Evidence for Widespread Genomic Methylation in the Migratory Locust, *Locusta migratoria* (Orthoptera: Acrididae)

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Abstract

The importance of DNA methylation in mammalian and plant systems is well established. In recent years there has been renewed interest in DNA methylation in insects. Accumulating evidence, both from mammals and insects, points towards an emerging role for DNA methylation in the regulation of phenotypic plasticity. The migratory locust (*Locusta migratoria*) is a model organism for the study of phenotypic plasticity. Despite this, there is little information available about the degree to which the genome is methylated in this species and genes encoding methylation machinery have not been previously identified. We therefore undertook an initial investigation to establish the presence of a functional DNA methylation system in *L. migratoria*. We found that the migratory locust possesses genes that putatively encode methylation machinery (DNA methyltransferases and a methyl-binding domain protein) and exhibits genomic methylation, some of which appears to be localised to repetitive regions of the genome. We have also identified a distinct group of genes within the *L. migratoria* genome that appear to have been historically methylated and show some possible functional differentiation. These results will facilitate more detailed research into the functional significance of DNA methylation in locusts.

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Introduction

DNA methylation is a taxonomically widespread epigenetic marker to which many non-exclusive functions have been attributed, among them genomic imprinting and the regulation of phenotypic plasticity [1–5]. It was originally considered doubtful that DNA methylation was biologically significant in insects, based upon the results of initial studies using *Drosophila melanogaster* adults [6]. Subsequent work confirmed the existence of methylecytosine in dipteran insects [7], but showed that it is present in very small amounts and, at least in *D. melanogaster*, limited to early embryonic developmental stages [8]. Similarly low levels of DNA methylation were found in the coleopteran *Tribolium castaneum* [9,10]. More recently, however, a growing body of research has indicated that in a number of insect species DNA methylation is both present at appreciable levels and regulates diverse and important biological processes. In honeybees (*Apis mellifera*) it is involved in caste differentiation [4] and learning and memory [11]. In mealybugs (*Planococcus citri*) it is thought to be responsible for genomic imprinting [2] and in the peach aphid (*Myzus persicae*) has been associated with pesticide resistance [12–14].

In contrast to the high levels of global methylation characteristic of vertebrates, it appears that DNA methylation in insects has a mosaic distribution throughout the genome, occurs predominantly within genes, and does not play a large role in the suppression of repetitive DNA [9,15]. Accordingly, the proportion of the genome that is methylated is typically reported to be much smaller in insects [1,16–18]. However, there are some examples of insect species with relatively higher levels of DNA methylation [19] and in one case this appears to be associated with the methylation of repetitive DNA [20]. Overall, DNA methylation in insects is still not well characterised, with the potential diversity in its functional roles remaining largely unexplored. It is also unclear how much variation exists across taxa in the amount and distribution of DNA methylation within the genome and its significance to organismal biology.

The migratory locust (*Locusta migratoria*) is an economically important species of insect with a globally widespread distribution [21]. Individuals display a striking polyphenism, exhibiting either a solitary or gregarious phase phenotype depending upon maternal epigenetic cues [22–24] and the population density that they have experienced [25,26]. The expression of these density-dependent phenotypic changes in behaviour and other traits can play a causal role in locust swarm formation [27–29]. Phase polyphenism is generally accepted to have arisen independently multiple times within locusts [30] and it is unclear to what extent the underlying molecular mechanisms are conserved between species. Until very recently, there was only one report of DNA methylation in locusts, with estimates of methyleycytosine content obtained using a rudimentary paper chromatography technique [31]. This is somewhat surprising given the significance of these species as models for the study of phenotypic plasticity. The presence of DNA
methylation and DNA methyltransferase encoding genes has now been confirmed in the desert locust Schistocerca gregaria [32], but the function of DNA methylation in locusts is yet to be determined. Here we provide novel evidence that the genome of the migratory locust is both methylated and contains genes encoding methylation machinery orthologs. We show that a fraction of the genome enriched in methyl-CpG sites contains repetitive sequences, including a SINE retrotransposon known to be differentially expressed between isolated and crowded locusts [33]. We also demonstrate that a subset of locust genes display evidence of historical methylation within their protein encoding sequences. The degree of both current and historical genomic methylation indicated by our results to be present in the migratory locust suggests the possible role of DNA methylation in regulating locust phenotypic plasticity as a promising avenue of future study.

Results

Methylation-specific restriction enzyme assays

Digestion of both *L. migratoria* egg and adult head DNA with *Hpa*II (5’-CCGG-3’), which is sensitive to methylation of the internal cytosine, revealed a pronounced high molecular weight fraction that largely disappeared in the methylation-insensitive *Msp*II (5’-CCGG-3’) digest, clearly indicating the presence of methylated CpG sites within this DNA (Figure 1a, b). In contrast, no distinction was evident between the *Hpa*II and *Msp*II digests of either *D. melanogaster*, the adult genome of which contains virtually no methylcytosine, or the honeybee, which possesses a functional CpG methylation system, but relatively low levels of genomic methylation (Figure 1a). A much larger proportion of the *Mus musculus* (mouse) DNA was observed to be resistant to *Hpa*II digestion than the insect DNA, consistent with the relatively high levels of CpG methylation observed in mammals. The majority of the high molecular weight fraction was cleaved by *Msp*I, and in this species, unlike the insect representatives, there was also a noticeable reduction in the average fragment length obtained from the *Msp*I digestion relative to the *Hpa*II digest. No obvious differences in CpG methylation were detected between the genomic DNA of eggs from isolated and crowded locusts using this method (Figure 1b). The restriction pattern of genomic DNA from crowded adult head tissue was comparable to that of egg tissue (Figure 1a, b).

Cloning and sequencing of methylcytosine enriched genomic DNA

The high molecular weight fraction of *L. migratoria* genomic DNA seen following digestion with *Hpa*II was not present for the *Msp*I digest, which indicates that this DNA is enriched in methylated CpG sites. In an effort to further characterise this portion of the genome, we purified the DNA and then cloned and sequenced a number of fragments. Omitting bacterial contaminants, a total of 49 clones were obtained, representing 40 unique sequences (Table 1). One third of these clones (31%) were found to be resistant to *Hpa*II digestion and thus contain DNA methylated by the *Hpa*II restriction enzyme, which is sensitive to methylation of the internal cytosine. The *Hpa*II digestion behaviour of these DNA sequences is comparable to that of methyl-CpG enriched DNA from a mouse (Figure 1a, b). Among the methyl-CpG enriched DNA samples, we have identified a number of locust sequences, a total of 49 clones were obtained, representing 40 unique sequences (Table 1). One third of these clones (31%) were found to be resistant to *Hpa*II digestion and thus contain DNA methylated by the *Hpa*II restriction enzyme, which is sensitive to methylation of the internal cytosine. The *Hpa*II digestion behaviour of these DNA sequences is comparable to that of methyl-CpG enriched DNA from a mouse (Figure 1a, b).

Table 1. Identity of cloned DNA sequences derived from methyl-CpG enriched portion of the *L. migratoria* genome.

| Sequence type          | Sequence class          | Number of clones (%) |
|------------------------|-------------------------|----------------------|
| Repetitive DNA         | 15 (31%)                |
| *Lm* retrotransposon   | 8 (16%)                 |
| Other transposon       | 3 (6%)                  |
| Tandem repeat          | 2 (4%)                  |
| Other                  | 2 (4%)                  |
| Unannotated locust EST | 9 (18%)                 |
| Unknown sequence       | 25 (51%)                |

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to undergo CpG depletion over evolutionary time [34,35]. The ratio of observed to expected CpG sites (CpG*O/E*) in a sequence can therefore be used to make inferences about the level of historic DNA methylation in regulating locust phenotypic plasticity as a promising avenue of future study.

Observed and expected CpG ratios

Due to the hypermutability of methylcytosine to thymine, sequences that are highly methylated in the germline are predicted to undergo CpG depletion over evolutionary time [34,35]. The ratio of observed to expected CpG sites (CpG*O/E*) in a sequence can therefore be used to make inferences about the level of historic DNA methylation in regulating locust phenotypic plasticity as a promising avenue of future study.

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DNA methylation that occurred in the germline. As no genome sequence information was available for *L. migratoria* at the time of the present study, analysis of CpG sites was performed using a data set of 12,148 unigenes downloaded from a publicly available EST database [36]. The distribution of unigene CpGΩ/E ratios is clearly bimodal (Figure 2; unimodal distribution rejected, p<0.05), with two peaks centred at approximately 0.4 and 0.9. This suggests that the *L. migratoria* genome contains two distinct populations of genes, those that have an observed CpG frequency close to that expected by chance and those in which the CpG content is much lower. A similar distribution of CpGΩ/E ratios was observed when 345 unigenes previously demonstrated to be differentially expressed between isolated and crowded *L. migratoria* [37] were examined separately (Figure 2b). This pattern was observed regardless of whether genes were more highly expressed in isolated or crowded locusts. Whereas CpGΩ/E shows a pronounced bimodal distribution, CpG¶Ω/E and all of the other dinucleotide pairs display unimodal gene frequencies for the complete unigene dataset (Figure S1). This implies that there has been little bias exerted upon the CpGΩ/E distribution by the GC content of the genes and that the bimodality of CpGΩ/E is likely to be attributable to the historic presence of methylcytosine in the germline. The existence of the low CpGΩ/E population of unigenes suggests that widespread historical gene body methylation has occurred in a CpG context in this species, and that this extends to some of the genes involved in regulating phase polyphenism. Although there is strong evidence of CpG depletion and historic methylation within the *L. migratoria* genome, caution should be exercised when interpreting precisely to what extent this occurs, as it is possible that the distribution of CpGΩ/E ratios has been influenced by positional biases in the distribution of ESTs within transcripts and methylated CpG sites within gene bodies [9]. Gene Ontology (GO) functional enrichment analyses (biological process) of a subset of 2,482 unigenes for which annotation data was available showed that transport, carbohydrate metabolism, proteolysis and peptidolysis, chitin metabolism and muscle development genes were overrepresented in the high CpGΩ/E group (p<0.05), whereas metabolism genes were significantly enriched in the low CpGΩ/E class of genes (Table 2).

### Identification of *L. migratoria* methylation machinery

DNA sequence data was obtained for two of the three animal methyltransferases, the DNA methyltransferase (Dmnt1) and the tRNA methyltransferase (Dmnt2) as well as the invertebrate methyl-CpG binding domain protein (MBD2/3). The corresponding amino acid sequences contained conserved motifs characteristic of these proteins (Figure 3). Two discrete *L. migratoria* Dmnt1 cDNA variants, designated Dmnt1 type 1 and type 2 and sharing 96.9% identity at the amino acid level (340/351 sites) were identified from two clones. Bayesian analysis of Dmnt1 sequences grouped all of the arthropod representatives together (Figure 3). Dmnt1a and Dmnt1b sequences were not found to cluster according to gene class; for insect species in which two Dmnt1 genes have been identified, greatest similarity was found between the two Dmnt1 variants within a species.

### Discussion

The present study provides the first evidence that the genome of *L. migratoria* contains genes encoding DNA methylation machinery, and demonstrates using two independent lines of evidence that the genome is methylated to a considerable degree. Differences in the pattern of restriction fragments observed following digestion of *L. migratoria* DNA with *HpaII/MspI*, which is indicative of the level of CpG methylation, were not as pronounced as for *M. musculus*, which exemplifies the high level of genomic methylation characteristic of mammals. The amount of cytosine methylation present for *L. migratoria* appears to be higher than that of *D. melanogaster* and *A. mellifera*, the latter of which has a functional DNA methylation system [10], although it is possible that this is partly due to variation in the distribution of *HpaII/MspI* sites within the genome as well as differences in the amount of CpG methylation. The proportion of methylcytosine in the genome of another locust, *S. gregaria*, has recently been estimated to be 1.3–1.9% in different nervous and endocrine tissues [32]. In agreement with the *L. migratoria* digest results, these values are intermediate to those calculated for *D. melanogaster* embryos and *A. mellifera* (0.1% and 0.3% respectively) [9] and mammals (4–6%) [38]. Digestion of *L. migratoria* DNA was similar to that reported for the stick insect *Medauroides extradentata* [20] and the mole cricket *Gryllotalpa fuscus* [39] and is reminiscent of that seen in higher deuterostome invertebrates [40]. Together with the finding that the cabbage moth (*Mamestra brassicae*) possesses relatively high levels of DNA methylation [19], these results highlight the variability in the amount of methylation present among insect genomes, the majority of which have been shown to be sparsely methylated to date.

Genome-wide analyses of methylation have been conducted at base pair resolution in two insects, the honeybee and the silk moth (*Bombyx mori*). In these species it has been shown that DNA methylation is localised primarily to gene bodies and at least in *B. mori*, sequences encoding small RNAs, whereas transposable elements are hypomethylated [9,16,41]. It appears that while there is evidence of gene body methylation in *L. migratoria*, CpG methylation is not limited to the regions described for *A. mellifera* and *B. mori*. As for the phasmid *M. extradentata* [20], substantial methylation of repetitive sequences is also likely to be present, as indicated by the 31% of clones from the methyl-CpG enriched portion of the *L. migratoria* genome found to contain repetitive DNA. Recent studies have indicated that DNA methylation is involved in stabilising gene expression [35] and regulating the production of alternative transcripts [42]. Despite these insights, the role of DNA methylation and its functional conservation throughout different taxa is still not fully understood. It is possible that in the Orthoptera and Phasmatodea DNA methylation plays a part in controlling repetitive elements in addition to influencing gene expression. The proportion of repetitive DNA in the *L. migratoria* genome has been estimated to be 30% [43], ten times more than of *A. mellifera* [44]. The most abundant sequence represented in the cloned methyl-CpG enriched *L. migratoria* DNA corresponds to a previously characterised SINE retrotransposon, *Lml* [45], which alone accounts for approximately 2% of the genome. Interestingly, this element has been found to show differential expression in the nervous tissue of isolated and crowded *L. migratoria* [46] and is a source of small RNAs [47]. The methylation of repetitive DNA in the Orthoptera and Phasmatodea could be related to the very large genome size of these species [48–50], although other aspects of organismal biology are also likely to be important in determining levels of genomic methylation [17]. Indeed, repetitive DNA is estimated to comprise 45% of the sparsely methylated *B. mori* genome [16,51].

Perhaps unsurprisingly, no clear difference was observed when comparing the *HpaII/MspI* digestion patterns of solitary phase locusts to those of gregarious ones. The lower sensitivity of this method, which was unable to detect the biologically significant CpG methylation present in the *A. mellifera* genome, means that only relatively dramatic shifts in genome-wide methylation are likely to be identifiable. Finer scale mapping may reveal
biologically meaningful heterogeneity in the methylomes of solitarious versus gregarious locusts. For example, bisulphite sequencing has already been used to investigate differences in gene methylation between phenotypic variants in both the honeybee and the pea aphid [1,4,42]. The evidence of historical methylation shown here for some genes known to be differentially expressed between isolated and crowded locusts suggests that this will be an worthwhile area of investigation.

Complementary DNA sequencing confirmed that the *L. migratoria* genome contains orthologs of the invertebrate methyl-binding protein MBD2/3 [52] and at least two of the three animal Dnmt's; the maintenance methyltransferase, Dnmt1 [53] and the tRNA methyltransferase, Dnmt2 [54]. It is possible that Dnmt3, a proposed *de novo* methyltransferase [53], is also present. However, work towards identifying this gene has been problematic thus far. It is interesting to note that Dnmt2 does not appear to occur in either the *S. gregaria* [32] or *L. migratoria* [36] EST databases. Further research is required to clarify whether this gene is conserved in locusts as it is in the Hymenoptera [18,53–59] and the aphid *A. pisum* [1], or like in the representatives of other insect orders (*T. castaneum* [55], *B. mori* [16], *D. melanogaster* [56,57], *Anopheles gambiae* [7] and *Pediculus humanus* [58]), appears to have been lost.

It is now becoming apparent that there is considerable interspecific variation in the number of Dnmt proteins that insects possess [59]. In some lineages Dnmt1 appears to have been lost entirely, while other species have up to three variants. Whereas humans possess two Dnmt1 isoforms encoded by a single gene [60], in insects multiple Dnmt1 isoforms are encoded by separate genes. The degree of polymorphism observed between the two *L. migratoria* Dnmt1 variants suggests that they correspond to different genes. The tree topology observed for the insect Dnmt1 proteins is in agreement with observed sequence differences and the previous observation that multiple copies of the gene appear to have arisen independently in several different insect lineages.

The *L. migratoria* genome displays evidence of widespread historical gene methylation in the germline, with unigenes falling into two distinct populations; low CpG$_{OE}$ genes which show evidence of historical methylation and high CpG$_{OE}$ genes which are likely to have been hypomethylated in the evolutionary past. In other species it appears that these historical estimates are likely to be a good predictor of current levels of gene methylation [35,61]. Thus, it may be that in *L. migratoria* many of the low CpG$_{OE}$ genes undergo present-day methylation, at least in the germline.

Analyses of genome-wide methylation patterns in a diverse range of organisms have revealed a parabolic relationship between levels of gene body methylation and transcription, with both silent and highly expressed genes typically exhibiting little or no methylation and genes with intermediate levels of expression the most strongly methylated [9]. It has therefore been suggested that a major function of DNA methylation is to prevent aberrant transcripts arising from basic biological process genes that have broad, moderate expression [35]. In accordance with this, metabolism was the most overrepresented term in the low CpG$_{OE}$ group of *L. migratoria* unigenes, similar to *A. mellifera* [62]. It remains to be seen if DNA methylation is involved in the phase-specific regulation of metabolism genes, which a previous study has identified as being differentially expressed between isolated and crowded fifth instar locusts [37], consistent with the differing metabolic requirements of solitarious and gregarious individuals [26,63].

The high CpG$_{OE}$ group of *L. migratoria* unigenes was functionally enriched with five gene categories, the expression of three of which (muscle development, proteolysis and peptidolysis and carbohydrate metabolism) varies according to locust phenotypic phases [37,65], but the regulatory mechanisms governing these differences remain elusive. Our results provide clear evidence of genomic methylation in *L. migratoria*, with methylation occurring in both repetitive and protein-encoding regions of the genome. They also show that metabolic process genes, which are known to be differentially expressed in isolated versus crowded locusts [37], are the most proportionally overrepresented genes in a population of protein-encoding sequences that are likely to have been historically methylated. DNA methylation has been implicated in phenotypic plasticity and genomic imprinting in a diverse range of species.

Table 2. Results of functional enrichment analysis.

| CpG$_{OE}$ class | GO term | EASE score | Benjamini |
|------------------|---------|------------|-----------|
| low              | Metabolism | <0.01 | <0.01 |
|                  | Protein transport | <0.01 | 0.04 |
|                  | Intracellular protein transport | <0.01 | 0.09 |
|                  | Biosynthesis | <0.01 | 0.15 |
|                  | Electron transport | <0.01 | 0.23 |
|                  | Transport | <0.01 | <0.01 |
| high             | Carbohydrate metabolism | <0.01 | <0.01 |
|                  | Proteolysis and peptidolysis | <0.01 | <0.01 |
|                  | Chitin metabolism | <0.01 | <0.01 |
|                  | Muscle development | <0.01 | 0.03 |

For both low and high CpG$_{OE}$ gene populations, the five most overrepresented GO terms are presented. The EASE score is a modified Fisher exact probability p-value, the Benjamini value is the raw EASE score corrected for multiple comparisons using the Benjamini-Hochberg false discovery rate method [72]. Significantly overrepresented functional categories (p<0.05) are shown in italics.

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Conclusions

The ecology, ethology and physiology of locust phase polyphenism are becoming well understood. The lack of locust genome sequence data available at present and the size of the genome, which is approximately 30 times larger than that of *D. melanogaster* [64], have posed specific challenges to understanding the molecular processes underlying locust phenotypic plasticity. Considerable progress has been made in identifying genes differentially expressed between the solitarious and gregarious phases [37,65], but the regulatory mechanisms governing these differences remain elusive. Our results provide clear evidence of genomic methylation in *L. migratoria*, with methylation occurring in both repetitive and protein-encoding regions of the genome. They also show that metabolic process genes, which are known to be differentially expressed in isolated versus crowded locusts [37], are the most proportionally overrepresented genes in a population of protein-encoding sequences that are likely to have been historically methylated. DNA methylation has been implicated in phenotypic plasticity and genomic imprinting in a diverse range of species.
Figure 3. Methylation machinery of *L. migratoria*. ClustalW alignment of (a) Dnmt1, (b) Dnmt2 and (c) MBD2/3 partial sequences. For clarity, only regions with homology to conserved motifs are presented. Sequences from a larger selection of species were used for phylogenetic analyses, but for simplicity, only a subset of these is shown here. The reported *S. gregaria* Dnmt1 sequence could not be aligned with the other Dnmt1 sequences presented here. Completely conserved residues are shaded black; 80–99% identical residues are dark grey; 60–79% identical residues are light grey;
including some insects [1–5]. Given that locust phase change is a form of phenotypic plasticity that in some species involves the epigenetic transfer of phase state from parent to offspring, investigating the possible role of DNA methylation in regulating phase change in locusts represents a promising new area of research in this economically significant group of insects.

Materials and Methods

Locust culture and experimental material

A *L. migratoria* culture was established as described earlier [66]. Isolated and crowded individuals were reared essentially as specified for *Schistocerca gregaria* [67,68]. For both isolated and crowded treatments, eggs were harvested from egg pods on the ninth day after oviposition, when locusts were in the final stages of embryonic development. Isolated eggs were derived from mothers that were the offspring of crowded parents but were raised in isolation from approximately two days prior to hatching and were mated with males reared in the same manner.

Methylation-specific restriction enzyme assays

DNA was extracted from *L. migratoria*, *D. melanogaster* and *A. mellifera* adult head tissue and *L. migratoria* egg tissue using Genomic Tips (Qiagen). *M. musculus* genomic DNA was obtained commercially (New England Biolabs) and was derived from the embryonic fibroblast cell line NIH 3T3. Restriction digestions with *Hpa*II and *Msp*I (New England Biolabs) were carried out for 4 hr at 37°C with 2 µg of genomic DNA and 10 U of enzyme in a final volume of 50 µl. Digested DNA was separated on 0.8% agarose/TAE gel and visualised using ethidium bromide staining.

Cloning and sequencing of methylcytosine enriched genomic DNA

Genomic DNA from the eggs of ten crowd-reared mothers was pooled for this analysis. The high molecular weight genomic DNA that appeared as a discrete band following *Hpa*II digestion and agarose electrophoresis was excised from the gel and purified with the Wizard SV Gel and PCR Clean-Up System (Promega). DNA was further purified and concentrated by ethanol precipitation, using standard methods. Approximately 0.35 µg of purified DNA was then digested with 20 U *TaqI* (Roche) for 4 hr at 65°C in a final volume of 50 µl. Digested DNA was ethanol precipitated and used to perform a ligation reaction with pUC19 DNA that had been linearised with *Nar*I (New England Biolabs) and treated with Antarctic phosphatase (New England Biolabs). Ligation was carried out with T4 DNA ligase (New England Biolabs) at 16°C overnight. Recombinant DNA was used to transform chemically competent *Escherichia coli* TOP10 cells (Invitrogen) and positive transformants were identified using colony PCR. Sequencing was performed by Macrogen, Inc. (Korea) and clones were identified by performing NCBI BLAST [69] searches for homologous sequences (blastn algorithm; default parameters). Search terms were initially restricted to insect EST sequences. For clones that could not be identified, search terms were expanded to the entire nucleotide collection. The identity of insect EST sequences that produced top hits with the *L. migratoria* clones was determined using the blastx algorithm (default parameters).

Observed and expected CpG ratios

Analysis was performed using a data set of 12,148 unigenes downloaded from a publicly available database, LocustDB; http://locustdb.genomics.org.cn/ [36]. The database is derived from seven different cDNA libraries, which were constructed using the whole bodies of crowded locusts and head, hind leg and midgut tissue from isolated and crowded individuals. The unigenes in the dataset have a mean length of 598 bp (range = 101–4,278 bp). CpG ratios were calculated as described previously [1,62,70], incorporating a correction for the presence of Ns in the sequence data (File S1). Bimodality of the CpG ratio was assessed using the statistical program NOCOM to test whether the gene frequency distribution differed significantly from a unimodal distribution [62,71]. A subset of 2,482 unigenes for which pre-existing GO biological process annotation data was available in LocustDB was examined further to investigate whether there was any proportional overrepresentation of functional categories in low versus high CpG gene populations. Analyses were performed essentially as described elsewhere [62], with the stand-alone DAVID application EASE used to perform statistical calculations [72]. Functional enrichment was assessed by comparing the number of occurrences of each GO term in low and high CpG gene groups to a background list of all annotated unigenes.

Identification of Dnmt and MBD sequences

Degenerate primers (Table 3) were designed based upon alignments of published gene orthologs from other species, either manually or using the program CODEHOP [73]. PCR fragments were amplified from cDNA template, cloned and sequenced using standard methods. RNA was extracted from locust tissue using Trizol reagent (Invitrogen). cDNA was synthesised using the

| Table 3. Primers used in this study. |
|-------------------------------------|
| **Gene** | **Primer** | **Sequence (5’-3’)** |
|----------|------------|----------------------|
| Dnmt1    | forward    | TGTGCCGGAAGACCACTGTYCARG |
|          | reverse    | CCCAGTGTGGTTCCNGCRRTTNC |
| Dnmt2    | forward    | ATGAGTCCTCCNTYCARCC |
|          | reverse    | TTCAACTCCCCNTNACRTYTCC |
| MBD      | forward    | CCGAAAAGGTGAGCCAAAGAAAG |
|          | reverse    | TGAGGTTGCCTGGAAACTCCTC |

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Supporting Information

**Figure S1** Observed to expected ratios for all dinucleotide combinations. The number of genes is shown on the y-axis. For clarity O/E values greater than 3.00, where present, are not represented; omitting this data did not affect the shape of any of the distributions. (TIF)

**Table S1** Calculation and analysis of observed and expected CpG ratios. (DOC)

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**Author Contributions**

Conceived and designed the experiments: DT-E KLR GAS SJS. Performed the experiments: KLR. Analyzed the data: KLR DT-E. Conceived and designed the experiments: DT-E KLR GAS SJS. Performed the experiments: KLR. Analyzed the data: KLR DT-E. Contributed reagents/materials/analysis tools: GAS. Wrote the paper: KLR DT-E NL GAS SJS.

**References**

1. Walsh TK, Brison JA, Robertson HM, Gordon K, Jaubert-Possamai S, et al. (2010) A functional DNA methylation system in the pea aphid, *Acyrthosiphon pisum*. Insect Mol Biol 19: 213–220.

2. Bougouis S, Prantera G (2003) Imprinted facultative heterochromatinization in mealbugs. Genera 117(2): 271–279.

3. Li E, Beard C, Jarmisch R (1993) Role for DNA Methylation in Genomic Imprinting. Nature 366(6453): 362–363.

4. Kucharski R, Malecka J, Foret S, Malecka R (2000) Nutritional control of reproductive status in honeybees via DNA methylation. Science 319(5871): 1827–1830.

5. Siny F, Weaver I, Meaney M (2007) Maternal care, the epigenome and phenotypic differences in behavior. Reprod Toxicol 24(1): 9–19.

6. Bird AP, Taggart MH (1980) Variable Patterns of Total DNA and cDNA Methylation in Animals. Nucleic Acids Res 8(7): 1485–1497.

7. Marhold J, Rothe N, Pauli A, Mund C, Kuehle K, et al. (2004) Conservation of DNA methylation in dipteran insects. Insect Mol Bio 13(2): 117–123.

8. Lyko F, Ramsahoye BH, Jarmisch R (2000) Development - DNA methylation in *Drosophila melanogaster*. Nature 409(6812): 538–540.

9. Zemach A, McDaniel IE, Silva P, Zilberman D (2010) Genome-Wide Evolutionary Analysis of Eukaryotic DNA Methylation. Science 329(6080): 916–919.

10. Brown SJ, Henry JK, Black WC, Denell RE (1990) Molecular Genetic Manipulation of the Red Flour Beetle - Genome Organization and Cloning of a Ribosomal-Protein Gene. Insect Biochem 20(2): 185–193.

11. Lockat GA, Belhamed P, Malecka R (2010) Involvement of DNA methylation in memory processing in the honey bee. Neuropeptides 21(12): 812-816.

12. Field LM, Devonshire AL, French-Constant RH, Forde BG (1989) Changes in DNA methylation during loss and reselection of insecticide resistance in peach-potato aphids, *Acyrthosiphon pisum*.(Sulz.). FEBS Lett 243(2): 323–327.

13. Feyereisen R, Siomi MK (1995) The role of DNA methylation in invertebrates: Developmental regulation or genome defense? Mol Cell Biol 15(7): 880–891.

14. Zemach A, McDaniel IE, Silva P, Zilberman D (2010) Genome-Wide Evolutionary Analysis of Eukaryotic DNA Methylation. Science 329(6080): 916–919.

15. Suzuki MM, Kerr ARW, De Sousa D, Bird A (2007) CpG methylation is targeted to transcription units in an invertebrate genome. Genome Res 17(5): 756–756.

16. Regey A, Lamb MJ, Jablonska E (1998) The role of DNA methylation in invertebrates: Developmental regulation or genome defense? Mol Biol Evol 15(3): 890–891.

17. Wang Y, Jorda M, Jones PL, Malecka R, Ling X, et al. (2006) Functional CpG methylation system in a social insect. Science 314(5793): 643–647.

18. Nandini M, Yopi N (2003) The genome of the lipophagous Lepidoptera *Mamestra brassicae* has a vertebrate-like content of methylcytosine. Genera 119(2): 107–91.

19. Krauss V, Eisenhardt C, Unger T (2009) The Genome of the Stick Insect *Schistocerca gregaria* Is Strongly Methylated within Genes and Repetitive DNA. PLoS ONE 4(9): e7293.

20. Food and Agriculture Organization of the United Nations (2010) Migratory Locust. [Internet] Available: http://www.fao.org/ag/locusts-CCA/en/1010/1018/1078/index.html. Accessed Dec 15 2010.

21. Food and Agriculture Organization of the United Nations (2010) Migratory Locust. [Internet] Available: http://www.fao.org/ag/locusts-CCA/en/1010/1018/1078/index.html. Accessed Dec 15 2010.

22. McCaffery AR, Simpson SJ, Islam MS, Rosenging P (1990) A gregarizing factor present in the egg pod foam of the desert locust *Schistocerca gregaria*. J Exp Biol 203(3): 347–363.

23. Ben Hamouda A, Ammar M, Ben Hamouda MH, Baouain A (2009) The role of egg pod foam and rearing conditions of the phase state of the Asian migratory locust *Locusta migratoria migratoria* (Orthoptera, Acrididae). J Insect Physiol 55(7): 617–623.

24. Tanaka S, Maeno K (2010) A review of maternal and embryonic control of phase-dependent progeny characteristics in the desert locust. J Insect Physiol 56(8): 911–918.

25. Uvarov BP (1966) Grasshoppers and Locusts. Vol. 1. Cambridge: Cambridge University Press. 481 p.

26. Peever MP, Simpson SJ (2009) Locus Phase Polyphenism: An Update. Advances in Insect Physiology 36: 1–272.

27. Gray LJ, Sword GA, Anstey ML, Clissold FJ, Simpson SJ (2009) Behavioural phase polyphenism in the Australian plague locust (*Chortoicetes terminifera*). Biol Lett 5(3): 306–309.

28. Sword GA, Lecoq M, Simpson SJ (2010) Phase polyphenism and preventative locust management. J Insect Physiol 56(6): 949–957.

29. Simpson SJ, Sword GA (2009) Phase polyphenism in locusts: Mechanisms, population consequences, adaptive significance and evolution. In Whitman D, Ananthakrishnan T, eds. Phenotypic Plasticity of Insects: Mechanisms and Consequences. Plymouth: Science Publishers Inc. pp 147–190.

30. Song H (2005) Phylogenetic perspectives on the evolution of locust phase polyphenism. J Orthoptera Res 14(2): 235–245.

31. Wyatt GR (1951) The Purine and Pyrimidine Composition of Deoxypentose Nucleic Acids. Biochem. J 48(5): 584–590.

32. Boerjan B, Sas F, Ernst UR, Tobbback J, Lemière F, et al. (2011) Locus phase polyphenism: Does epigenetic precede endocrine regulation? Gen Comp Endocrinol 173(1): 120–128.

33. Guo W, Wang XH, Zhao DJ, Yang PC, Kang L (2010) Molecular cloning and temporal-spatial expression of I element in gregarious and solitary locusts. J Insect Physiol 56(8): 945–948.

34. Bird AP (1980) DNA Methylation and the Frequency of CpG in Animal DNA. Nucleic Acids Res 8(7): 1499–1504.

35. Suzuki MM, Kerr ARW, De Sousa D, Bird A (2007) CpG methylation is targeted to transcription units in an invertebrate genome. Genome Res 17(5): 625–631.

36. Ma ZY, Yu J, Kang L (2006) LocusDDE: a relational database for the transcriptome and biology of the migratory locust (*Locusta migratoria*). BMC Genomics 7: 11.

37. Kang L, Chen XY, Zhou Y, Liu BW, Zheng W, et al. (2008) The analysis of large-scale gene expression correlated to the phase changes of the migratory locust. Proc Natl Acad Sci U S A 101(51): 17611–17615.

38. Adams RLP (1990) DNA Methylation - the Effect of Minor Bases on DNA Protein Interactions. Biochem J 265(2): 309–320.

39. Sarkar S, Gupta VS, Hendre RR, Rao SRV (1992) 5-Methylcytosine content in mealybugs. Genetica 117(2): 271–279.

40. Karthik S, Thirunarayan V, Sahay J, Ananthakrishnan T, eds. Epigenetics: Invertebrates: Developmental regulation or genome defense? Mol Cell Biol 20(3): 347–363.

41. Adams RLP (1990) DNA Methylation - the Effect of Minor Bases on DNA Protein Interactions. Biochem J 265(2): 309–320.

42. Karthik S, Thirunarayan V, Sahay J, Ananthakrishnan T, eds. Epigenetics: Invertebrates: Developmental regulation or genome defense? Mol Cell Biol 20(3): 347–363.
