Loss of Gene Body Methylation in *Eutrema salsugineum* Is Associated with Reduced Gene Expression

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Abstract

Gene body methylation (gbM) is typically characterized by DNA methylation in the CG context within coding regions and is associated with constitutive genes that have moderate to high expression levels. A recent study discovered the loss of gbM in two plant species (*Eutrema salsugineum* and *Conringia planisiliqua*), illustrating that gbM is not necessary for survival and reproduction. The same paper stated there was no detectable effect of gbM loss on gene expression (GE). Here, we reanalyzed the GE data and accounted for experimental variability in expression level estimates. We show that the loss of gbM in *E. salsugineum* is associated with a small but highly significant decrease in GE relative to the closely related species *Arabidopsis thaliana*. Our results are consistent with various evolutionary analyses that suggest gbM has a function, perhaps as a homeostatic effect on GE.

Key words: gene body methylation, Brassicaceae, gene expression.

Since its first genomic characterization in *Arabidopsis thaliana* (Cokus et al. 2008; Lister et al. 2008), gene body methylation (hereafter gbM) has been a puzzling phenomenon. Typically, plants methylate cytosines in three contexts: CG, CHG, and CHH (where H = A, C, or T). When all three contexts are methylated, gene expression (hereafter GE) is silenced. In contrast, *A. thaliana* gbM is characterized by elevated methylation levels in the CG context, but neither in the CHG nor in the CHH context of coding regions. This high CG methylation does not silence genes. In fact, gbM genes tend to be immediately expressed (Zhang et al. 2006; Lister et al. 2008) and expressed across more tissues than Lmethylated (UM) genes (Lister et al. 2008; Kawakatsu et al. 2016). Importantly, not all genes have gbM; only 20% of *A. thaliana* genes contain CG methylation above background levels (Takuno and Gaut 2012).

The puzzling aspect about gbM is its function. Some argue that gbM must have some function (Zilberman 2017), based on four observations rooted in evolutionary analyses. First, gbM levels are highly conserved between orthologs (Takuno and Gaut 2013; Seymour et al. 2014), even across ~300 My of land plant evolution (Takuno et al. 2016). Second, gbM genes tend to evolve more slowly than UM genes (Takuno and Gaut 2012, 2013) with lower levels of polymorphism (Takuno et al. 2017), consistent with their enrichment for important functions (Zhang et al. 2006; Takuno and Gaut 2012). Third, most gbM genes are not particularly GC rich, suggesting that elevated cg methylation levels are not a simple consequence of cytosine availability (Takuno and Gaut 2013). In fact, gbM is maintained against mutational biases that decrease GC content over time (Takuno and Gaut 2013). Finally, gbM status seems to have a modest effect on GE, based on an evolutionary comparison between closely related *Arabidopsis* species (Takuno et al. 2017). This comparison focused on a small subset of orthologs that did not have conserved gbM levels between species. It revealed that gbM genes tend to be more highly expressed than their UM orthologous counterpart, but the overall trend was not convincingly significant. In another study comparing *A. thaliana* Swedish accessions, gbM genes were found to be more heavily methylated in northern latitudes, which was associated with a higher expression level (Dubin et al. 2015).

Other studies have suggested gbM has no function and is instead a by-product of transposition and/or methylation pathways (Roudier et al. 2009; Teixeira and Colot 2009; Kawakatsu et al. 2016; Bewick and Schmitz 2017). This argument is consistent with observations that: 1) the loss of gbM in an *A. thaliana* mutant does not substantially alter GE (Roudier et al. 2009; Bewick et al. 2016) and 2) gbM variation across *A. thaliana* accessions does not strongly affect GE (Kawakatsu et al. 2016). Two recent papers have seemingly strengthened the argument against gbM functionality, because they found that two flowering plants (*Eutrema salsugineum* and *Conringia planisiliqua*) have no gbM throughout their genome. The identification of these species is important for two reasons. First, they have provided important insights into the mechanism that produces gbM, which had been mysterious (Bewick et al. 2017). Second, they provide *prima facie* evidence against gbM function, because these two plant species seem to exist happily without it. In support of this argument, Bewick et al. (2016) compared GE between genes that have gbM in *A. thaliana* and their UM orthologs in *E. salsugineum*. They concluded that the two sets of genes had similar transcription levels, again suggesting that gbM has no function.

Here, we use the data of Bewick et al. (2016) to revisit GE analyses and to test whether an effect of gbM can be
detected. We began by defining gbM genes and UM genes in *A. thaliana* (see Materials and Methods) using the same statistical approach and thresholds as Bewick et al. (2016). We then identified 12,189 orthologous UM genes in *E. salsugineum*, based on best-hits between species, again using the same methods as Bewick et al. (2016). These ortholog pairs were then separated into two groups (table 1) for inclusion in GE analyses. Group 1 consists of 4,221 ortholog pairs that changed gbM status between species—that is, the gene was gbM in *A. thaliana* and UM in *E. salsugineum*. Group 2 includes 7,968 ortholog pairs that did not change gbM status, because they were UM in both species. These Group 2 genes can be viewed as a “control” set and are necessary for comparative analyses.

<table>
<thead>
<tr>
<th>Ortholog Pairs</th>
<th>Methylation Status in <em>A. thaliana</em></th>
<th>Methylation Status in <em>E. salsugineum</em></th>
<th>Change in Methylation Status</th>
<th>Number of Orthologous Genes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group 1 gbM</td>
<td>UM</td>
<td>Yes</td>
<td>4,221</td>
<td></td>
</tr>
<tr>
<td>Group 2 UM</td>
<td>UM</td>
<td>No</td>
<td>7,968</td>
<td></td>
</tr>
</tbody>
</table>

Note.—The first group consists of genes that have changed methylation status between *A. thaliana* and *E. salsugineum*. The second group has conserved methylation status as UM genes.

Our result is important for at least two reasons. First, by utilizing *E. salsugineum*, a species naturally devoid of gbM, we have studied the effect of gbM loss from 4,221 Group 1 genes simultaneously. A previous paper took a similar approach but could study only hundreds of genes (Takuno et al. 2017). It is worth noting, however, that both studies have detected a modest but consistent association between gbM and GE, with a trend toward higher expression levels for gbM genes. Second, by contrasting the GE effects between two groups—that is, genes that did and did not differ in gbM status (table 1)—we have effectively controlled for experimental effects. We suspect that our conclusions differ from Bewick et al. (2016) in part due to the use of Group 2 as a control to contrast to Group 1 genes.

We believe these results to bolster the argument, which is largely based on evolutionary analyses, that gbM has some function, specifically with regard to effects on GE. Although the loss of gbM in *E. salsugineum* is associated with only a small reduction in GE, the effect is significant and notable across 4,221 genes, even if the effect is small at the scale of individual genes. We note that natural selection can act on functional genomic features that have small effects on fitness, such as codon usage bias (Duret and Mouchiroud 1999), as long as the species effective population size (*N_e*) is large enough for selection to act efficaciously. One reason why *E. salsugineum* lost gbM, whereas most angiosperms have preserved it, could be the relatively small *N_e* of this species and its specialized halophytic niche (Wang et al. 2018).

The question remains, however, as to the exact nature and mechanism linking gbM and GE. Zilberman (2017) argues persuasively that the primary effect of gbM is likely to be homeostatic, by (for example) preventing aberrant transcription within genes or restricting access to histone H2A.7, which is associated with gene responsiveness. If the effect of gbM is to better regulate GE, it may have been missed up to now due to technical limitations. Indeed, current estimates of GE are averaged across thousands of cells, and because the effect of gbM is small, it may be necessary to consider single cell approaches to better understand it (Ji et al. 2015; Zilberman 2017). Whatever the function, the effect of gbM is modest enough to be difficult to detect on experimental time-scales, but strong enough to be conserved among plant orthologs.

### Materials and Methods

#### Data Set
We used data from Bewick et al. (2016) and Niederhuth et al. (2016) from five Brassicaceae species: *A. thaliana*, *Arabidopsis lyrata*, *Brassica rapa*, *Capsella rubella*, and *E. salsugineum*. The second group has conserved methylation status as UM genes.
The methylation data consisted of the methylation status of each cytosine based on a binomial test (Lister et al. 2008). The RNA-seq data were reported by Bewick et al. (2016) as the FPKM level for each gene for six *A. thaliana* replicates (three leaf and three aerial tissue) and two *E. salsugineum* replicates from leaf tissue (supplementary table S1, Supplementary Material online).

**gbM Inference**

For each gene, the methylation state was inferred using the same approach as Bewick et al. (2016) and Niederhuth et al. (2016), for details see Supplementary Material.

**Analysis of Expression and Methylation Levels**

GE levels were analyzed using a linear regression model with mixed effects using the R package *lme4* (Bates et al. 2015). FPKM expression levels were log transformed. To account for interGE variability, a random gene effect was included in the model (see eq. 1). Similarly, a random tissue effect was included (eq. 1) to account for variability in expression between leaf and aerial RNA-seq data. The aim of this model was to compare the expression level of gbM genes in *A. thaliana* to genes that lost gbM in *E. salsugineum*, while taking into account any global difference in expression levels between the two species. In order to achieve this, we defined a fixed effect called "gene Group" (table 1). A species fixed effect was also used in the model to account for global differences in expression levels between the two species. The resulting model was written to examine an effect of gene Group on expression level after taking the species effect into account—that is,

\[
\log(FPKM + 1) \sim \text{Species} \ast \text{gene Group} \\
+ (1|\text{Tissue}) + (1|\text{Gene}) 
\]

(1)

Significance for fixed effects and their interaction were determined by comparing the fit of the full model to nested models that first removed the interaction and then removed one effect at a time. *P*-values for each effect and their interaction were determined by comparing the fit of the full model to nested models that first removed the interaction and then removed one effect at a time.
interaction were computed via Wald-statistics approximation using sjPlot R package (Lüdecke 2018).

**Supplementary Material**

Supplementary data are available at *Molecular Biology and Evolution* online.

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**References**


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