CG (A) and CHG (B) contexts are highly correlated with methylation levels of cytosines in the same sequence context across increasing genomic distances, the x-axis refers to distance between adjacent positions and shows the correlation of positions separated by 0 to 100,000 bp. Cytosines in the CHH context (C) are not strongly correlated and this relationship is not affected by increasing genomic distance.



Figure 6. Spatial autocorrelation of methylation over short genomic distances. Methylation levels of cytosines in the CG (A) and CHG (B) contexts are highly correlated with methylation levels of cytosines in the same sequence context across increasing genomic distances, the x-axis refers to distance between adjacent positions and shows the correlation of positions separated by 0 to 1,000 bp. Cytosines in the CHH context (C) are strongly correlated only over very short (<100 bp) genomic regions, beyond this the degree of correlation is not affected by increasing genomic distance.



Figure 7. Average levels of methylation across protein coding genes and flanking regions. Average methylation in *L. sativa* for 10,000 bp preceding transcription start site, (TSS), average methylation over gene bodies where methylation averaged for each 100th of genes at least 1,000 bp long, and average methylation for 10,000 bp down stream of poly-A signal in CG (A), CHG (B) and CHH (C) contexts. *L. sativa* and *L. serriola* results do not differ appreciably, *L. serriola* results not shown.



Figure 8. Average levels of methylation across resistance genes and flanking regions. Average methylation of resistance genes in *L. sativa* for 1,000 bp preceding transcription start site, average methylation over gene bodies where methylation averaged for each 100th of genes at least 1,000 bp long, and average methylation for 1,000 bp down stream of poly-A signal in CH (A), CHG (B) and CHH (C) contexts. *L. sativa* and *L. serriola* results do not differ appreciably, *L. serriola* results not shown.



Figure 9: Distribution of DMCs across genomic regions. The majority of DMCs are located in repetitive regions in CG (A), CHG (B), and CHH (C) sequence contexts. The majority of DMCs in repetitive regions are located in CG (D), CHG (E), and CHH (F) sequence contexts in long terminal repeat (LTR) retrotransposons families.



Figure 10. Relative number of DMCs by sequence context and proximity of feature to annotated repetitive regions. The number of DMCs per protein coding gene (A.) or upstream region (B.) does not differ appreciably by proximity to repetitive regions.



Figure 11. Percent methylation of DMCs in *L. sativa* and *L. serriola* by sequence context and proximity of feature to annotated repetitive regions. Percentage methylation of DMCs by sequence context and genotype for protein coding genes within 1 kb of repeats (A) and not within 1 kb of repeats (B); for upstream regions of protein coding genes within 1 kb of repeats (C) and upstream regions not within 1 kb of repeats (D) Error bars represent 95% confidence interval.



Figure 12. Distribution of DVCs by genomic region. The majority of differentially variable cytosines are located in repetitive regions in CG (A), CHG (B), and CHH (C) sequence contexts. The majority of differentially variable cytosines in repetitive regions are located in CG (D), CHG (E), and CHH (F) sequence contexts in long terminal repeat (LTR) retrotransposons families.



Figure 13. The locations of DVCs and DMCs showed low to moderate correlation of abundance across the genome. Spearman correlation of the frequency of protein coding genes (mrna_counts) and repetitive elements (repeat_counts) and single nucleotide polymorphisms (snp_counts) with all differentially methylated cytosines combined (all_mc_counts) and all differentially variable cytosines combined (all_var_counts) (B) or differentially methylated cytosines by sequence context (cg_mc_counts, chg_mc_counts) and differentially variable cytosines by sequence context (cg_var_counts, chg_var_counts, chh_var_counts) (A).



Figure 14. Distribution of sites of conserved methylation by genomic region. Distribution of conserved methylated cytosines between *L. sativa* and *L. serriola* in CG (A), CHG (B), and CHH (C) sequence contexts in gene bodies, repetitive elements or other genomic regions, not known to contain genes or repetitive regions.



Figure 15. Percent methylation at conserved sites by genomic region. Sites of conserved methylation found within genes or repetitive regions have extremely high levels of methylation, whereas very low levels of methylation are found at sites of conversed methylation found in other genomic regions, not known to contain genes or repetitive regions.



Figure 16. Frequency of sites of conserved methylation by average methylation level. The majority of positions with highly conserved methylation states were found at positions which had very low levels of methylation. The graph shows the frequency of conserved positions arranged in order of increasing average methylation level. Positions with highly conserved methylation levels are those positions where the variability of percent methylation between biological replicates of each genotype was in the lowest 25% for that genotype and the methylation percentage between biological replicates of *L. sativa* and biological replicates of *L. serriola* differed by less than 20%.



Figure 17. Frequency of DVCs by sequence context. The majority of DVCs between *L*. *sativa* and *L. serriola* are found in the CHH context.



Figure 18. Proportion of DVCs that are more variable in *L. sativa* or *L. serriola*. Most differentially variable cytosines have higher levels of variability in *L. sativa* than in *L. serriola*.



Tables

Table 1. Protein coding genes with frequent occurrence of DMCs. The location column indicates whether or not the gene is located within 1 kb of an annotated repetitive element.

mRNA ID	Location	Freq.	Average %	Average %	Gene Ontology	KEGG ID
		DMC	IIIC L. Sauva	serriola		
Lsat_1_v5_gn_5_157381	one kb	53	42.52687331	16.65685226	GO:0004553: hydrolase activity, hydrolyzing O- glycosyl compounds: Molecular Function	endo-1,4-beta- mannosidase (EC:3) K01567 [EC:3]
Lsat_1_v5_gn_3_81720	more than one kb	23	51.26227875	2.423321597		
Lsat_1_v5_gn_4_121121	more than one kb	18	40.47586824	9.118170134	GO:0004497: monooxygenase activity: Molecular Function GO:0005506: iron ion binding: Molecular Function GO:0009055: electron carrier activity: Molecular Function GO:0020037: heme binding: Molecular Function	CYP82C4 electron carrier/ heme binding / monooxygenase/ oxygen binding K00517 [EC:1.14]
Lsat_1_v5_gn_5_60241	one kb	18	86.64087226	4.058201301		
Lsat_1_v5_gn_4_120360	more than one kb	17	32.75042683	61.64468393	GO:0004497: monooxygenase activity: Molecular Function GO:0005506: iron ion binding: Molecular Function GO:0009055: electron carrier activity: Molecular Function GO:0020037: heme binding: Molecular Function	CYP82C4 electron carrier/ heme binding / monooxygenase/ oxygen binding K00517 [EC:1.14]
Lsat_1_v5_gn_4_186001	more than one kb	15	46.26032561	87.42532016		
Lsat_1_v5_gn_8_52201	more than one kb	14	35.47441458	8.743896814		EFR EFR (EF-TU RECEPTOR) ATP binding / kinase/ protein

mRNA ID	Location	Freq.	Average %	Average %	Gene Ontology	KEGG ID
		DMC	mC L. sativa	serriola		
						serine/threonine kinase K13428 LRR receptor-like serine/threonine- protein kinase EFR [EC:2.7.11.1]
Lsat_1_v5_gn_8_51501	more than one kb	12	44.96402592	11.99408829	GO:0004888: transmembrane receptor activity: Molecular Function GO:0005524: ATP binding: Molecular Function GO:0006915: apoptosis: Biological Process GO:0007165: signal transduction: Biological Process GO:0031224: intrinsic to membrane: Cellular Component	
Lsat_1_v5_gn_8_51520	more than one kb	11	44.04897287	1.09796669		EFR EFR (EF-TU RECEPTOR) ATP binding / kinase/ protein serine/threonine kinase K13428 LRR receptor-like serine/threonine- protein kinase EFR [EC:2.7.11.1]
Lsat_1_v5_gn_7_103821	more than one kb	10	25.11171772	72.65490565		hypothetical protein LOC100252654 K10302 F-box protein 22
Lsat_1_v5_gn_1_116300	more than one kb	9	51.25746895	1.266484043		
Lsat_1_v5_gn_2_37560	more than one kb	9	3.489673888	86.27979218	GO:0016020: membrane: Cellular Component GO:0030001: metal ion transport: Biological Process GO:0046873: metal ion transmembrane transporter activity: Molecular Eunction	

mRNA ID	Location	Freq.	Average %	Average %	Gene Ontology	KEGG ID
		DMC	mC <i>L. sativa</i>	mC L. serriola		
Lsat_1_v5_gn_2_79801	more than one kb	8	70.64564469	1.383656089	GO:0004672: protein kinase activity: Molecular Function GO:0004674: protein serine/threonine kinase activity: Molecular Function GO:0005488: binding: Molecular Function GO:0005524: ATP binding: Molecular Function	protein kinase, putative K00924 [EC:2.7.1]
Lsat_1_v5_gn_2_131781	more than one kb	8	62.66904651	22.30709298		
Lsat_1_v5_gn_4_118880	one kb	7	30.52014604	0.831021859	GO:0004252: serine-type endopeptidase activity: Molecular Function	clpP ATP- dependent Clp protease proteolytic subunit K01358 ATP-dependent Clp protease, protease subunit [EC:3.4.21.92]
Lsat_1_v5_gn_7_15020	one kb	7	9.758959529	90.23934699	GO:0003677: DNA binding: Molecular Function GO:0003700: sequence- specific DNA binding transcription factor activity: Molecular Function GO:0005515: protein binding: Molecular Function GO:0005634: nucleus: Cellular Component GO:0006355: regulation of transcription, DNA- dependent: Biological Process	Pbx4, Edg4 pre-B- cell leukemia homeobox 4 K09355 pre-B-cell leukemia transcription factor
Lsat_1_v5_gn_4_75780	one kb	7	1.69929683	86.35457316		pex7 WD40 repeat- containing protein K13341 peroxin-7
Lsat_1_v5_gn_7_95881	more than one kb	7	43.32149422	3.53940139	GO:0003677: DNA binding: Molecular Function	rpoB RNA polymerase beta subunit (EC:2.7.7.6) K03043 DNA-

mRNA ID	Location	Freq.	Average %	Average %	Gene Ontology	KEGG ID
		DMC	mC <i>L. sativa</i>	mC <i>L</i> . serriola		
				serriora	GO:0003899: DNA-directed RNA polymerase activity: Molecular Function	directed RNA polymerase subunit beta [EC:2.7.7.6]
Lsat_1_v5_gn_1_49901	more than one kb	6	0.757508579	93.33249082		hypothetical protein LOC100247694 K11583 protein phosphatase 2 (formerly 2A), regulatory subunit B"Scaffold=Lsat_1_v 5 g 1 2399
Lsat_1_v5_gn_2_34800	more than one kb	6	54.35380201	6.387451436	GO:0009521: photosystem: Cellular Component GO:0009767: photosynthetic electron transport chain: Biological Process GO:0016020: membrane: Cellular Component GO:0016168: chlorophyll binding: Molecular Function	psbC photosystem II 44 kDa protein K02705 photosystem II CP43 chlorophyll apoprotein
Lsat_1_v5_gn_5_36381	one kb	6	0	94.0648275		
Lsat_1_v5_gn_7_7020	one kb	6	10.02364161	64.59990596		
Lsat_1_v5_gn_4_120300	more than one kb	5	39.69122293	5.909350423	GO:0004497: monooxygenase activity: Molecular Function GO:0005506: iron ion binding: Molecular Function GO:0009055: electron carrier activity: Molecular Function GO:0020037: heme binding: Molecular Function	CYP82C4 electron carrier/ heme binding / monooxygenase/ oxygen binding K00517 [EC:1.14]
Lsat_1_v5_gn_1_10481	more than one kb	5	1.670997948	60.01399436	GO:0003677: DNA binding: Molecular Function GO:0003824: catalytic	DNA polymerase I (POL I) K02335 DNA polymerase I [EC:2.7.7.7]
mRNA ID	Location	Freq. DMC	Average % mC <i>L. sativa</i>	Average % mC <i>L</i> . serriola	Gene Ontology	KEGG ID
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					activity: Molecular Function	
Lsat_1_v5_gn_4_144541	one kb	5	98.54095195	10		
Lsat_1_v5_gn_4_183940	more than one kb	5	71.83309789	46.59597312		
Lsat_1_v5_gn_6_37880	more than one kb	5	49.46149523	3.544937009		
Lsat_1_v5_gn_8_102340	more than one kb	5	97.74642394	41.81293883		
Lsat_1_v5_gn_1_8140	more than one kb	4	87.70722568	4.131184885	GO:0004497: monooxygenase activity: Molecular Function GO:0005506: iron ion binding: Molecular Function GO:0009055: electron carrier activity: Molecular Function GO:0020037: heme binding: Molecular Function	CYP81D3 electron carrier/ heme binding / monooxygenase/ oxygen binding K00517 [EC:1.14]
Lsat_1_v5_gn_4_23500	more than one kb	4	86.26581025	13.99313738		hypothetical protein K10610 DNA damage-binding protein 1
Lsat_1_v5_gn_4_154701	more than one kb	4	82.49343656	0.823882901		hypothetical protein LOC100260814 K01873 valyl-tRNA synthetase [EC:6.1.1.9]Scaffold =Lsat_1_v5_g_4_13 7
Lsat_1_v5_gn_3_92480	one kb	4	79.76345118	0	GO:0004672: protein kinase activity: Molecular Function GO:0004674: protein serine/threonine kinase activity: Molecular Function GO:0005524: ATP binding: Molecular Function	SNRK2.2 (SNF1- RELATED PROTEIN KINASE 2.2) kinase/ protein kinase K00924 [EC:2.7.1]
Lsat_1_v5_gn_4_54820	one kb	4	6.34674667	98.43807557	GO:0006629: lipid metabolic process: Biological Process	Zeta-carotene desaturase, chloroplast precursor, putative (EC:1.14.99.30) K00514 zeta- carotene desaturase

mRNA ID	Location	Freq. DMC	Average % mC <i>L. sativa</i>	Average % mC <i>L</i> .	Gene Ontology	KEGG ID
				serrioia		[EC:1.14.99.30]
Lsat_1_v5_gn_2_95981	one kb	4	95.73928962	0		
Lsat_1_v5_gn_3_17360	more than one kb	4	57.83039578	1.229783353		
Lsat_1_v5_gn_4_167780	more than one kb	4	41.02307881	70.2750932		
Lsat_1_v5_gn_7_67161	one kb	4	58.23682653	0.733750942		
Lsat_1_v5_gn_8_105980	one kb	4	73.80332287	10.27888622		
Lsat_1_v5_gn_8_125081	more than one kb	4	32.34516424	97.99682517		
Lsat_1_v5_gn_8_94660	one kb	4	48.62625699	47.15131292		

GO ID	GO description	Subset count	Genome count	Raw p-value	Adj. p-value
GO:0008289	lipid binding: Molecular Function	1	1	0	0
GO:0051287	NAD or NADH binding: Molecular Function	2	3	1 1894E-06	7 43376E-05
CO:0048028	quinone binding: Molecular	2	20	5 20646E 05	0.002211022
GO:0000521	runction	2	10	0.000125020	0.002211025
GO:0009521	photosystem: Cellular Component photosynthetic electron transport	2	10	0.000135029	0.002813112
GO:0009767	chain: Biological Process	2	10	0.000135029	0.002813112
GO:0016168	Function	2	10	0.000135029	0.002813112
GO:0003855	3-dehydroquinate dehydratase activity: Molecular Function	1	3	0.000335334	0.004657422
GO:0006467	protein thiol-disulfide exchange: Biological Process	1	3	0 000335334	0 004657422
	phosphoenolpyruvate carboxylase		2	0.0000000000	0.001657122
GO:0008964	tricarboxylic acid cycle: Biological	1	3	0.000335334	0.004657422
GO:0006099	Process	1	4	0.000665949	0.006936966
GO:0006334	Process	1	4	0.000665949	0.006936966
GO:0008250	oligosaccharyltransferase complex: Cellular Component	1	4	0.000665949	0.006936966
GO:0004556	alpha-amylase activity: Molecular Function	1	5	0.00110211	0 00918425
GO.0001550	shikimate kinase activity:		-	0.00110211	0.00010125
GO:0004765	endoplasmic reticulum membrane:	1	5	0.00110211	0.00918425
GO:0005789	Cellular Component	2	19	0.001015632	0.00918425
	NADH or NADPH: Molecular		20	0.001105105	0.0000 (1.005
GO:0016651	Function serine-type endopeptidase activity:	2	20	0.001185497	0.009261695
GO:0004252	Molecular Function	5	115	0.0014697	0.010799674
	diphosphooligosaccharide-protein				
GO:0004579	glycotransferase activity: Molecular Function	1	6	0.001641551	0.010799674
	oxidoreductase activity, acting on				
GO:0016671	as acceptor: Molecular Function	1	6	0.001641551	0.010799674
GO:0004888	transmembrane receptor activity: Molecular Function	7	210	0.00192669	0.01146839
CO:0021224	intrinsic to membrane: Cellular	7	200	0.001970177	0.0114(920
GO:0031224	shikimate 5-dehydrogenase	/	209	0.0018/01//	0.01140839
GO:0004764	activity: Molecular Function	1	7	0.002282039	0.01296613
GO:0004659	Molecular Function	1	11	0.005811072	0.031581912
GO:0016760	cellulose synthase (UDP-forming) activity: Molecular Function	2	36	0.006550426	0.034116801
GO:0004519	endonuclease activity: Molecular	1	14	0.009414681	0.042029824
00.0004515	photosynthetic electron transport	1	17	0.009414001	0.042029024
GO:0009772	in photosystem II: Biological Process	1	14	0.009414681	0.042029824
GO:0016853	isomerase activity: Molecular Function	1	14	0.009414681	0.042029824
GO:0030077	plasma membrane light-harvesting	1	14	0.009414681	0.042029824

Table 2. Enriched gene ontology terms for protein coding genes containing one or more

 DMCs.

GO ID	GO description	Subset count	Genome count	Raw p-value	Adj. p-value
	complex: Cellular Component				
GO:0019684	photosynthesis, light reaction: Biological Process	1	15	0.010787375	0.044746869
GO:0001522	pseudouridine synthesis: Biological Process	1	18	0.015392923	0.044746869
GO:0004143	diacylglycerol kinase activity: Molecular Function	1	17	0.013778379	0.044746869
GO:0005643	nuclear pore: Cellular Component	1	17	0.013778379	0.044746869
GO:0006855	drug transmembrane transport: Biological Process	2	49	0.015278703	0.044746869
GO:0006952	defense response: Biological Process	6	238	0.014121969	0.044746869
GO:0008131	primary amine oxidase activity: Molecular Function	1	16	0.012242574	0.044746869
GO:0008171	O-methyltransferase activity: Molecular Function	1	17	0.013778379	0.044746869
GO:0009308	amine metabolic process: Biological Process	1	16	0.012242574	0.044746869
GO:0009522	photosystem I: Cellular Component	1	18	0.015392923	0.044746869
GO:0015238	drug transmembrane transporter activity: Molecular Function	2	49	0.015278703	0.044746869
GO:0015297	antiporter activity: Molecular Function	2	49	0.015278703	0.044746869
GO:0015977	carbon fixation: Biological Process	1	18	0.015392923	0.044746869
GO:0016165	lipoxygenase activity: Molecular Function	1	17	0.013778379	0.044746869
	oxidoreductase activity, acting on single donors with incorporation of molecular oxygen, incorporation of two atoms of				
GO:0016702	oxygen: Molecular Function	1	17	0.013778379	0.044746869
GO:0005507	Function	4	139	0.016669006	0.047355131
GO:0009579	thylakoid: Cellular Component	1	19	0.017084369	0.04745658
GO:0007165	signal transduction: Biological Process	7	308	0.01782667	0.048442037
GO:0004629	phospholipase C activity: Molecular Function	1	20	0.01885091	0.049090912
GO:0009451	RNA modification: Biological Process	1	20	0.01885091	0.049090912

GO ID	GO description	Subset count	Genome count	Raw p-value	Adj. p-value
	endoplasmic reticulum membrane: Cellular				
GO:0005789	Component	2	19	0.000124423	0.003919194
	protein thiol-disulfide exchange: Biological				
GO:0006467	Process	1	3	7.97151E-05	0.003919194
	oligosaccharyltransferase complex: Cellular				
GO:0008250	Component	1	4	0.000158886	0.003919194
	serine-type endopeptidase activity:				0.004150550
GO:0004252	Molecular Function	4	115	0.000340708	0.004170553
GO:0004556	alpha-amylase activity: Molecular Function	1	5	0.000263907	0.004170553
	dolichyl-diphosphooligosaccharide-protein				
GO:0004579	glycotransferase activity: Molecular Function	1	6	0.000394512	0.004170553
	oxidoreductase activity, acting on a sulfur				
~~ ~~ ~~ ~	group of donors, disulfide as acceptor:				
GO:0016671	Molecular Function	1	6	0.000394512	0.004170553
00.0004650	prenyltransferase activity: Molecular			0.001.400007	0.010154000
GO:0004659	Function	1	11	0.001422087	0.013154302
GO:0004519	endonuclease activity: Molecular Function	1	14	0.002328997	0.017234575
GO:0016853	isomerase activity: Molecular Function	1	14	0.002328997	0.017234575
GO:0005643	nuclear pore: Cellular Component	1	17	0.003445371	0.023177948
GO:0015977	carbon fixation: Biological Process	1	18	0.003862895	0.023821184
	double-stranded RNA binding: Molecular				
GO:0003725	Function	1	21	0.005248288	0.025891553
00.0004/00	phospholipase C activity: Molecular		•	0.00.15(1500	
GO:0004629	Function	1	20	0.004764592	0.025891553
GO:0009536	plastid: Cellular Component	1	21	0.005248288	0.025891553
	acid phosphatase activity: Molecular				
GO:0003993	Function	1	26	0.007985868	0.035337434
GO:0006629	lipid metabolic process: Biological Process	4	240	0.008435545	0.035337434
	phosphoric diester hydrolase activity:				
GO:0008081	Molecular Function	1	27	0.008595592	0.035337434
GO:0010181	FMN binding: Molecular Function	1	28	0 009225524	0 035930987
55.0010101	i mini omanig. molecular i anction	*	20	0.007223324	0.000000000

Table 3. Enriched gene ontology terms for DMCs in protein coding genes within 1kb of an annotated repetitive element.

mRNA id	Freq. DMC	Average % mC Sativa	Average %mC Serriola	Gene Ontology	KEGG ID
Lsat_1_v5_gn_5_60241	35	69.92	22.87		
Lsat_1_v5_gn_4_127800	17	70.59	24.45		
Lsat_1_v5_gn_4_127821	17	70.59	24.45		
Lsat_1_v5_gn_8_116481	7	83.82	31.70	GO:0006508: proteolysis: Biological Process	hypothetical protein LOC100242026 K01365 cathepsin L [EC:3.4.22.15]
Lsat_1_v5_gn_9_115440	7	58.03	7.02	GO:0016651: oxidoreductase activity, acting on NADH or NADPH: Molecular Function GO:0048038: quinone binding: Molecular Function GO:0051287: NAD or NADH binding: Molecular Function	ndhH NADH dehydrogenase 49 kDa subunit K05579 NADH dehydrogenase I subunit 7 [EC:1.6.5.3]
Lsat_1_v5_gn_8_33421	7	97.10	19.11		nucleobase:cation symporter K03457 nucleobase:cation symporter-1, NCS1 family
Lsat_1_v5_gn_6_10001	7	73.87	2.94		
Lsat_1_v5_gn_5_137180	6	62.76	21.97	GO:0005488: binding: Molecular Function GO:0005743: mitochondrial inner membrane: Cellular Component GO:0006810: transport: Biological Process	Hypothetical protein CBG03436 K05863 solute carrier family 25 (mitochondrial carrier adenine nucleotide translocator)
Lsat_1_v5_gn_7_83821	5	41.96	39.39	GO:0004553: hydrolase activity, hydrolyzing O- glycosyl compounds: Molecular Function GO:0005618: cell wall: Cellular Component GO:0005975: carbohydrate metabolic process: Biological Process GO:0006073: cellular glucan metabolic process: Biological Process GO:0016762: xyloglucan:xyloglucosyl transferase activity: Molecular Function	Brassinosteroid-regulated protein BRU1 precursor, putative (EC:2.4.1.207) K08235 xyloglucan:xyloglucosyl transferase [EC:2.4.1.207]
Lsat_1_v5_gn_5_157381	4	39.08	21.86	GO:0004553: hydrolase activity, hydrolyzing O- glycosyl compounds: Molecular Function	endo-1,4-beta-mannosidase (EC:3) K01567 [EC:3]
Lsat_1_v5_gn_2_3840	4	0.00	58.90		hypothetical protein LOC100243760 K02116 ATP synthase protein IScaffold=Lsat 1 v5 g 2 5839
Lsat_1_v5_gn_9_51020	4	76.16	5.42	GO:0009055: electron carrier activity: Molecular Function GO:0016020:	petB cytochrome b6 K02635 cytochrome b6

Table 4. Frequency of DMCs in regions upstream of protein coding genes and within 1kb of a predicted repetitive element.

mRNA id	Freq.	Average	Average	Gene Ontology	KEGG ID
	DMC	% mC Sativa	%mC Serriola		
				membrane: Cellular	
				GO:0016491:	
				oxidoreductase activity:	
Last 1 v5 on 9 150591	4	02.91	0.00	Molecular Function	notossium ion transportar
LSat_1_v5_gi1_6_159561	4	95.61	0.00	ion transport: Biological	family protein K03549 KUP
				Process GO:0015079:	system potassium uptake
				potassium ion	protein
				transporter activity:	
		(7) 7		Molecular Function	
Lsat_1_v5_gn_1_101	4	65.43	95.08	GO:0006855: drug	
				transport: Biological	
				Process GO:0015238:	
				drug transmembrane	
				Molecular Function	
				GO:0015297: antiporter	
				Function GO:0016020:	
				membrane: Cellular	
Leat 1 v5 on 3 1800	4	19.81	29.99	Component	
$Lsat_1_v5_gn_5_1800$	4	73 71	29.99		
Lsat_1_v5_gn_4_100200		05.25	20.02		
Lsat_1_v5_gn_7_100541	4	85.35	1.92		
Lsat_1_v5_gn_7_38301	3	33.67	2.98	GO:0009522:	psaA photosystem I P700
				Component	K02689 photosystem I P700
				GO:0009579: thylakoid:	chlorophyll a apoprotein A1
				GO:0015979	
				photosynthesis:	
I sat 1 v5 on 2 89801	3	24.70	0.36	Biological Process	rbcI ribulose_1.5_
Lsat_1_v5_gi1_2_69601	5	24.70	0.50	magnesium ion binding:	bisphosphate
				Molecular Function	carboxylase/oxygenase large
				GO:0009536: plastid: Cellular Component	subunit (EC:4.1.1.39) K01601 ribulose-bisphosphate
				GO:0015977: carbon	carboxylase large chain
				fixation: Biological	[EC:4.1.1.39]
Lsat_1_v5_gn_1_42401	3	86.31	5.12	1100055	
Lsat_1_v5_gn_4_10101	3	95.95	0.00		
Lsat_1_v5_gn_4_185040	3	64.93	30.30		
Lsat_1_v5_gn_4_186001	3	97.04	31.46		
Lsat_1_v5_gn_5_168120	3	36.13	7.31		
Lsat_1_v5_gn_5_60261	3	72.36	38.20		
Lsat_1_v5_gn_2_96380	2	85.10	54.17	GO:0004672: protein	APK2A APK2A (PROTEIN
				kinase activity: Molecular Function	KINASE 2A) ATP binding / kinase/ protein kinase/ protein
				GO:0004674: protein	serine/threonine kinase
				serine/threonine kinase	K00924 [EC:2.7.1]
				Function GO:0005524:	

mRNA id	Freq.	Average	Average	Gene Ontology	KEGG ID
	DMC	% mC Sativa	%mC Serriola		
				ATP binding: Molecular	
Lsat_1_v5_gn_4_49540	2	92.61	5.56	GO:0005215:	Aquaporin PIP2.2, putative K09872 aquaporin PIP
				Molecular Function	K09072 uquuponn i n
				GO:0006810: transport: Biological Process	
Lsat_1_v5_gn_6_80221	2	75.46	0.00		ATLUP2 ATLUP2 beta-
					synthase K01853 cycloartenol synthase [EC:5.4.99.8]
Lsat_1_v5_gn_3_73580	2	100.00	21.29	GO:0004497: monooxygenase	CAld5H/F5H1, CYP84A10 coniferylaldehyde 5-
				activity: Molecular	hydroxylase K09755 ferulate-
				iron ion binding:	5-ilydroxylase [EC.1.14]
				GO:0009055: electron	
				carrier activity: Molecular Function	
				GO:0020037: heme	
				Function	
Lsat_1_v5_gn_8_11480	2	90.42	14.71	GO:0016872: intramolecular lyase	Chalconeflavonone isomerase, putative (EC:5.5.1.6) K01859
				activity: Molecular Function	chalcone isomerase
Lsat_1_v5_gn_8_42261	2	96.03	30.00	GO:0005622:	CO CO (CONSTANS)
				Component	transcription regulator/ zinc ion
					binding K12135 zinc finger protein CONSTANS
Lsat_1_v5_gn_9_22560	2	78.71	30.68	GO:0006508: proteolysis: Biological	hypothetical protein K01285 lysosomal Pro-X
L + 1 5 4 7040	2	(0.20	40.01	Process	carboxypeptidase [EC:3.4.16.2]
Lsat_1_v5_gn_4_/040	2	60.20	48.81	Molecular Function	AP-1 complex subunit gamma-
				GO:0006886: intracellular protein	1
				transport: Biological Process GO:0016192	
				vesicle-mediated	
				Process GO:0030117:	
				membrane coat: Cellular Component	
				GO:0030131: clathrin	
Leet 1	2	49.52	1.(1	Cellular Component	how the direct
Lsat_1_v5_gn_4_1/5880	2	48.53	1.01	acid binding: Molecular	LOC100247996 K13128 zinc
				Function	finger CCHC domain- containing protein 8
Lsat_1_v5_gn_8_36821	2	82.61	50.00	GO:0003677: DNA binding: Molecular	hypothetical protein
				Function GO:0003899:	DNA-directed RNA
				DINA-directed RNA polymerase activity:	[EC:2.7.7.6]
Lsat_1_v5_gn_2_100901	2	100.00	0.00	Molecular Function	leucine-rich repeat
					transmembrane protein kinase, putative (EC:1.3.1.74)
					K13420 LRR receptor-like

mRNA id	Freq. DMC	Average % mC Sativa	Average %mC Serriola	Gene Ontology	KEGG ID
					serine/threonine-protein kinase FLS2 [EC:2.7.11.1]
Lsat_1_v5_gn_1_111240	2	90.24	23.60		
Lsat_1_v5_gn_2_43920	2	92.26	9.03		
Lsat_1_v5_gn_2_62741	2	18.18	95.00	GO:0003676: nucleic acid binding: Molecular Function GO:0004519: endonuclease activity: Molecular Function GO:0006308: DNA catabolic process: Biological Process	
Lsat_1_v5_gn_2_98441	2	34.44	1.63		
Lsat_1_v5_gn_3_112240	2	88.54	27.54		
Lsat_1_v5_gn_3_138461	2	97.20	1.03		
Lsat_1_v5_gn_4_176941	2	53.44	47.31		
Lsat_1_v5_gn_4_180340	2	93.73	32.14		
Lsat_1_v5_gn_4_82360	2	69.85	0.00		
Lsat_1_v5_gn_5_105761	2	43.40	2.60		
Lsat_1_v5_gn_5_109401	2	83.47	76.04		
Lsat_1_v5_gn_5_121221	2	38.69	4.12		
Lsat_1_v5_gn_5_128341	2	45.76	95.65		
Lsat_1_v5_gn_5_154760	2	87.33	0.00		

Table 5.	Enriched gene ontolo	gy terms for	DMCs upstream	of protein	coding	genes	and
within 1	kb of an annotated re	petitive eleme	ent.				

GO ID	GO description	Subset count	Genome count	Raw p-value	Adj. p-value
GO:0006073	cellular glucan metabolic process: Biological Process	1	1	0	0
GO:0016762	xyloglucan:xyloglucosyl transferase activity: Molecular Function	1	1	0	0
GO:0030131	clathrin adaptor complex: Cellular Component	1	1	0	0
GO:0009522	photosystem I: Cellular Component	2	18	3.40369E-05	0.000470334
GO:0009579	thylakoid: Cellular Component	2	19	4.03144E-05	0.000470334
GO:0051287	NAD or NADH binding: Molecular Function	1	3	3.72573E-05	0.000470334
GO:0006308	DNA catabolic process: Biological Process	1	4	7.43415E-05	0.000743415
GO:0005742	mitochondrial outer membrane translocase complex: Cellular Component	1	6	0.000184992	0.001618681
GO:0016872	intramolecular lyase activity: Molecular Function	1	8	0.000343718	0.002673365
GO:0009521	photosystem: Cellular Component	1	10	0.000549846	0.003207436
GO:0009767	photosynthetic electron transport chain:	1	10	0.000549846	0.003207436

GO ID	GO description	Subset count	Genome count	Raw p-value	Adj. p-value
	Biological Process				
GO:0016168	chlorophyll binding: Molecular Function	1	10	0.000549846	0.003207436
GO:0016020	membrane: Cellular Component	7	539	0.000690914	0.00345457
GO:0030117	membrane coat: Cellular Component	1	11	0.000670477	0.00345457
GO:0015979	photosynthesis: Biological Process	2	51	0.00079771	0.003722646
GO:0004519	endonuclease activity: Molecular	1	14	0.001101644	0.00453618
GO:0016853	isomerase activity: Molecular Function	1	14	0.001101644	0.00453618
GO:0031461	cullin-RING ubiquitin ligase complex: Cellular Component	1	15	0.001268185	0.004931829
GO:0004190	aspartic-type endopeptidase activity: Molecular Function	2	71	0.002079611	0.006616145
GO:0005743	mitochondrial inner membrane: Cellular	2	72	0.002164466	0.006616145
GO:0015079	potassium ion transmembrane transporter	1	20	0.002268393	0.006616145
GO:0015977	carbon fixation: Biological Process	1	18	0.001835129	0.006616145
GO:0016651	oxidoreductase activity, acting on	1	20	0.002268393	0.006616145
GO:0048038	quinone binding: Molecular Function	1	20	0.002268393	0.006616145
GO:0009536	plastid: Cellular Component	1	21	0.002501376	0.006734475
GO:0016192	vesicle-mediated transport: Biological Process	2	75	0.002431838	0.006734475
GO:0004252	serine-type endopeptidase activity: Molecular Function	2	115	0.008027475	0.020068686
GO:0005975	carbohydrate metabolic process: Biological Process	2	114	0.007838015	0.020068686
GO:0004970	ionotropic glutamate receptor activity: Molecular Function	1	41	0.009326821	0.021762582
GO:0005234	extracellular-glutamate-gated ion channel	1	41	0.009326821	0.021762582
GO:0004221	ubiquitin thiolesterase activity: Molecular Function	1	46	0.011637708	0.025457487
GO:0042802	identical protein binding: Molecular	1	46	0.011637708	0.025457487
GO:0006511	ubiquitin-dependent protein catabolic	1	48	0.012625282	0.025535109
GO:0006855	drug transmembrane transport: Biological Process	1	49	0.013132342	0.025535109
GO:0015238	drug transmembrane transporter activity: Molecular Eurotion	1	49	0.013132342	0.025535109
GO:0015297	antiporter activity: Molecular Function	1	49	0.013132342	0.025535109
GO:0004650	polygalacturonase activity: Molecular	1	51	0.014172668	0.026107546
GO:0016887	ATPase activity: Molecular Function	2	141	0.013916917	0.026107546
GO:0006810	transport: Biological Process	4	413	0.016017268	0.028030219
GO:0006813	potassium ion transport: Biological Process	1	54	0.015797668	0.028030219
GO:0004553	hydrolase activity, hydrolyzing O-	3	275	0.016766635	0.028625963
GO:0006508	proteolysis: Biological Process	3	300	0.022257123	0.037095205
GO:0003899	DNA-directed RNA polymerase activity: Molecular Function	1	66	0.023038469	0.037504484

GO ID	GO description	Subset count	Genome count	Raw p-value	Adj. p-value
GO:0005618	cell wall: Cellular Component	1	73	0.027778767	0.044193493

Table 6. Gene ontology analysis of genes containing highly conserved methylation among replicates and between genotypes.

GOID	GO description	Subset count	Genome count	Raw p-value	Adj. p-value
	mitochondrial electron				
	transport, cytochrome c to				
GO:0006123	O:0006123 oxygen: Biological Process		2	0	0
	lipid binding: Molecular				
GO:0008289	Function	1	1	0	0
	ATP synthesis coupled proton				
GO:0015986	transport: Biological Process	3	24	3.20849E-07	5.88223E-06
	hydrogen ion transmembrane				
	transporter activity:				
GO:0015078	Molecular Function	3	32	1.07E-06	1.47125E-05
	proton-transporting two-				
	sector A l Pase complex,				
CO:0022177	Collular Component	2	12	2 70225E 06	4 17159E 05
00.0033177	cytochrome c ovidase	2	15	5.79255E-00	4.1/138E-03
GO:0004129	activity: Molecular Function	2	14	4 81823E-06	4 41671E-05
00.000412)	NAD or NADH binding:	2	14	4.010251-00	4.41071L-05
GO:0051287	Molecular Function	1	3	1 71898E-05	0.000135063
00.0051207	chloroplast: Cellular	1	5	1.710701-05	0.000155005
GO:0009507	Component	1	6	8.55458E-05	0.00052278
	ATP synthesis coupled	-			
	electron transport: Biological				
GO:0042773	Process	1	6	8.55458E-05	0.00052278
	microtubule motor activity:				
GO:0003777	Molecular Function	2	67	0.000578356	0.002164938
	NADH dehydrogenase				
	(ubiquinone) activity:				
GO:0008137	Molecular Function	1	13	0.000439986	0.002164938
	photosynthetic electron				
CO.0000772	transport in photosystem II:	1	14	0.000512514	0.0021(4029
GU:0009772	GO:0009//2 Biological Process		14	0.000512514	0.002164938
	nroton transport: Biological				
GO:0015991	Process	2	60	0.000418179	0.002164938
nhotosynthesis light reaction:		-	00	0.000110175	0.002101950
GO:0019684	Biological Process	1	15	0.000590438	0.002164938
plasma membrane light-					
	harvesting complex: Cellular				
GO:0030077	Component	1	14	0.000512514	0.002164938
	phosphatidylinositol				
	phosphate kinase activity:				
GO:0016307	Molecular Function	1	16	0.000673731	0.002315949
	oxidoreductase activity,				
CO 001((51	acting on NADH or NADPH:	1	20	0.0010(000)	0.002220151
GO:0016651	Molecular Function	1	20	0.001060086	0.003239151
CO:0048038	quinone binding: Molecular	1	20	0.001060086	0.003230151
00.0048038	U:0048038 Function		20	0.001000080	0.003239131
GO:0008375 activity: Molecular Eurotion		1	34	0.003062342	0.008864674
extracellular region: Cellular		-		0.003002372	0.0000007077
GO:0005576 Component		1	35	0.003242876	0.008917909
	lipid transport: Biological				
GO:0006869 Process		1	46	0.005545107	0.013862768

GO ID	GO description	Subset count	Genome count	Raw p-value	Adj. p-value	
	identical protein binding:					
GO:0042802	Molecular Function	1	46	0.005545107	0.013862768	
	photosynthesis: Biological					
GO:0015979 Process		1	51	0.006777992	0.016208243	
	structural constituent of					
	ribosome: Molecular					
GO:0003735 Function		3	331	0.008488664	0.018675061	
	ribosome: Cellular					
GO:0005840	Component	3	331	0.008488664	0.018675061	
	aspartic-type endopeptidase					
GO:0004190 activity: Molecular Function		1	71	0.01280657	0.027090822	

Table 7. Enriched gene ontology terms for protein coding genes containing DVC.

GO ID	O ID GO description		Genome count	Raw p-value	Adj. p-value
	mitochondrial electron transport,				
GO:0006123	CO:0006122 CO:0006122 Process		2	0	0
GU:0006123 Process		2	2	0	0
GO:0009060	Process	1	1	0	0
GO:0004129	cytochrome-c oxidase activity: Molecular Function	2	14	6 54545E-07	7 41817E-06
00.000112)	RNA-directed DNA polymerase	_		0.0 10 10 2 07	7.1101712.00
GO:0003964	activity: Molecular Function	1	6	2.28035E-05	0.000155064
	ATP synthesis coupled electron				
GO:0042773	transport: Biological Process	1	6	2.28035E-05	0.000155064
CO:0006461	Piological Process	1	0	5 45092E 05	0.00020626
00.0000401	respiratory chain complex IV	1	9	5.45985E-05	0.00020020
GO:0008535	assembly: Biological Process	1	8	4.24991E-05	0.00020626
	heme transporter activity: Molecular				
GO:0015232	Function	1	9	5.45983E-05	0.00020626
GO:0015886	heme transport: Biological Process	1	9	5.45983E-05	0.00020626
	NADH dehydrogenase (ubiquinone)				
GO:0008137	activity: Molecular Function	1	13	0.000117922	0.000400933
CO 001500(ATP synthesis coupled proton		24	0.000412(27	0.001270515
GO:0015986	transport: Biological Process	1	24	0.000413637	0.0012/8515
	transporter activity: Molecular				
GO:0015078	Function	1	32	0.000738651	0.002092844
GO:0015979	photosynthesis: Biological Process	1	51	0.001870407	0 004891834
00.0013717	microtubule motor activity:	1	51	0.0010/010/	0.001071051
GO:0003777	Molecular Function	1	67	0.003202742	0.007259549
	DNA-directed RNA polymerase				
GO:0003899	activity: Molecular Function	1	66	0.003109594	0.007259549
GO:0004190	aspartic-type endopeptidase activity: Molecular Function	1	71	0.00358829	0.007625117
	structural constituent of ribosome:				
GO:0003735	Molecular Function	2	331	0.00818573	0.015461935
GO:0005840	GO:0005840 ribosome: Cellular Component		331	0.00818573	0.015461935
GO:0006412 translation: Biological Process		1	119	0.009761785	0.017468457
GO:0003723	GO:0003723 RNA binding: Molecular Function		197	0.025249811	0.04129632
	iron ion binding: Molecular				
GO:0005506	Function	2	508	0.025506551	0.04129632
CO:000055	electron carrier activity: Molecular	2	524	0.020004007	0.044704070
00:0009055	runction	2	334	0.028984986	0.044/949/9

Chromosome	Start	End	Context	Gene ID	GO ID	KEGG ID
Lsat_1_v6_lg_4	350455078	350458757	CG	Lsat_1_v5_gn_4_ 152881	GO:0004672: protein kinase activity: Molecular Function GO:0004674: protein serine/threonine kinase activity: Molecular Function GO:0005515: protein binding: Molecular Function GO:0005524: ATP binding: Molecular Function	FLS2 FLS2 (FLAGELLIN- SENSITIVE 2) ATP binding / kinase/ protein binding / protein serine/threonine kinase/ transmembrane receptor protein serine/threonine kinase K13420 LRR receptor-like serine/threonine- protein kinase FLS2 [EC:2.7.11.1]
Lsat_1_v6_lg_4	350455078	350458757	CHG	Lsat_1_v5_gn_4_ 152881	GO:0004672: protein kinase activity: Molecular Function GO:0004674: protein serine/threonine kinase activity: Molecular Function GO:0005515: protein binding: Molecular Function GO:0005524: ATP binding: Molecular Function	FLS2 FLS2 (FLAGELLIN- SENSITIVE 2) ATP binding / kinase/ protein binding / protein serine/threonine kinase/ transmembrane receptor protein serine/threonine kinase K13420 LRR receptor-like serine/threonine- protein kinase FLS2 [EC:2.7.11.1]
Lsat_1_v6_lg_4	394333444	394338489	СНН	Lsat_1_v5_gn_4_ 167780		
Lsat_1_v6_lg_4	394333444	394338489	СНН	Lsat_1_v5_gn_4_ 167780		
Lsat_1_v6_lg_4	394333444	394338489	CG	Lsat_1_v5_gn_4_ 167780		
Lsat_1_v6_lg_4	394333444	394338489	CHG	Lsat_1_v5_gn_4_ 167780		
Lsat_1_v6_lg_4	394333444	394338489	CHG	Lsat_1_v5_gn_4_ 167780		
Lsat_1_v6_lg_4	394333444	394338489	СНН	Lsat_1_v5_gn_4_ 167780		
Lsat_1_v6_lg_4	394333444	394338489	CHG	Lsat_1_v5_gn_4_ 167780		
Lsat_1_v6_lg_4	394333444	394338489	CHG	Lsat_1_v5_gn_4_ 167780		
Lsat_1_v6_lg_4	394333444	394338489	CHG	Lsat_1_v5_gn_4_ 167780		
Lsat_1_v6_lg_4	432409453	432409740	CG	Lsat_1_v5_gn_4_ 183940		
Lsat_1_v6_lg_7	191848981	191850978	CHG	Lsat_1_v5_gn_7_ 96181		
Lsat_1_v6_lg_7	191848981	191850978	СНН	Lsat_1_v5_gn_7_ 96181		
Lsat_1_v6_lg_7	191848981	191850978	CHG	Lsat_1_v5_gn_7_ 96181		
Lsat_1_v6_lg_7	191848981	191850978	СНН	Lsat_1_v5_gn_7_ 96181		
Lsat_1_v6_lg_9	50682250	50685381	CHG	Lsat_1_v5_gn_9_ 38740		
Lsat_1_v6_lg_9	50682250	50685381	CG	Lsat_1_v5_gn_9_ 38740		

Table 8. Genes containing at least one cytosine which is covered by at least 10 reads and fully methylated in each biological replicate of both *L. sativa* and *L. serriola*.

Chromosome	Start position	End position	Context	Gene ID	GO ID	KEGG ID
Lsat_1_v6_lg_9	50682250	50685381	CG	Lsat_1_v5_gn_9_ 38740		
Lsat_1_v6_lg_9	50682250	50685381	CG	Lsat_1_v5_gn_9_ 38740		
Lsat_1_v6_lg_9	50682250	50685381	CG	Lsat_1_v5_gn_9_ 38740		
Lsat_1_v6_lg_9	50682250	50685381	CG	Lsat_1_v5_gn_9_ 38740		
Lsat_1_v6_lg_9	50682250	50685381	CG	Lsat_1_v5_gn_9_ 38740		
Lsat_1_v6_lg_9	50682250	50685381	CG	Lsat_1_v5_gn_9_ 38740		
Lsat_1_v6_lg_9	50682250	50685381	CG	Lsat_1_v5_gn_9_ 38740		
Lsat_1_v6_lg_9	88567292	88574707	CG	Lsat_1_v5_gn_9_ 60620		
Lsat_1_v6_lg_9	88567292	88574707	CG	Lsat_1_v5_gn_9_ 60620		
Lsat_1_v6_lg_9	88567292	88574707	CG	Lsat_1_v5_gn_9_ 60620		
Lsat_1_v6_lg_9	88567292	88574707	CG	Lsat_1_v5_gn_9_ 60620		
Lsat_1_v6_lg_9	88567292	88574707	CHG	Lsat_1_v5_gn_9_ 60620		
Lsat_1_v6_lg_9	88567292	88574707	CG	Lsat_1_v5_gn_9_ 60620		
Lsat_1_v6_lg_9	88567292	88574707	CHG	Lsat_1_v5_gn_9_ 60620		
Lsat_1_v6_lg_9	88567292	88574707	CG	Lsat_1_v5_gn_9_ 60620		
Lsat_1_v6_lg_9	88567292	88574707	CG	Lsat_1_v5_gn_9_ 60620		
Lsat_1_v6_lg_9	143112314	143121896	СНН	Lsat_1_v5_gn_9_ 80520		
Lsat_1_v6_lg_9	143112314	143121896	СНН	Lsat_1_v5_gn_9_ 80520		
Lsat_1_v6_lg_9	143112314	143121896	СНН	Lsat_1_v5_gn_9_ 80520		
Lsat_1_v6_lg_9	143112314	143121896	CG	Lsat_1_v5_gn_9_ 80520		
Lsat_1_v6_lg_9	143112314	143121896	CG	Lsat_1_v5_gn_9_ 80520		
Lsat_1_v6_lg_9	143112314	143121896	CG	Lsat_1_v5_gn_9_ 80520		
Lsat_1_v6_lg_9	143112314	143121896	СНН	Lsat_1_v5_gn_9_ 80520		
Lsat_1_v6_lg_9	143112314	143121896	СНН	Lsat_1_v5_gn_9_ 80520		
Lsat_1_v6_lg_9	143112314	143121896	СНН	Lsat_1_v5_gn_9_ 80520		
Lsat_1_v6_lg_9	143112314	143121896	СНН	Lsat_1_v5_gn_9_ 80520		
Lsat_1_v6_lg_9	143112314	143121896	СНН	Lsat_1_v5_gn_9_ 80520		
Lsat_1_v6_lg_9	143112314	143121896	CG	Lsat_1_v5_gn_9_ 80520		
Lsat_1_v6_lg_9	143112314	143121896	СНН	Lsat_1_v5_gn_9_ 80520		

CHAPTER 3

MODIFIED REDUCED REPRESENTATION BISULFITE SEQUENCING FOR PLANT GENOMES

Introduction

Reduced representation bisulfite sequencing (RRBS) allows a cost-effective whole genome survey suitable for assessing methylation in large genomes, and for assessing methylation variation across populations of individuals, tissues and treatments. The majority of RRBS studies, including the one example of RRBS of a plant genome [135], utilize the restriction endonuclease MspI. MspI is the most widely used enzyme in RRBS as its cleavage is insensitive to methylation in the predominant context for mammalian methylation, CG. However, MspI cleavage is blocked by methylation in the outer C of its recognition sequence (mCCGG). Unlike in mammalian species, the CHG (H = C, A, or T) context can account for a significant percentage of genome wide methylation in plants [23,25–28,30,54,107,136]. Methylation of proximal cytosines have been shown to be correlated [25,136], introducing the possibility that use of MspI in plants would bias calls of differential methylation.

Methods

To determine if this was the case we performed *in silico* digests of *Arabidopsis* genomes with known methylation states using MspI, a hypothetical MspI unaffected by methylation in its recognition site, and BssSI (C^ACGAG) and BsoBI (C^YCGRG). BssSI and BsoBI are insensitive to methylation in their recognition sequences and generate 5' overhangs that are filled in during the end-repair step of library preparation providing an internal control for bisulfite non-conversion. We based the *in silico* digestions of *Arabidopsis thaliana* on an existing data set to most realistically model the variability in read coverage and methylation distribution through the genome. The data set generated by Dowen et al. (2012) included four libraries: two biological replicates in control condition and two biological replicated treated with salicylic acid [40]. Reads were trimmed to remove adapters using Trimmomatic (v 0.32) [137], aligned to the *Arabidopsis thaliana* genome (TAIR 10) using Bismark (v 0.13.0) [98] and Bowtie2 (v 2-2.1.0) [100]. The number of C's and T's in the reads covering each reference cytosine were used as the reference methylation state for generating the *in silico* RRBS libraries.

In silico libraries were generated by identifying MspI, BssSI, and BsoBI recognition sequences located 200 to 700 bp apart in the reference genome; these positions flank the potential sequence fragments in the *in silico* RRBS library. For libraries representing MspI's actual cleavage, the total number of reads for each fragment was equal to the minimum number of methylated reads covering the outer C's of the fragment's flanking restriction sites. For libraries representing an unbiased MspI digest and a double digest with BssSI and BsoBI the total number of reads for each fragment

was equal to the minimum number of total reads covering the outer C's of the fragment's flanking restriction sites. The proportion of methylation for each C within the fragment was taken to be the mode of 1,000 samplings from a binomial distribution with p = proportion of all reads that were methylated in the reference methylation library for that position and n = the total number of reads for the fragment. To detect differential methylation we removed from consideration positions which were covered by fewer than 4 reads in the whole genome data set, and positions where the variance in proportion of methylation between treatments was less than the 25 percentile. For each library, processing the CG, CHG and CHH contexts independently, we used the R function *prop.test* from the *stats* package to test each covered position for significant differences in the percent methylation in the between control and treatment conditions. Multiple test correction was performed using the method of Benjamini and Hochberg [138]. Positions with adjusted p-values less than 0.05 were considered to be differentially methylated cytosines (DMCs).

Results and discussion

In silico analysis

In silico libraries covered 658,776, 1,919,848 and 870,462 in methylationsensitive MspI, methylation-insensitive MspI, and BssSI/BsoBI libraries respectively, representing 2.5%, 7.3% and 3.3% of cytosines covered by at least 4 reads in all replicates in the WGBS reference (Figure 1). We identified DMC between the salicylic acid treatment and control conditions in the methylation-sensitive MspI, methylation-

insensitive MspI, and BssSI/BsoBI in silico RRBS libraries. The BssSI/BsoBI digest covered slightly more genomic cytosines than the methylation-sensitive MspI digest, and the BssSI/BsoBI digest detected a much larger percentage of the DMC detected from the same genomics positions using the WGBS libraries (Figure 3). The *in silico* libraries representing the actual methylation-sensitive behavior of MspI detects 38-44% - 13% of the DMC detected using the same positions from the WGBS libraries, whereas the DMC detected in BssSI/BsoBI libraries represent 63-75% of DMC detected from the same positions in the WGBS libraries (Figure 3A). We repeated the analysis with a minimum read depth 10 reads per position, again more DMC were detected in BsoBI/BssSI RRBS libraries than actual MspI libraries (Figure 3B). However, the percentage of WGBS DMC detected in RRBS libraries varied with read depth and was lower at the higher read depth. DMC which were detected only in the RRBS libraries were found at positions with significant differences in the read coverage between DMC and RRBS libraries. There was no apparent sequence context bias between DMCs detected in RRBS-only, DMCs detected in both libraries or DMCs detected in only WGBS.

Reduced representation bisulfite sequencing of L. serriola

Additionally, we performed RRBS in the common weed *Lactuca serriola* using a double digest of BssSI and BsoBI. We generated libraries for each of three biological replicates of *L. serriola* grown without fertilizer and parental plants grown in control conditions (CN1, CN2, and CN3), and three biological replicates grown for two generations in without fertilizer (NN1, NN2, and NN3). DNA extractions and library construction methods are fully described in [139], important modifications to standard

protocols are noted here: 1.4 μ g of purified DNA was digested for 10 hours with 30 units of BssSI and 30 units of BsoBI (New England Biolabs, Ipswich, MA), enzymes were heat inactivated and reactions cleaned-up, prior to library preparation using NEBNext® UltraTM DNA Library Prep Kit for Illumina® and NEBNext® Multiplex Oligos for Illumina® (Methylated Adaptors) according to manufacturer's instructions (New England Bio- labs, Ipswich, MA). Ligation products were purified and libraries were size selected with a 1.5% Blue Pippin agarose gel cassette for fragment sizes between 250-600 bp (Sage Science, Beverly, MA). Each library was PCR amplified for 13 cycles in a single 50 μ l reaction containing 20 μ l of bisulfite treated sample. Paired-end sequencing (2x100) was performed on a HiSeq 2000 at the University of Massachusetts Boston Center for Personalized Cancer Therapy Genomics Core. Sequence QC and alignment methods are fully described in [139].

There were 1,325,486 cytosines that were covered by at least ten reads in all replicates, representing 1% of all cytosines in the *L. serriola* genome, and 62% of cytosines covered by at least 10 reads in WGBS of *L. serriola* [136,139]. Mean methylation levels of RRBS libraries were consistent between biological replicates in both conditions (Figure 4) and with levels reported in WGBS of *L. serriola* [136]. The majority of covered cytosines are found in the CHH context 62%, with 22% and 16% of cytosines found in the CG and CHG contexts respectively. The majority of methylated cytosines were located in repetitive regions (84%), 4% of cytosines were found in protein coding genes and 10% in unannotated regions (Figure 6). Though mean methylation levels did not appreciably differ between the two treatment conditions, the samples did

contain distinguishing differences in proportion methylation across covered cytosine positions as shown by hierarchical clustering (Figure 5). RRBS provides a significant cost-savings over WGBS while providing reproducible information on the methylation status of a significant subset the plant genome. However, the choice of enzymes is critical to the success of this technique. Here we introduce a method of selecting restriction endonucleases suitable for use in RRBS of plant genomes. We show *in silico* data predicting improved performance of RRBS using BssSI and BsoBI relative to the traditional enzyme of choice, MspI, and performed RRBS of *L. serriola* using BssSI and BsoBI covering with at least ten reads and 62% of cytosines covered by at least 10 reads in WGBS of *L. serriola* with reproducible methylation values between biological replicates.

Figures

Figure 1. Number of cytosines covered by *in silico* MspI and BssSI/BsoBI libraries. The number of cytosines in *Arabidopsis thaliana* covered in 200-700 bp fragments were calculated for *in silico* digests given MspI's actual cleavage, theoretical digest with an MspI not blocked by methylation in the C^mCG context, and double digest of BssSI and BsoBI.



Figure 2. Distribution of cytosines in gene bodies, repetitive elements or other genomic regions for *in silico* digests with MspI, ideal MspI, BssSI and BsoBI.



Figure 3. Percentage of significant DMCs by sequence context in *in silico* libraries. The percentage of significantly differentially methylated cytosines by sequence context was calculated for *in silico* MspI methylation sensitive libraries, MspI methylation insensitive libraries, and BssSI/BsoBI libraries relative to those detected in WGBS libraries considering positions found in both the RRBS and WGBS libraries.



Α.





Figure 4. Median genome-wide levels of methylation by sequence context for RRBS of *L. serriola*. Samples *L. serriola* plants were grown without fertilizer and parental plants grown in control conditions (CN1, CN2, and CN3) or without fertilizer (NN1, NN2, and NN3).



Figure 5. Hierarchical clustering of methylation in the CG (A), CHG (B), and CHH (C) contexts of *L. serriola* grown in different treatment conditions. In red are approximately unbiased probabilities generated by pvclust, in green are boot strap probabilities. CN1, CN2, and CN3 represent biological replicates grown in without fertilizer and parental plants grown in control conditions; N1, N2, and N3 represent biological replicates grown in no-fertilizer conditions.



Figure 6. Distribution of cytosines in *L. serriola* by genomic region. The proportion of cytosine positions covered with at least 10 reads in all replicates and conditions was determined for the CG (A), CHG (B), and CHH (C) sequence contexts in *L. serriola* gene bodies, repetitive elements or other genomic regions.



CHAPTER 4

REDUCED REPRESENTATION BISULFITE SEQUENCING OF *L. SERRIOLA* AND *L. SATIVA* SALINAS WITH DIFFERING FAMILY HISTORIES OF NUTRIENT DEPRIVATION

Introduction

Transmission of specific methylation in response to a stress has led some to suggest that DNA methylation is a mechanism for Lamarkian inheritance of acquired characteristics [140], while others suggest that acquired methylation serves to maintain phenotypic stochasticity in genetically homogeneous populations [74]. Both the Lamarkian and the stochastic variation models for the evolutionary effect of methylation could have profound consequences for the evolution of plants. In a Lamarkian sense, transmission of acquired methylation may pre-adapt offspring to the environment and result in a competitive advantage for individuals. Inheritance of stress associated DNA methylation could be directed towards the stress that is encountered. There are several examples in plants of transgenerational priming, improved fitness of offspring of stressed parents, including examples in radishes [141], *Arabidopsis* [42,46,142], monkey flowers [143], tomato [42], and tobacco [62]. In some cases these transgenerational benefits can

be reduced or eliminated in plants deficient in RNA directed DNA methylation or treated with a methylation inhibitor [42,46]. Alternately stochastic variation of methylation could explain the competitive success of invasive plants in highly disturbed environments. Feinberg & Irizarry (2010) modeled evolutionary consequences under different selection conditions and found that within a fixed environment the genotype with the greatest expected value for the desirable trait and the lowest stochastic variation was favored but in a variable environment the highly variable genotype was favored [74]. Indeed, Latzel et al. (2013) found that epigenetic diversity in *Arabidopsis* was associated with increased plant productivity (biomass) in environments challenged with plant competitors and pathogens [75].

Plants methylomes show significant effects of environmental stimuli. Natural populations of clonal plants have shown significant differences in methylation in different environments. For example, Gao et al. (2010) found differentially methylated loci in individual alligator weed clones grown in aquatic and terrestrial "common gardens" habitats regardless of the particular geography and habitat from which the plant was originally collected, suggesting that DNA methylation plays a role in adapting individuals to diverse habitats [144]. Results of controlled abiotic stress treatments have been very specific to the plant variety and experimental design. The duration of the stress and the time between when the plant experienced the stress and when the tissue was sampled may be important, but largely unconsidered, variables when relating different stress methylation studies, as methylation changes can be induced within a few hours of stress treatment [45,47]

and a large proportion of induced changes may revert with time [48]. For example, salt treatment of rice varieties showed global hypomethlyation in both sensitive and resistant varieties [44,45], though the degree and speed at which the changes accumulated differed by variety [45]. In contrast, salt stressed *A. thaliana* showed global hypermethylation, and local hypomethylation of abiotic stress response genes, and progeny of salt stressed plants showed increased germination and root length when grown in salt media [46].

Here we use RRBS to compare the acquisition and inheritance of methylcytosine between two *Lactuca* sp. with differing abilities to adapt to disturbed environments. Modern production of commercial lettuce requires moderate rates of nitrogen and phosphorous application. The domesticated variety *L. sativa* cv. Salinas was developed by the USDA in the 1980s and is the one of the mostly widely used elite cultivars in the breeding of modern crisphead lettuce varieties [93]. *Lactuca serriola* is a hardy weed commonly found beside highways and in other human disturbed environments. *L. serriola* is found on all continents with the exception of Antarctica, and is a common weed found throughout the lower 48 states [83]. The accession of *L. serriola* used in the present work (UC96US23) was originally derived from a plant growing in the parking lot of an abandoned gas station in Davis, CA.

Methods

Plant Growth, sample collection and DNA extraction

Lactuca sativa cv. Salinas and Lactuca serriola (UC96US23) seeds were obtained from the Richard Michelmore at the University of California Davis and the Compositae Genome Project (http://compgenomics.ucdavis.edu). To reduce variation due to the maternal effect of different growing conditions for the different sources of seeds used in this study, a progenitor generation was planted, grown and self-pollinated prior to the start of this experiment. Phenotypic measures of the next "parental" generation (S0) were collected. Plants were bagged to ensure self-fertilization, and seeds collected. Each biological replicate in the offspring generation (S1) was derived from a different individual in the parental generation. Seeds from this S1 generation were sterilized according to the following procedure: 1 mL 20% bleach solution and one drop Tween 20 were added to 25 seeds in a 2 mL microcentrifuge tube and gently agitated for 5 minutes. After a quick spin, detergent solution was decanted and 1 mL autoclaved, deionized water added and tubes gently agitated for 5 minutes. This process was repeated for a total of 10 rinses. The seeds were refrigerated overnight at 4°C. Seeds were planted in commercial potting soil (Fafard Growing Mix 2: 70% Canadian sphagnum peat, 30% perlite and vermiculite) that had been autoclaved (25 minutes wet cycle) each of the two days preceding planting for a total of 2 treatments separated by approximately 24 hrs. The autoclaved soil was thoroughly moistened with autoclaved deionized water, prior to filling half- gallon nursery pots. Sterilized seeds were then planted 2 seeds per container,

at approximately one-quarter in. depth, covered with aluminum foil, then refrigerated at 4°C for 5 days.

Plants were randomly assigned positions within a 72 square grid in a Coviron® PGW36 Plant Growth Chamber at the University of Massachusetts Boston. Standard growth conditions were 16 hours of 800 μ mol/m²/s intensity light at 23°C and 8 hours dark at 18°C. For the first two weeks in the growth chamber plants were watered 6 days per week with autoclaved deionized water.

Thereafter control plants (C) were watered 2 times per week with autoclaved deionized water and once with autoclaved deionized water supplemented with Peter's 20-20-20 all-purpose fertilizer at a concentration of 120 parts per million. Plants assigned to the nutrient deprived (N) treatment were watered 3 times per week with autoclaved deionized water and no fertilizer. Tissue was collected from three or four biological replicates of *L. sativa* and *L. serriola* in each treatment. In order to minimize variation between samples due to developmental differences, leaf tissue was collected when the first individual flowers of the secondary inflorescence are visible but still closed [145]. Samples were collected at a consistent time of day, between 1 and 2 hours prior to daybreak, to minimize variation in stress-related transcriptomes [94,95]. Two half-inch leaf discs were placed in sterile containers and immediately immersed in liquid nitrogen. *Reduced representation bisulfite library preparation and sequencing*

Library preparation proceeded as in [146] with the following exceptions. 1.4 μ g of purified DNA was digested for 10 hours with 30 units of BssSI and 30 units of BsoBI. Enzymes were heat inactivated and reactions cleaned-up, prior to end repair, A-tailing and ligation of methylated adapters according to manufacturer's instructions (NEB, Ipswich, MA). Ligation products were purified and libraries were size selected with a 1.5% Blue Pippin agarose gel cassette for fragment sizes between 250-600 bp. (Sage Science, Beverly, MA) Bisulfite treatment and clean-up were performed as above. Each library was PCR amplified for 13 cycles in a single 50 μ l reaction containing 20 μ of bisulfite treated sample. Paired-end sequencing (2x100) was performed on a HiSeq 2000 at the University of Massachusetts Boston Center for Personalized Cancer Therapy Genomics Core. Read were trimmed using Trimmomatic and overlapping paired end reads merged as described for whole genome bisulfite sequencing. The bisulfite non-conversion rate was estimated by aligning reads to the *L. sativa* chloroplast genome as described for whole genome bisulfite sequencing.

The genome assemblies of *L. sativa* (v6) and *L. serriola* (v6) were generously provided by the Compositae Genome Project. Sequences were bisulfite converted using Bismark's bismark_genome_preparation. Trimmed reads were aligned to the bisulfite converted and indexed genome using Bismark and bowtie2 using the two step process and counts of methylated and unmethylated reads for each position in the genome were generated as described for whole genome bisulfite sequencing [136]. For comparisons of DMCs between *L. sativa* and *L. serriola*, reads were aligned to the *L. sativa* (v6) genome. For computation of genome-wide methylation levels and DMC detection between samples within a genotype, *L. sativa* reads were aligned the *L. sativa* (v6) genome and *L. serriola* reads were aligned the *L. sativa* (v6) genome and *L. serriola* reads were aligned the *L. sativa* (v6) genome and *L. serriola* reads were aligned the *L. sativa* (v6) genome and *L. serriola* reads were aligned the *L. sativa* (v6) genome and *L. serriola* reads were aligned the *L. sativa* (v6) genome and *L. serriola* reads were aligned the *L. sativa* (v6) genome and *L. serriola* reads were aligned the *L. sativa* (v6) genome and *L. serriola* reads were aligned the *L. sativa* (v6) genome and *L. serriola* reads were aligned the *L. sativa* (v6) genome and *L. serriola* reads were aligned the *L. sativa* (v6) genome and *L. serriola* reads were aligned the *L. sativa* (v6) genome.

Detection of differential methylation

Only genome positions with at least ten reads in each replicate of each sample, which did not co-localize with known SNPs between *L. sativa* and *L. serriola* and which had greater than zero variance in percent methylation across all replicates were retained for further analysis. Reads were analyzed in R using MethylSig [103]. Local information was included in the estimation of variance but not local methylation level. The differentially methylated cytosines (DMCs) with q-value <0.05 and a methylation difference >= 20% were considered significant. The mRNA and predicted repetitive features which overlapped with these DMC were identified using bedtools intersect.

Detection of differentially variable methylation between L. sativa and L. serriola

We utilized the iEVORA algorithm to test the null hypothesis of equal variances of proportion of methylation between biological replicates of *L. sativa* and *L. serriola* using a q-value threshold of 0.001 [104,147]. The predicted protein coding genes and repetitive features which overlapped with these DMC were identified using bedtools intersect.

Results

Relative contribution of environment and genotype to DNA methylation

Lactuca sativa and *L. serriola* differ in their ability to reproduce in disturbed environments. Here we exposed both species to controlled, with fertilizer, conditions (C), and nutrient-stressed, without fertilizer, conditions (N). In the starting parental (S0) generation the effects of nutrient stress were not apparent until approximately one month after planting, and thus it was not surprising that controlled and nutrient deprived plants of the parental generation did not significantly differ in time to germination or time to development of first through fourth leaves (results not shown). *Lactuca serriola* plants in controlled conditions flowered significantly earlier than *L. sativa* in controlled conditions (Wilcoxon rank sum test, n=3, p-value = 0.0021). *Lactuca sativa* and *L. serriola* plants grown with nutrient stress flowered significantly later than plants grown in controlled conditions (Wilcoxon rank sum test, n = 3, *L. sativa* p-value = 0.0361 and *L. serriola* pvalue = 0.0361).

Lactuca sativa, but not *L. serriola*, S1 seedlings were significantly affected by the treatment of the parental generation. *Lactuca sativa* seedlings whose parents were nutrient deprived (n=11) had significantly lower above ground wet weight biomass (median = 0.227 g, median absolute deviation (mad) = 0.1586, Wilcoxon rank sum test, p-value = 0.0353, Figure 1A.) and fewer leaves (median = 4 g, mad = 0, p-value = 0.0035; Figure 1B.) than the offspring of non-stressed, controlled parents (n=19). However, these seedling differences associated with parental treatment did not translate to significant differences in time to flowering between the treatments groups (Figure 2).

Methylation signals in L. serriola and L. sativa in control and no-fertilizer conditions

To examine if methylation differences associated with growth in nutrient deprived conditions persisted to flowering in *L. sativa* and *L. serriola* we performed reduced representation bisulfite sequencing of two *L. sativa* and two *L. serriola* individuals which had been grown for two generations without fertilizer (NN). These generations and their parental generations were grown concurrently with the plants

sampled for WGBS sequencing and their parental generation [136]. The plants sampled for WGBS were grown in control conditions for two generations (CC) [136]. Hierarchical clustering using the methylation percentages at positions in NN and CC *L. sativa* and *L. serriola* biological replicates generated distinct, high confidence clusters by treatment and library type, with replicates of each genotype forming distinct sub-clades (Figure 3).

We identified DMCs between NN and CC samples of both *L. serriola* and *L. sativa*. In both *L. serriola* and *L. sativa* the median methylation percent across all DMCs was significantly higher in CC relative to NN conditions (Wilcoxon rank sum test, p-values « 0.001). For both *L. serriola* and *L. sativa*, the median methylation percentage of DMCs both within and upstream of protein coding genes was significantly higher in NN samples relative to CC controls (Wilcoxon rank sum test, p-values « 0.001), but significantly lower within annotated repetitive elements in NN samples relative to CC samples (Table 1). In each case the majority of DMC were found in the CHH context.

There were 667 positions which were differentially methylated in both NN *L*. *sativa* and NN *L*. *serriola* relative to their conspecific controls, these common DMC represent 28.9% of all unique DMC in these samples (Figure 4). The direction of difference in mean methylation percent between NN and CC samples was consistent for all of the shared DMC's, and 81% of the DMC had higher methylation levels in CC. Genotype-specific DMC represent a larger percentage of DMC within *L. sativa* (63%) than in *L. serriola* (43%). 48 of the common DMC were located within or upstream of 10 protein coding genes; seven of the genes encode proteins of unknown function, one is annotated as involved in ATP synthesis coupled proton transport, one as having protein serine/threonine activity, one as a structural constituent of ribosome (Table 4).

We identified 47,848 DMCs between *L. serriola* NN and *L. sativa* NN samples, over half (63.6%) of which had higher percent methylation in *L. sativa*. The median methylation percentage was significantly higher in *L. sativa* (80.6%) than in *L. serriola* (52%) (Wilcoxon rank sum test, p-values « 0.001) at the genome level and in all genomic regions analyzed including annotated repetitive elements (Table 1). Though there were many positions (19,843) that were significantly differentially variable between *L. sativa* and *L. serriola* under nutrient deprived conditions, only two of these positions were also differentially variable between *L. sativa* and *L. serriola* in control conditions [136].

Signals of trans-generational methylation in L. serriola under no-fertilizer conditions

We performed RRBS of *L. serriola* grown in the following two-generation stress treatments: samples grown in control conditions whose parents were grown in nutrient stress (NC); samples grown in nutrient stress conditions whose parents were also grown in nutrient stress conditions (NN); and samples grown in nutrient stress conditions whose parents were grown in control conditions (CN). WGBS sequencing was previously [136] generated for a fourth treatment in which both parents and offspring were grown under controlled conditions (CC). Three independent biological replicates were generated for each stress treatment. All libraries were aligned to the *L. serriola* genome sequence. The positions covered with at least 10 reads in all WGBS and RRBS samples were retained, returning information on 22,679, 18,585, 73,271 cytosines in the CG, CHG, and CHH contexts respectively. The median methylation percentage for cytosines in the CG context
were 86.96% (mad 11.87%), 94.7% (mad 7.86%), 94.62% (mad 7.97%), 95.12% (mad 7.23%) for plants grown in the CC, NC, CN and NN environments. The CC samples with no history of nutrient stress clustered distinctly from all other samples (Figure 5) though it is not possible with the current data to distinguish biological and technical variation due to the method of sequencing.

We identified DMCs between *L. serriola* NN, CN, and NC samples relative to CC samples. There was no genome-wide statistical difference in median percent methylation of DMCs within protein coding regions detected among the three treatments with a history of nutrient stress (NN, CN, or NC). However, the median methylation percentages showed significant genomic position dependent differences relative to controls. The median percentage methylation of DMC in protein coding genes within 1 kb of an annotated repetitive element was significantly lower in the samples with nutrient deprivation life histories than in controls, but methylation levels at DMC in protein coding genes more than 1 kb from an annotated repetitive element were significantly higher (Wilcoxon rank sum test, p<0.05, Table 2).

Within *L. serriola* samples which had themselves been subjected to stress treatment (CN and NN), 4,012 positions were differentially methylated relative to controls and 4,011 these positions had the same direction of change (hypo- or hypermethylation) relative to controls. These DMC represent 43.9% of all unique DMC in these samples (Figure 7 A.), and 52% of the common DMC had higher methylation levels in CC. 227 of the common DMC were located within or upstream of 57 protein coding genes of unknown function (List 1). Most (76.1%) of the DMC associated with current stress are also found in NC, and all of the DMC positions had the same direction of difference from controls (Figure 8). Approximately 6% of the common DMC were located within or upstream of 46 protein coding genes of unknown function (List 2). It is also notable that the number of DMC are positively correlated with the severity of stress in terms of immediacy and generational duration; relative to the CC treatment NN samples had the most DMC (7,377), followed by CN (5,765), and NC samples (5,212) (Figure 9).

For all contexts and conditions the coefficient of variation for methylation percentage at a position is inversely related to the average percent methylation, reflecting the importance of high and invariant methylation, possibly in silencing transposable elements (Figure 10). We utilized the iEVORA algorithm to test the null hypothesis of equal variances in the proportions of methylation between biological replicates of stressed and control L. serriola using a q-value threshold of 0.001 [104,147]. The methylation levels at differentially variable cytosines (DVC) tended to be high, more than 60% methylation, in all treatment vs. control contrasts (Figure 10 A.). Variability between biological replicates in stress treatments was lower than in controls (Figure 10 B.). However, in the current stress treatments (NN, CN) DVCs tended to be completely methylated and invariable in one of the two treatments (Figure 10). There were approximately equal number of DVCs which were more variable in the two current stress treatments (NN, CN) than control (186 DVCs), and less variable in both current stress treatments relative to untreated controls (203 DVCs), approximately 12 and 14% respectively of all unique DVCs found in both treatments. Methylation

levels at DVC positions in CN samples were more similar to controls and the coefficient of variation took on a range of more similar values (Figure 10 A.).

Discussion

The hypermethylation observed in *L. sativa* relative to *L. serriola* grown in controlled conditions [136] was maintained under stress conditions, *L. serriola* was hypomethylated relative to *L. sativa* when comparing between RRBS libraries of *L. sativa* and *L. serriola* grown without fertilizer for two generations (NN). The median level of methylation of DMC positions within NN *L. sativa* samples were consistently and significantly higher than in NN *L. serriola* samples in all genomic regions analyzed including annotated repetitive elements (Table 1). The methylation patterns of *L. sativa* and *L. serriola* NN and CC conditions were more similar by treatment and library type than by genotype (Figure 4).

In both *L. serriola* and *L. sativa*, DMCs in NN samples were globally hypomethylated relative to controls, though methylation levels of DMC differed significantly by genomic region. DMC within protein coding genes and upstream regions were significantly more methylated than in control samples, but significantly less methylated within annotated repetitive elements of NN samples relative to controls (Table 1). Several factors suggest that these significant patterns of hyperand hypomethylation could be beneficial, targeting higher rates of homologous recombination, and the associated higher rates of mutation, to less deleterious regions of the genome. Increases in homologous recombination frequency have been reported in plants subjected to abiotic stress and correlated with increased fitness in stressful environments [46]. Recombination hotspots are associated with high mutation frequency [148,149] and, in some plant genomes, enriched in LRR resistance genes [148]. Additionally, the frequency of crossing over events in a region is negatively correlated with its methylation levels as been shown in altered methylation patterning of *met1* mutants [150,151] and reduce recombination at recombination hotspots where constructs target methylation to these regions [150].

The parental generation's nutrient deprivation status resulted in significant differences in the S1 number of leaves and above ground biomass in *L. sativa* but not *L. serriola* (Figure 2). These gross phenotypic differences in the S1 based on parental treatment were not detectable at maturity in plants grown in control conditions, however, we found evidence that treatment specific methylation signals persisted in offspring of nutrient deprived parents grown in control conditions. In contrast, Secco et al. did not see persistence of methylation signals in offspring of inorganic phosphate deficient plants [152], suggesting a possible stress dose dependency of methylation persistence. This is supported by our comparative analysis of *L. serriola* with differing life histories of nutrient deprivation. The number of DMC detected relative to controls were positively correlated with the severity of stress in terms of immediacy and generational duration (Figure 9). This finding highlights the importance of considering the duration of the stress and the time between the application of the stress and when the tissue was sampled when comparing methylation studies.

In L. serriola we found apparent conservation of the positions and relative methylation difference of DMC in NN, CN, and NC samples relative to controls. There were 4,012 positions which were differentially methylated in both NN and CN L. serriola relative to controls, 43.9% of all DMC in these samples (Figure 7 A.). Most (76%) of these DMC were also differentially methylated in the S1 of stressed parents which had not themselves been subject to stress (NC) (Figure 8). Additional work is required to determine if this apparent conservation represents biological signal or a technical artifact due to comparing RRBS to WGBS libraries. Hierarchical clustering analysis including only the RRBS libraries does not distinguish between treatments (NN, CN, NC) suggests a high degree of relatedness between the biological replicates of different treatments (Figure 11). If confirmed the conservation of these stress associated sites even in samples which had not themselves been subject to stress implies the transgenerational transmission of stress associated differential methylation. Previous studies have suggested that inheritance of methylation may be inconsistent among siblings, though where the inherited methylation signals are present they have been associated with beneficial performance. Using low resolution methylation sensitive amplified polymorphism (MSAP), Kou et al. found offspring of nitrogen deprived plants which showed the altered mC pattern of their parents performed better in nitrogen deprivation than their siblings which did not inherit the modified mC pattern [153]. Likewise MSAP patterns in heavy metal stressed rice showed cases of transmission of the parental modification to the progeny as well as to the next selfed generation, and beneficial performance of offspring of stressed individuals with heavy

metal treatment relative to offspring of non-treated controls [154]. Beneficial effects of adaptive transgenerational priming has been reported to be reduced by treatment with methylation inhibitors [42,46], Dicer loss of function mutants [42,46] and with loss of function Pol IV, the RNA polymerase which produces transcripts from which sRNA are derived [42]. Interestingly, Boyko et al (2010) found offspring of salt stressed *Arabidopsis* were globally hypermethylated relative to offspring of non-stressed controls when grown in control conditions, but were hypomethylated relative to offspring of non-stressed parent when grown on salt [46]. The hypomethylation was also associated with better growth on salt media, but the beneficial effect of parental treatment was reduced when plants were treated with a methylation inhibitor [46]. A possible explanation could that the local hypomethylation is due to the activity of a DNA glycosylase such as ROS1 whose activity is positively regulated by DNA methylation [52]. This mode of action would be complementary with the hypothesis of sRNA of RdDM as the memory mechanism of transgenerational inheritance of modifications [155].

In addition to beneficial plant phenotypes associated with average methylation levels, diversity in methylation has been positively correlated with in plant productivity [75]. We previously found that most positions (86%) having significant differences in variability between *L. sativa* and *L. serriola* had more variable methylation percentages in the domestic variety *L. sativa* [136]. Similarly, increased methylation diversity was seen in domesticated soybean relative to wild soybeans [113]. It is interesting to note that one of the most significant positive measures of productivity observed previously associated with epigenetic diversity was increased plant density, a trait under positive selection in modern agriculture [156]. Here we see that when both genotypes were grown for two generations in nutrient deprived conditions we found the slight majority (59%) of DVCs were more variable among *L. serriola* replicates. Variability does not appear to be a conserved characteristic of particular genomic loci as the vast majority of positions which were differentially variable between the genotypes in the control conditions [136] were not differentially variable in the nutrient deprivation conditions. This shift in the relative variability of *L. sativa* and *L. serriola* under stressful conditions suggests the possibility that change in variability of methylation in response to changing environment could be a characteristic of stress adaptation.

Our work suggests several roles for acquisition and inheritance of methylation in the evolution of *Lactuca* sp. response to stress. Both genotypes exhibited patterns of hypermethylation within gene bodies and hypomethylation over repetitive elements in treatment conditions relative to conspecific controls, which suggests a beneficial role for stress associated methylation in targeting stress associated higher rates of homologous recombination, and the associated higher rates of mutation, to intergenic regions of the genome. We also found that changes in relative methylation levels at DMCs are less affected by environment than are changes in relative variability of methylation at DVCs. Though there were significant differences in methylation levels between *L. sativa* and *L. serriola*, in both treatment and controlled conditions most DMC were hypermethylated at DMC in *L. sativa* relative to *L. serriola* and there were a significant number of DMC in nutrient deprived conditions that were found in both genotypes, with the same direction of difference, relative to conspecific controls. In contrast, the frequency of DVC which were more variable in *L. sativa* relative to *L. serriola*, shifted between controlled and nutrient stressed conditions and there was very little overlap between DVC positions in the genotypes relative to controls. We found suggestions that abiotic stress associated methylation may be transmitted between generations with fidelity.

Figures

Figure 1. Development characteristics of *L. sativa* and *L. serriola* in one month old seedlings by parental treatment. The offspring of a parental generation grown in either nutrient deprived (N) or controlled (C) conditions had genotype dependent differences in wet weight (A.) and number of leaves (B.) as one month old seedlings. Dots represent individual seedlings and bars represent the median value per treatment.



Figure 2. Effect of parental stress treatment on days to flowering of next generation in *L*. *sativa* and *L*. *serriola* by parental treatment. Days to flowering for *L*. *sativa* and *L*. *serriola*, where parental and offspring generations were grown in control conditions (CC), parental generation in control conditions offspring generation in treatment (CN), parental generation in treatment conditions offspring generation in control conditions (NC), and parental and offspring generations were grown in treatment conditions (NN).



Figure 3. Hierarchical clustering of methylation of *L. sativa* and *L. serriola* grown in different treatment conditions. Shown are hierarchical clustering of methylation at positions having sufficient read support in CG (A), CHG (B), and CHH contexts. In red are approximately unbiased probability values generated by R package pvclust, in green are boot strap probabilities. C1 and C2 represent biological replicates grown in control conditions, N1 and N2 represent biological replicates grown in no-fertilizer conditions.



Figure 4. Venn diagram of DMCs found in NN *L. sativa* and NN *L. serriola* relative to their conspecific controls.



Figure 5. Hierarchical clustering of methylation at positions having sufficient read support in CG (A), CHG (B), and CHH (C) contexts. In red are approximately unbiased probability values generated by R package pvclust, in green are boot strap probabilities. CC1, CC2, and CC3 represent biological replicates grown in control conditions having parents also grown in control conditions, CN1, CN2, and CN3 represent biological replicates grown in control conditions, NC1, NC2, and NC3 represent biological replicates in control conditions whose parental plants were grown in no-fertilizer conditions, NN1, NN2, and NN3 represent biological replicates grown without fertilizer for two consecutive generations.



Figure 6. Total number of DMCs in *L. serriola* with different family histories of nutrient deprivation. The graph shows the total number of DMCs detected in *L. serriola* grown in nutrient deprived conditions for two generations (NN), *L. serriola* grown in nutrient deprived conditions and offspring of parents grown in controlled conditions (CN), or *L. serriola* grown in control conditions and *L. serriola* offspring of nutrient deprived parents (NC). 76% of the positions which were differentially methylated in both of the current stress treatments (NN, CN) are also differentially methylated in unstressed offspring of stressed parents (NC).



Figure 7. Relative methylation levels in nutrient deprived and control *L. sativa* and *L. serriola* plants. In both *L. sativa* (A.) and *L. serriola* (B.) DMCs in nutrient deprived individuals (NN) were hypomethylated relative to controls (CC).



Figure 8. Relative methylation levels in nutrient deprived *L. sativa* and *L. serriola* by genomic region. DMCs between NN *L. sativa* and NN *L. serriola* were consistently hypomethylated in *L. serriola*.





Figure 9. Relationship of the variance in methylation and mean methylation in *L. serriola* at cytosines covered by more than 10 reads in all samples.

Figure 10. Relationship of average methylation levels and variance in methylation in *L. serriola* with different family histories of nutrient deprivation. Average methylation levels (A) and variance in methylation (B) are shown for NC, CN and NN treatments in *L. serriola*.



Figure 11. Hierarchical clustering of methylation at positions having sufficient read support in CG (A), CHG (B), and CHH (C) contexts. CC1, CC2, and CC3 represent biological replicates grown in control conditions having parents also grown in control conditions, CN1, CN2, and CN3 represent biological replicates grown in without fertilizer and parental plants grown in control conditions, NC1, NC2, and NC3 represent biological replicates in control conditions whose parental plants were grown in no-fertilizer conditions, NN1, NN2, and NN3 represent biological replicates grown without fertilizer for two consecutive generations.



Tables

Table 1. Comparison of median methylation percentages over genomic regions of *L*. *sativa* and *L*. *serriola* in NN and CC conditions. Each comparison considers only positions with at least 10 reads in both samples.

	median, NN sativa	median, CC sativa	p- value	median, NN serriola	median, CC serriola	p- value	median, NN sativa	median, NN serriola	p- valu e
mRNA	24.23	1.15	0.00	30.46	3.06	0.00	40.75	24.07	0.06
mRNA, within 1 kb	21.43	0.73	0.00	NA	NA	NA	50.70	40.48	0.00
mRNA, beyond 1 kb	24.23	1.20	0.00	28.43	3.01	0.00	33.08	19.87	0.00
Upstream	26.53	1.58	0.00	42.30	3.34	0.00	46.05	25.76	0.00
Upstream, within 1 kb	21.43	0.73	0.00	NA	NA	NA	51.01	40.48	0.00
Upstream, beyond 1 kb	27.14	1.65	0.00	42.00	3.34	0.00	43.94	23.53	0.00
Repetitive Elements	7.18	65.62	0.00	5.99	60.00	0.00	57.69	44.61	0.00

	median,	median,	p-	median,	median,	p-	median,	median,	p-
	NN	CC	value	CN	CC	value	NC	CC	value
	serriola	serriola		serriola	serriola		serriola	serriola	
mRNA	29.31	38.93	0.92	31.97	40.09	0.52	30.95	46.39	0.02
mRNA, within 1 kb	31.04	46.39	0.03	35.53	46.84	0.00	34.12	49.77	0.00
mRNA, beyond 1 kb	21.86	2.26	0.00	25.00	2.92	0.00	22.47	6.56	0.03
Upstream	39.89	21.15	0.14	41.55	24.71	0.49	23.05	18.75	0.21
Upstream, within 1 kb	38.28	40.90	0.61	NA	NA	NA	27.16	59.15	0.70
Upstream, beyond 1 kb	40.54	19.55	0.12	41.92	24.66	0.46	23.05	18.57	0.25
Repetitive Elements	24.62	26.88	0.06	22.98	49.99	0.00	23.85	49.18	0.00

Table 2. Comparison of median methylation percentages over genomic regions of *L*. *serriola* in NN, CN and NC and CC conditions. Each comparison considers only positions with at least 10 reads in both samples.

Table 3. Comparison of median methylation percentages over genomic regions of *L*. *serriola* in different conditions, by proximity to annotated repetitive elements. Each comparison considers only positions with at least 10 reads in both samples.

	mRNA, within 1 kb	mRNA, beyond 1 kb	p-value	upstream, within 1 kb	upstream, beyond 1 kb	p-value
NN serriola						
(NNCCser)	31.04	21.86	0.29	38.28	40.54	0.67
CC serriola						
(NNCCser)	46.39	2.26	0.00	40.90	19.55	0.23
CN serriola						
(CNCCser)	35.53	25.00	0.28	15.79	41.92	0.38
CC serriola						
(CNCCser)	46.84	2.92	0.00	44.13	24.66	0.80
NC serriola						
(NCCCser)	34.12	22.47	0.73	27.16	23.05	0.97
CC serriola						
(NCCCser)	49.77	6.56	0.00	59.15	18.57	0.30

Table 4. Protein coding genes containing DMC within upstream or within gene bodies that were found in both NN *L. sativa* and NN *L. serriola* relative to their conspecific controls.

mRNA ID	Gene ontology terms				
Lsat_1_v5_gn_1_24401					
Lsat_1_v5_gn_3_17360					
Lsat_1_v5_gn_3_17381	GO:0015078: hydrogen ion transmembrane transporter activity: Molecular Function GO:0015986: ATP synthesis coupled proton transport: Biological Process GO:0015991: ATP hydrolysis coupled proton transport: Biological Process GO:0033177: proton-transporting two-sector ATPase complex				
Lsat 1 v5 gn 3 22600	GO:0003735: structural constituent of ribosome: Molecular Function GO:0005622: intracellular: Cellular Component GO:0005840: ribosome: Cellular Component				
Lsat 1 v5 gn 4 102681					
Lsat_1_v5_gn_4_145480					
Lsat 1 v5 gn 4 152881	GO:0004672: protein kinase activity: Molecular Function GO:0004674: protein serine/threonine kinase activity: Molecular Function GO:0005515: protein binding: Molecular Function GO:0005524: ATP binding: Molecular Function				
Lsat_1_v5_gn_4_63021					
Lsat_1_v5_gn_4_65980					
Lsat_1_v5_gn_6_37840					

Lists

List 1. Identifiers of 57 protein coding genes of unknown function which were associated with DMC found in *L. serriola* samples subjected to stress.

Lser_1_v1_gn_1_57860	Lser_1_v1_gn_8_46860	Lser_1_v1_gn_4_57800
Lser_1_v1_gn_1_67861	Lser_1_v1_gn_8_6600	Lser_1_v1_gn_4_71541
Lser_1_v1_gn_1_34840	Lser_1_v1_gn_8_58961	Lser_1_v1_gn_4_76341
Lser_1_v1_gn_2_160	Lser_1_v1_gn_8_65781	Lser_1_v1_gn_4_82241
Lser_1_v1_gn_2_36280	Lser_1_v1_gn_8_73660	Lser_1_v1_gn_4_98880
Lser_1_v1_gn_2_4440	Lser_1_v1_gn_8_83821	Lser_1_v1_gn_5_33080
Lser_1_v1_gn_2_5240	Lser_1_v1_gn_8_23320	Lser_1_v1_gn_5_64780
Lser_1_v1_gn_2_17381	Lser_1_v1_gn_8_30321	Lser_1_v1_gn_6_43921
Lser_1_v1_gn_3_35600	Lser_1_v1_gn_9_32460	Lser_1_v1_gn_6_45761
Lser_1_v1_gn_3_760	Lser_1_v1_gn_1_43960	Lser_1_v1_gn_6_27020
Lser_1_v1_gn_3_65721	Lser_1_v1_gn_1_59160	Lser_1_v1_gn_7_57641
Lser_1_v1_gn_3_9461	Lser_1_v1_gn_2_21861	Lser_1_v1_gn_7_23081
Lser_1_v1_gn_3_9481	Lser_1_v1_gn_2_5221	Lser_1_v1_gn_7_31001
Lser_1_v1_gn_4_43060	Lser_1_v1_gn_2_1800	Lser_1_v1_gn_8_44021
Lser_1_v1_gn_4_16001	Lser_1_v1_gn_2_13661	Lser_1_v1_gn_8_6001
Lser_1_v1_gn_5_40201	Lser_1_v1_gn_3_55720	Lser_1_v1_gn_8_6020
Lser_1_v1_gn_5_47480	Lser_1_v1_gn_3_63980	Lser_1_v1_gn_8_70301
Lser_1_v1_gn_7_38501	Lser_1_v1_gn_3_67420	Lser_1_v1_gn_8_10340
Lser_1_v1_gn_7_44920	Lser_1_v1_gn_3_1920	Lser_1_v1_gn_9_55720

List 2. Identifiers of 48 protein coding genes of unknown function which were associated with DMC found in all *L. serriola* samples either presently subjected to nutrient deprivation or whose parents were subjected to nutrient deprivation.

Lser_1_v1_gn_1_57860	Lser_1_v1_gn_7_44920	Lser_1_v1_gn_4_76341
Lser_1_v1_gn_1_67861	Lser_1_v1_gn_8_46860	Lser_1_v1_gn_4_98880
Lser_1_v1_gn_1_34840	Lser_1_v1_gn_8_58961	Lser_1_v1_gn_5_33080
Lser_1_v1_gn_2_160	Lser_1_v1_gn_8_83821	Lser_1_v1_gn_5_64780
Lser_1_v1_gn_2_36280	Lser_1_v1_gn_8_23320	Lser_1_v1_gn_6_43921
Lser_1_v1_gn_2_17381	Lser_1_v1_gn_8_30321	Lser_1_v1_gn_6_45761
Lser_1_v1_gn_3_35600	Lser_1_v1_gn_9_32460	Lser_1_v1_gn_6_27020
Lser_1_v1_gn_3_760	Lser_1_v1_gn_1_59160	Lser_1_v1_gn_7_57641
Lser_1_v1_gn_3_65721	Lser_1_v1_gn_2_21861	Lser_1_v1_gn_7_23081
Lser_1_v1_gn_3_9461	Lser_1_v1_gn_2_13661	Lser_1_v1_gn_7_31001
Lser_1_v1_gn_3_9481	Lser_1_v1_gn_3_55720	Lser_1_v1_gn_8_44021
Lser_1_v1_gn_4_43060	Lser_1_v1_gn_3_63980	Lser_1_v1_gn_8_70301
Lser_1_v1_gn_4_16001	Lser_1_v1_gn_3_67420	Lser_1_v1_gn_8_10340
Lser_1_v1_gn_5_40201	Lser_1_v1_gn_3_1920	Lser_1_v1_gn_9_55720
Lser_1_v1_gn_5_47480	Lser_1_v1_gn_4_57800	
Lser_1_v1_gn_7_38501	Lser_1_v1_gn_4_71541	

CHAPTER 5

CONCLUSION

The research presented in this dissertation increases our understanding of the relationship between DNA methylation in plants and environmental conditions through bisulfite sequencing of closely related accessions of *Lactuca*. We have shown that the methylomes of domesticated L. sativa and its conspecific wild and weedy relative, L. serriola, are very similar at the genome scale under non-stressed conditions. Both the domesticated and wild genotypes have genomic-region specific patterns of hypo- and hyper-methylation under stress conditions which may direct stress associated increases in homologous recombination away from gene coding regions. Both genotypes also have conserved methylation signatures around pathogen response related genes when grown in unstressed control conditions. The genotypes also showed environment specific differences in the relative abundance of differentially variable positions, suggesting genotype specific interactions between variability in DNA methylation and environmental stress. Together these findings suggest that epigenetic modifications are an additional source and mechanism of genomic variation which may be isolated and adapted for improvement of crops.

Relationship of global methylation levels and gene regions

Though both *L. sativa* and *L. serriola* both have very high average levels of methylation, similar to other large plant genomes, a significant number of DMCs are more highly methylated in *L. sativa* than *L. serriola*. The differences between this self-crossing dicot wild-domestic pair do not support an obvious relationship between methylation levels and a plant's domestication status that transcends diverse plant families. Methylation levels between closely related wild and domesticated monocots are not correlated with domestication status [54] and methylation levels at DMCs between domesticated corn and wild teosinte tend to be less methylated in the domestic varieties [157].

Within the methylomes of *L. sativa* and *L. serriola* are interesting characteristics suggesting a role for DNA methylation in plant-microbe interactions. We found striking patterns of methylation around resistance genes, highly conserved methylation states in pathogen response related genes, and gene ontology enrichment of plant-microbe interaction related terms among genes with differentially methylated cytosines between *L. sativa* and *L. serriola*. DMCs that distinguish these genotypes are related to important gene ontology categories known to affect fitness. DMCs between *L. sativa* and *L. serriola* were found in genes enriched for gene ontology terms related to: photosynthesis, biosynthesis of aromatic amino acids, signaling and lipid signaling, and transport; these terms are also implicated in plant microbe interactions [158–160].

Methylation levels around protein coding genes in *Lactuca* sp. have characteristic patterns shared by most angiosperms and conserved and striking methylation patterns

around genes related to pathogen response. Like most other angiosperms, methylation levels upstream and downstream of protein coding genes in *Lactuca* sp. are relatively high, dipping dramatically at transcription start and end sites. However, the average levels of methylation in *Lactuca* sp. take on strikingly different patterns around annotated disease resistance genes. The up- and downstream regions of resistance genes have much higher average levels of methylation in the CHH context than the genome wide average across all predicted protein coding genes. The accumulation of CHH methylation is particularly noteworthy in the regions 100-400 bp upstream of the transcription start site. Small RNAs were found to accumulated in these regions of resistance genes in Arabidopsis, and the active demethylation of these regions by ROS1 upon pathogen challenge was shown to be required for pathogen resistance [41]. As the average levels of methylation over these regions do not vary between unchallenged L. sativa and L. *serriola*, genotypes with divergent pathogen resistance phenotypes, it would be interesting to compare the relative rates of active methylation and demethylation around resistance genes in *L. sativa* and *L. serriola* during the course of active pathogen infection. Previous work profiling the DNA methylation of salt resistant and tolerant rice genotypes found that rates of change in methylation, but not average methylation levels, differed between salt tolerant and sensitive rice varieties [45].

An additional distinctive feature of *Lactuca* sp. resistance genes is the strong spike in methylation levels in all sequence contexts found at the 3' end of the genes. Similar spikes in methylation had previously been seen in Arabidopsis resistance gene RPP7 [161]. In RPP7 the 3' methylation was associated with an intronic transposable

element, accumulation of Histone H3 lysine9 di-methylation (H3K9me2) and alternative polyadenylation of the gene [161]. Sequence analysis of the 3' intronic regions of these genes in *Lactuca* sp. could determine if similar transposon "domestication" explains the observed accumulation of 3' methylation in lettuce. Likewise, transcripts of these genes could be analyzed for isoform production and temporal changes in methylation levels under pathogen challenge. Though the *L. sativa* and *L. serriola* average levels of methylation around resistance genes do not significantly differ, analysis of differential rates of methylation and demethylation of resistance genes and flanking regions could highlight differences in pathogen responsiveness which could complement traditional gene based crop improvement strategies.

The relationship between average methylation levels and the variability of methylation levels between biological replicates was highly dependent on genomic region. At most genomic positions average methylation levels are inversely related to the variability of methylation at that position. Sites of highly conserved methylation were defined as being among the 25% least variable positions between biological replicates of both *L. sativa* and *L. serriola*. Genes with highly conserved methylation states were highly methylated, even though, genome wide, most positions with highly conserved states had low or no methylation. This seemingly contrary result was explained by looking at the frequency of occurrence of conserved methylation states by genomic region. The majority of positions with highly conserved methylation in all sequence contexts. Most of the conserved positions found in annotated repetitive regions

were in the CG and CHG context and had high levels of methylation in line with genome wide averages for these features. Genes containing positions with highly conserved methylation states had unusually high levels of methylation (>90% methylation in all sequence contexts), particularly notable in the CHH context, suggesting that these sites may be targets of RdDM. Annotated genes with the highest frequency of positions with highly conserved methylation states controlling for variation in gene length included endo-1,4-beta-mannosidase, a cell wall degrading enzyme, Flagellin-sensitive 2 (FLS2), an important sensor of pathogen attack, and F-type H+-transporting ATPase subunit C, a conserved transport ATPase. The frequency of conserved methylation states within FLS2 is particularly interesting given previous work correlating expression of FLS2 in ROS1 dependent DNA de-methylation [41]. FLS2 is strongly induced by the bacterial flagellin N-terminal epitope flg22, and triggers transcriptional regulation of stress response genes. Flg22 exposure also results in ROS1 dependent DNA de-methylation and expression of a specific subset of transposable elements [41]. Given the highly conserved and highly methylated state of FLS2 in non-pathogen challenged *Lactuca* sp. and the interaction of FLS2 with de-methylation dependent pathogen resistance – it could be illuminating to assess the methylation in this gene and the class of R-genes targeted by ROS1 over the course of pathogen infection.

The effect of altered methylation levels on plant-microbe interactions and microbial community composition warrants further research. The work to date has focused on the interaction of particular loss of function DNA methylation mutants on the plants' susceptibility to particular pathogens [40,41,62,69,84] or the plant methylation

signature of genotype specific interactions with a particular beneficial microbe [162]. The natural milieu in which the microbes and plant communicates have evolved is much more complex than the introduction of a single beneficial or pathogenic bacteria or fungi in a controlled environment. The plant microbiome can be an important contributor to plant fitness [163] and may be an important link in the interaction of epigenetic diversity of plant populations and their environments with fitness traits such as biomass density, competition and pathogen resistance [75].

Acquisition and variability of mC in *L. sativa* and *L. serriola* in nutrient limited conditions

Methylation levels in both nutrient deprived and control samples of *L. sativa* were higher than the corresponding treatment in *L. serriola*. Though *L. sativa* and *L. serriola* differed in methylation levels, there were consistent patterns of methylation between the two conditions. More than half of the DMCs in stressed *L. serriola* relative to controls were also found in *L. sativa* relative to controls with the same direction of difference relative to controls. In both *Lactuca* sp. the plants grown in nutrient limited conditions were hypermethylated in gene bodies and hypomethylated over repetitive elements relative to control plants. In both biotic and abiotic stress response, differential methylation has been associated with genes and upstream regions that are in close proximity to repetitive elements [40,70]. A similar general role for TE associated DNA methylation of nearby genes is found in *Lactuca* as the methylation levels of genes in both *L. sativa* and *L. serriola* differ based on their proximity to annotated repetitive

elements. Methylation levels are significantly higher in genes located within 1000 bp of an annotated repetitive element, possibly due to TE targeted methylation which may introduce selectable variation in gene expression.

We found a significant correspondence between positions of stress associated differences in mean methylation levels between genotypes, but a significant shift in the positions and relative abundance of differentially variable cytosines between the two genotypes under stress conditions. Growth in nutrient limited conditions resulted in accumulation of more DMCs within L. sativa relative to con-specific controls than L. serriola, suggesting the domestic methylome was more affected by growth in nutrient limited conditions. Though the total number of positions differed dramatically, more than half of the DMC seen in stressed L. serviola relative to controls are also seen in L. sativa relative to controls with the same direction of difference relative to controls. In contrast, the relative variability at sites of differentially variable methylation shifted between the two genotypes in control and nutrient deprived treatments. Most DVCs were more variable in L. sativa than L. serriola under control condition but were less variable in L. sativa under nutrient limited conditions. The relative shift toward more variability under nutrient limited conditions in *L. serriola* is an interesting finding in light of the relative adaptation of L. serriola to marginalized and highly disturbed environments. Recent work identified an association of increased diversity of methylation with increased plant productivity in pathogen and competition challenged environments [75]. The drivers of differential variability are unclear, though variability does not appear to be characteristic of particular genomic regions as sites of differentially variable methylation between

biological replicates of *L. sativa* and *L. serriola* in control conditions were conserved in nutrient limited conditions. It would be interesting to study the relative activity and fidelity of the methyltransferases in the two genotypes to further dissect possible sources of variability.

Conclusions

This work identified several ways in which differential methylation levels and conservation of methylation states between these two genotypes are associated with gene regions and functions implicated in plant-microbe interactions. Thought there were particular positions having significantly different average methylation levels, there were also positions in genes related to plant microbe interactions whose methylation states were highly conserved. These instances suggest the value of closely monitoring these loci over the time course of acute stress, particularly over the time course of pathogen exposure and infection. This work also identified an interaction between stress environment and the relative frequency of differentially variable cytosines between the genotypes. This work allows future researchers to be more targeted in their approach to dissect the role of DNA methylation in the stress adaptive phenotypes of these two *Lactuca* sp.. Research may be targeted to key genes and genomic regions involved in pathogen response and more efficiently analyze the time course component of acquired DNA methylation modifications. Additionally the increase in relative abundance of highly variable positions in stress conditions within the relatively stress-adapted genotype, is in line within initial research in Arabidopsis suggesting a positive adaptive

role for increased epigenetic diversity in the absence of genetic diversity [75]. Competition experiments between populations of differing levels of epigenetic diversity within genetically homogeneous populations of *L. sativa* and *L. serriola* would help distinguish the relative contribution of epigenetic diversity and genotype to the stressadaptation of the wild species.

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