The Drosophila G9a gene encodes a multi-catalytic histone methyltransferase required for normal development

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ABSTRACT

Mammalian G9a is a histone H3 Lys-9 (H3–K9) methyltransferase localized in euchromatin and acts as a co-regulator for specific transcription factors. G9a is required for proper development in mammals as g9a/C0/g9a/C0 mice show growth retardation and early lethality. Here we describe the cloning, the biochemical and genetical analyses of the Drosophila homolog dG9a. We show that dG9a shares the structural organization of mammalian G9a, and that it is a multi-catalytic histone methyltransferase with specificity not only for lysines 9 and 27 on H3 but also for H4. Surprisingly, it is not the H4–K20 residue that is the target for this methylation. Spatiotemporal expression analyses reveal that dG9a is abundantly expressed in the gonads of both sexes, with no detectable expression in gonadectomized adults. In addition we find a low but clearly observable level of dG9a transcript in developing embryos, larvae and pupae. Genetic and RNAi experiments reveal that dG9a is involved in ecdysone regulatory pathways.

INTRODUCTION

Modifications of histones are an important mark for transcriptional regulation during embryonic development. The protruding tales of the histones are modified by acetylation, phosphorylation, ubiquitination and arginine and lysine methylation, and the combinations are hypothesized to form a histone code (1,2). The best-characterized substrates for lysine methylation in eukaryotic cells are histone proteins, although methylation of several non-histone proteins, such as the tumor suppressor p53, has been reported as well (3).

Histone H3 has been shown to be methylated on lysine residues K4, K9, K27, K36 and K79 whereas in histone H4, K20 is methylated (4,5). Each of these lysine side chains can be mono-, di- or tri-methylated by histone lysine methyltransferases (HKMTases), which, except for Dot1 (6), carry a catalytic SET [Su(var), Enhancer of Zeste, Trithorax] domain (7). The SET domain is a conserved ~130 amino acid sequence, which is flanked by the less conserved pre-SET and post-SET regions at the amino and C-termini, respectively. The specificity of a HKMTase, as well as the number of methyl residues that attaches to a lysine residue, depends on the structure of the HKMTase or the presence of additional co-factor proteins (8). On the other hand Ezh2 requires the presence of the co-factors suppressor of zeste-12 (SUZ12) and embryonic ectoderm development (Eed) for tri-methylation of H3–K27 (9). The HKMTase ERG-associated protein (ESET) di-methylates H3–K9, but is converted into a tri-methylating enzyme by its association with a mouse-activating transcription-factor-associated modulator (mAM) (10). The methylated histones recruit proteins that carry CHROMO, TUDOR or WD40 domains and are capable of specific interactions with differently methylated lysine residues reviewed in Ref. (11). This recruitment step is likely to define a unique functional readout for individual lysine methylations. Thus, tri-methylation of lysine 9 in histone H3 by Suv39H1 and Suv39H2 creates a binding site for the chromodomain-containing heterochromatic protein HP1 which is thought to induce heterochromatin formation (12).

Mammalian G9a mono- and di-methylates H3–K9 at euchromatic loci (14,15), and has recently also been found at heterochromatic loci (16). In g9a/g9a mice H3–K9
methylation is drastically reduced resulting in severe growth retardation and early lethality (17). The loss of G9a primarily affects the methylation of H3–K9 in euchromatic regions (14). G9a is the major euchromatic histone H3–K9 methyltransferase in higher eukaryotes but in Drosophila the euchromatic H3–K9 HMTase has not been characterized. Although the H3–K9 methylation is strongly reduced in Su(var)3–9 null mutants, a small amount of H3 molecules remain methylated at K9 suggesting the existence of other K9 specific HMTases in Drosophila (18).

There are several reports demonstrating the silencing effects from H3–K9 methylation, including the inactive X chromosome of female mice and humans (19), and developmentally regulated genes (20).

In a search for SET domain containing genes in Drosophila that might code a K9 specific HMTase, we performed a bioinformatics search of the Drosophila melanogaster genome and found the gene CG2995 which share significant homology to mammalian G9a. In this paper we describe the cloning, and the biochemical and genetic analyses of CG2995. We show that it encodes a histone methyltransferase specific for H3–K9, K27 and H4, and that it shares the structural organization of mammalian G9a. Therefore, we suggest that CG2995 is renamed dG9a. It adds up to three methyl groups to unmethylated H3 and H4. Our results indicate a role for dG9a in germ cell formation. Using RNAi we show that dG9a is critical for development, very likely by being involved in the ecdysone regulated gene expression.

MATERIALS AND METHODS

Fly handling and generation of transgenic flies

All genetic crosses were carried out at 25°C. Fly lines were obtained from the Bloomington Drosophila stock centre.

Generation of double stranded (ds) RNA was performed by using the phBS and pUds-GFP vectors as described in Ref. (21). A 756 bp fragment of dG9a cDNA was PCR amplified with the 2995UBamHI (5′-CAAGGATCTGTCGCATTGTGTT-3′) and 2995right (5′-GGATCCTGTCGCATTGTGTT-3′) primers.

Transgenic flies were generated by P-element mediated transformation, and nine independent lines on different chromosomes of female mice and humans (19), and developmentally regulated genes (20).

In a search for SET domain containing genes in Drosophila that might code a K9 specific HMTase, we performed a bioinformatics search of the Drosophila melanogaster genome and found the gene CG2995 which share significant homology to mammalian G9a. In this paper we describe the cloning, and the biochemical and genetic analyses of CG2995. We show that it encodes a histone methyltransferase specific for H3–K9, K27 and H4, and that it shares the structural organization of mammalian G9a. Therefore, we suggest that CG2995 is renamed dG9a. It adds up to three methyl groups to unmethylated H3 and H4. Our results indicate a role for dG9a in germ cell formation. Using RNAi we show that dG9a is critical for development, very likely by being involved in the ecdysone regulated gene expression.

RT–PCR

Tests and ovaries from adults were dissected in Ringer’s solution (6.5 g NaCl, 0.14 g KCl, 0.2 g NaHCO3, 0.12 g CaCl2 and 0.01 g NaH2PO4 per liter). Total RNA was isolated from indicated tissues or stages by the use of TRIzol® reagent (Invitrogen), and 5 µg of total RNA were reverse transcribed with SuperScript III RNase H-free reverse transcriptase (RT) (Invitrogen). A random primer pd(N)6 was used for first-strand synthesis. PCR was performed with 2995left (5′-GATGAAAGAAGAAGTGCACA-3′, located in exon 5) and 2995right (5′-GATGAAAGAAGAAGTGCACA-3′, located in exon 9) primers and with rp49 primers (23) as loading control for 35 cycles at an annealing temperature of 56°C.

Immunostainings and immunofluorescence

The anti-dG9a polyclonal antiserum was raised in rabbits (Eurogentec S.A) against a synthetic peptide containing dG9a residues 1623–1637. The antiserum was affinity purified.

Polytene chromosomes from the salivary glands of third instar larvae were prepared and stained essentially as described in Ref. (24).

Biowinformatics tools

Database searches were performed using BLASTP, TBLASTN and PSI-BLAST (http://www.ncbi.nlm.nih.gov/BLAST/). Protein domains were identified using the programs RPS-BLAST (NCBI) and ProfileScan (http://hits.isb-sib.ch/cgi-bin/PFSCAN) searching the Pfam-A, Prosite profiles and SMART databases (NCBI). Nuclear localization signal was detected by using PredictNLS (http://cubic.biocolumbia.edu/predictNLS/). Amino acid sequence alignments were created using ClustalX 1.8 (http://www-igbmc.u-strasbg.fr/BioInfo/ClustalX) with default parameters and manual adjustments from GeneDoc 2.6.001 (http://www.psc.edu/biomed/genedoc/).

Whole-mount in situ hybridization

RNA in situ hybridization using digoxigenin-labeled antisense RNA probes was performed as described previously (25,26). A cDNA containing dG9a was linearized with
BamHI and used as template to make a 674 bp dG9a RNA probe.

Western analysis

Nuclear extract from 0–12 h dechorionated embryos (27) was separated on a SDS–PAGE (8%). Proteins were transferred onto PVDF membrane (Amersham) and probed with an anti-dG9a antibody (1:500) using standard procedures. Secondary antibodies conjugated to HRP (Amersham) were used according to the manufacturer’s instructions. Detection of antibody signals was performed with chemiluminescence (Pierce).

Generation of baculovirus, viral transfection and dG9a purification

Amino acids 789–1637 of dG9a was PCR amplified with the primers 2995 SpeIFlag (5'-GACTACAAGGACGACGATGACAAAGTTTGCTATGTCCAGAGCCCCTC-3', FLAG tag underlined) and 2995 KpnI (5'-TGCGGTACCCGTACGGTTCATTTTCTC-3') cloned into pFastBac(Tm)1 (Invitrogen) as a SpeI/KpnI fragment. Site-directed mutagenesis of flag-dG9a (789–1637 amino acid) was performed using the QuickChange kit (Stratagene) with primers CG2995H1536KBstIFwd (5'-ATGGGAATATGTAACCAGGGTGGGCAATCTA-3' and CG2995H1536KBstIrew (5'-CTTCACACGACGTCTGGAACG-3').

FLAG tagged dG9a in pFastBac(TM)1 was transformed into DH10Bac (Invitrogen) and bacmid purified according to protocol (Invitrogen). A monolayer of SF9 (Spodoptera frugiperda) was transfected with 1.2·10^7 mCi/ml and recombinant viruses were used for test expression. Cell extracts were checked, 48 h post-transfection, for fusion protein expression using anti-FLAG monoclonal antibody (Sigma). For routine protein expression using anti-FLAG monoclonal antibody and recombinant viruses were used for test expression. Cell extracts were checked, 48 h post-transfection, for fusion protein expression using anti-FLAG monoclonal antibody (Sigma). For routine protein expression using anti-FLAG monoclonal antibody and recombinant viruses were used for test expression.

Histone purification and nucleosome assembly

Recombinant Drosophila histones were expressed and purified from Escherichia coli BL21(DE3)pLys, and reconstituted into octamers as described previously (28). Recombinant histone H3 carrying the mutations K9A, K27A or K9A/K27A were expressed from plasmids kindly provided by D. Reinberg, and histone H4 carrying a K20A mutation was expressed from a plasmid given by D. Reinberg, and histone H4 carrying a K20A mutation was expressed from a plasmid given by D. Reinberg, and histone H4 carrying a K20A mutation was expressed from a plasmid given by D. Reinberg. Native histones were purified from 0–12 h Drosophila embryos essentially as described in Ref. (30). Nucleosomes were reconstituted on circular pBS(KS) (Stratagene) by salt dialysis over night from 2 to 0.1 M NaCl (31).

Histone methyltransferase assay

Histone methyltransferase assays were performed as described in Ref. (29). In short, 100 ng of eluted dG9a was mixed with 1 or 2 μg of histone H3, H4, octamer or nucleosomes and S-adenosyl-[methyl-^3H]-methionine (25 μCi/ml) (Amersham) in a buffer containing 12.5 mM Tris–HCl, pH 8.8, 1 mM DTT, 50 mM NaCl, 50 ng/μl BSA and 2.5 mM MgCl2. The reaction was incubated at 30°C for 1 h and stopped by adding SDS–PAGE loading buffer. The histones were separated by 15% SDS–PAGE, Coomassie stained, amplified and dried. The autoradiograph was developed after 1 and 2 weeks. For experiments with mouse G9a, we used 250 ng of protein and the incubation time was 30 min. Exposure time for autoradiograph was 1 and 2 days.

MALDI-TOF analysis

Methylation reactions were carried out as described above with 0.5 μg of histone H3 or H4 and 250 μM of S-Adenosylmethionine (New England BioLabs). The reaction was stopped by addition of SDS–PAGE loading buffer and the histones were separated by 15% SDS–PAGE. The Coomassie stained bands corresponding to H3 and H4 were excised and subjected to chemical modification to derive free amino groups of lysine residues as described previously (33). H3 and H4 were digested over night with 100 ng of sequencing-grade trypsin (Promega) in a total volume of 40 μl according to manufacturer’s protocol. In order to purify the methylated peptides from contaminating salts or acrylamide the peptide solution was passed over a pipette tip. The methylated peptides from contaminating salts or acrylamide the peptide solution was passed over a pipette tip. The methylated peptides from contaminating salts or acrylamide the peptide solution was passed over a pipette tip.

Edcsyne feeding experiments

The feeding experiments were performed essentially as described in Ref. (36).

RESULTS

CG2995 is the Drosophila homolog of the mouse G9a HKMTase

We performed a bioinformatics search of the D. melanogaster genome with the Su(var)3–9, E(Z) and Trithorax SET domains and found novel genes encoding putative SET proteins. Performing a BLASTP search with the SET domain of one of these proteins, the annotated CG2995 protein,
against the mouse and human database identified it as the Drosophila homolog of the G9a protein. This SET domain alone shares 61% identity (76% similarity) with the corresponding domain of the mouse G9a protein (Figure 1A). In comparison, the SET domain of CG2995 shares 45% identity with the SET domain of the Su(var)3–9 protein of Drosophila, suggesting that CG2995 is the only homolog of G9a in the Drosophila genome. This is emphasized when looking at the pre-SET/SET/post-SET regions, where it is notable that CG2995 is more similar to mouse G9a than to Su(var)3–9 or dSET2. Thus, dSu(var)3–9 versus G9a shows 37% identity, 55% similarity, dSET2 versus G9a displays 36% identity, 50% similarity, and CG2995 versus G9a has 47% identity, 68% similarity. CG2995 is located as the third gene in region 1A1 on the X chromosome, and a further comparative analysis of the CG2995 protein with the mouse G9a shows that the CG2995 has 33% identity and 49% similarity to the mouse protein (1172–1263 amino acid). The fly protein is longer at the N-terminus but otherwise shares the same module organization as its mouse counterpart. The CG2995 protein contains multiple putative domains in addition to the SET domain, like the adjacent cysteine-rich regions [the pre-SET (also called SAC); (37)], and conserved cysteine residues in the C-terminal region of the SET domain that corresponds to the post-SET domain (Figure 1A), which has shown to be required for enzymatic activity (38). Within the SET domain, a H(R)ΦΦNHSC motif (where Φ indicates a hydrophobic residue) has previously been shown to be an important catalytic site. For SUV39H1 protein, a histidine-to-arginine mutation of the first histidine (His^{320}) in the HΦΦNHSC^{320} motif resulted in a 20-fold higher catalytic activity (38). This observation suggests that the H(R)ΦΦNHSC motif is correlated with the HKMTase activity. The CG2995 protein contains a RΦΦNHSC^{1532} motif (Figure 1A, underlined), together with another motif reported to be needed for HKMTase activity, GE(x)ŚY, located in the C-terminal end of the SET domain (38; Figure 1A, underlined).

In addition, the CG2995 protein harbors contiguous copies of a 33-amino acid repeat (Figure 1B). This repeat, originally identified in the Notch protein of Drosophila and known as the ankyrin repeat, is also found in G9a and in a number of other proteins involved in intracellular protein–protein interactions (39).

An AT-hook also is found in the N-terminus part of the CG2995 protein. The AT-hook is a small DNA-binding protein motif that was first described in the high mobility group non-histone chromosomal protein HMG-I(Y). Since its discovery, this motif has been observed in other DNA-binding proteins from a wide range of organisms. Furthermore, AT-hook motifs are frequently associated with known functional domains seen in chromatin proteins and in DNA-binding proteins (e.g. histone folds, homeodomains and zinc fingers). In general, it appears that the AT-hook motif is an auxiliary protein motif co-operating with other DNA-binding activities and facilitating changes in the structure of the DNA either as a polypeptide on its own [e.g. HMG-I(Y)] (40) or as part of a multidomain protein.

Figure 1. The domain organization is conserved between dG9a and G9a. (A) Alignment of SET domains and flanking cysteine-rich regions of mouse and Drosophila dG9a protein. The degree of conservation is distinguished at four levels (100, 80 and 60%, and not conserved), where 100% has the darkest shade of grey. The upper and lower case letters in the consensus line indicate 100 and 80% conservation within all groups, respectively. Numbers in the consensus line represent conserved similarity groups as defined by the Blossum 62 scoring table. The conserved R(H)ΦΦNHSC and the FDYG motifs are underlined. (B) Domain organization within Drosophila protein dG9a. An AT-hook and an ankyrin motif are found in addition to the SET domain.
[e.g. Swi2p in Saccharomyces cerevisiae or HRX (ALL-1) in Homo sapiens] (41). It is most interesting that this motif seems to be specific to known or predicted chromosomal/DNA-binding proteins, suggesting that it may act as a versatile minor groove tether (41). A nuclear localization signal is found in the N-terminal of the protein. In conclusion, CG2995 has a high level of similarity to mouse G9a and we suggest CG2995 is the Drosophila homolog of G9a and will refer to it as dG9a.

Full-length cDNA was cloned by RACE and RT–PCR. This cDNA revealed that the dG9a gene consists of 10 exons with a 4911 bp open reading frame (ORF) encoding a protein of 1637 amino acid. The coding region ends by an in-frame stop codon that is followed by a poly(A) signal 1173 downstream, suggesting that it is full-length and consistent with the annotated sequence in FlyBase (http://flybase.bio.indiana.edu; Figure 1B). Northern analysis also showed that there is only one transcript of expected size (data not shown).

The dG9a protein localizes to euchromatin
Antibodies specific to dG9a were generated by immunization of rabbits with a peptide corresponding to the last 14 amino acids (1623 through 1637 amino acid). This antibody recognized a band of the predicted size of ~180 kDa (Figure 2A). The localization of dG9a protein was investigated by analysis of polytene chromosomes from salivary glands (Figure 2B). The immunostaining showed discrete banding pattern in euchromatic regions with no staining observed in the chromocenter.

Spatiotemporal expression of dG9a
To investigate the spatiotemporal expression of dG9a we first used semi-quantitative RT–PCR. As shown in Figure 3A, a low but measurable amount of dG9a transcript is present in 0–3-h-old embryos. In 3–6-h-old embryos the expression of dG9a is barely detectable, indicating that the dG9a transcripts seen in 0–3-h-old embryos are of maternal origin. Between 6 and 21 h of embryogenesis the expression of dG9a is low but clearly discernible, and about the same level of expression is observed throughout larval development, with a slightly elevated expression during the third larval instar. Then, in 12–46-h-old pupae there is no or very little expression of dG9a. In adult flies the expression of dG9a is restricted to the gonads in both sexes (Figure 3A, last four lanes). However, we cannot rule out the possibility that dG9a is expressed in one or more tissues of the gonadectomized flies, but at a level too low to be detected by the RT–PCR settings used here.

Next, in order to study the spatiotemporal expression of dG9a in more detail, we stained ovaries and embryos with the dG9a antibody. The immunostainings revealed that dG9a is expressed in all cells of the ovary, including the germarium (Figure 3C) where especially the nurse cells,
Figure 3. Spatiotemporal expression of dG9a. (A) Developmental RT–PCR shows that dG9a is maternally deposited in the egg, and that there is moderate expression during the larval development. dG9a is present in all developmental stages investigated. (B–K) dG9a is present from the very start of oogenesis through the end of oogenesis in wild-type ovarioles. Anterior is to the left, posterior to the right. dG9a in red (right column) and the nuclei is counter stained with DAPI in blue (left column). (B and C) The early stages of oogenesis development. The dG9a protein is present from the very start. (D and E) Stage 10B ovaries. dG9a localizes to nuclei in both nurse and follicle cells. An accumulation of protein is observed in the region where the anterior polar cells and the centripetal follicle cells are located, arrowheads and in the posterior follicle cells, arrow. (F and G) Stage 11. Shortly after centripetal migration (stage 10B), the nurse cells rapidly transfer their contents into the oocyte (stage 11) then begin to degenerate and undergo apoptosis (stages 12–14). (H and I) Stage 12. Dumping complete, no or very little dG9a is detectable in the degenerating nurse cell nuclei, but is still present in the follicle cells. Notice the accumulation of dG9a protein in the extreme posterior part of the egg, arrowheads, where the posterior polar cells located. (J and K) Stage 14. The egg is fully developed and dG9a protein is maternally deposited. (L–S) Lateral views of wild-type embryos hybridized with digoxigenin-labeled RNA probes (L and N with Nomarski optics) or with a dG9a antibody (M, O, Q and S). Anterior is to the left and dorsal is up. The nuclei are counter stained with DAPI in blue. (L and M) Embryo at syncytial blastoderm stage (stage 4, ~1.5–2.5 h). dG9a is localized to the nuclei. In early embryos the message and the protein are ubiquitously distributed due to its maternal contribution. (N and O) Embryo during germband extension (stage 9). (P and Q) Stage 12. In late-stage embryos, expression is strongest in the CNS and the neuroectoderm. (K and S) Stage 13. Surface view of embryo at the completion of germband shortening.
which undergo a dramatic endoreplication, stain heavily. It also clearly shows that dG9a is localized to the nucleus. In stage 10B egg chambers it appears that dG9a is markedly upregulated in what appears to be the centripetal follicle cells (Figure 3E, arrowheads). At this stage, a moderate upregulation is also discernible in the posterior follicle cells (arrow). During stage 11 nurse start dumping their content into the oocyte, which is revealed by an accumulation of dG9a at the border between the growing oocyte and the degenerating nurse cells (Figure 3G). At this stage an increased expression of dG9a is also found in the posterior follicle cells (Figure 3G, arrow head). In stage 12 the amount of dG9a has increased considerably and has started to move into the oocyte. An accumulation in the posterior follicle cells is now prominent (Figure 3I, arrow). At stage 14, dG9a appears to be evenly distributed in the mature oocyte (Figure 3K).

In blastoderm embryos (stage 4, ~1.5–2.5 h) dG9α transcript as well as protein are present in the syncytial nuclei (Figure 3L and M). During stages 9 and 12, dG9α expression appears to be more abundant in the central nervous system (CNS) and the neuroectoderm (Figure 3N, O and Q). Figure 3S shows a surface view of embryo at the completion of germband shortening, with all cells evenly stained.

Expression and purification of recombinant histone methyltransferase dG9α

In order to investigate the enzymatic properties of dG9α we expressed a FLAG tagged N-terminal fragment (789–1637 amino acid) using a baculovirus expression system. The purified dG9α was soluble and had the expected molecular size of 95 kDa (Figure 4A). To confirm that dG9α has HKMTase activity we incubated it in presence of H3-S-adenosyl-methionine (SAM) and different substrates (Figure 4B). dG9α methylates H3 and H4 present as free histones but had no detectable activity on nucleosomal arrays. In contrast to the recombinant dG9α, mouse G9α only methylates H3 even when other histones are present (Figure 4C). In order to exclude the possibility that the unexpected H4 HMT activity is due to a contaminating activity co-purifying with the recombinant dG9α we expressed the enzyme carrying a point mutation within the SET domain (H1536K) that dramatically impairs its catalytic activity. The mutated enzyme was not able to methylate either H3 or H4 indicating that both methylations are a result of dG9α activity (Figure 4D). In order to determine the substrate specificity we performed a quantitative MALDI-TOF analysis of H3 molecules methylated by dG9α. Similarly to the activity of the mouse ortholog (42), dG9α methylates exclusively K9 and K27 with K9 being the preferred substrate in wild-type (wt) H3. As shown for mG9α (43) dG9α is able to add up to three methyl groups to H3 (Figure 4H). This finding is confirmed by using H3 molecules carrying a lysine to alanine replacement at position 9 and 27 or both (Figure 4E). We observe decreased methylation efficiency on H3 K9A and H3 K27A compared to wt H3. In a filter binding assay we observed a 70% reduction when K9 was mutated and a 50% reduction when K27 was mutated. When both H3 lysine residues were mutated (K9A and K27A) we observe a lower activity (efficiency of 27%) indicating that in absence of K9 and K27 dG9α is also able to methylate other lysines. Mouse G9α also showed a decreased activity (27%) towards the double mutant (K9A/K27A) (Figure 4F). When we use highly active mG9α (43), it methylates wt H3 and the H3 molecules carrying a single mutation on K9 or K27 with a similar efficiency. However, as we use relatively long reaction times we can not exclude the possibility that K9 is methylated faster than K27, which explains the larger differences observed in previous publications (16,44).

Interestingly dG9α was also able to methylate histone H4 (Figure 4B). This is in marked contrast to the mouse G9α where we do not observe such an activity (Figure 4C) (42). The only lysine residue of H4 shown to be methylated in vivo is K20 and the first HKMTase identified with this activity was hPR-Set7/dSET8 (45,46). Other HKMTases in Drosophila shown to methylate H4 lysine 20 are Ash1 and Suv4–20 (47,48). Ash1 is in addition able to methylate lysine 4 and 9 in histone H3 (47). From these experiments we concluded that dG9α also is a multi-catalytic histone methyltransferase with specificity for lysine 9 and 27 on H3 and possibly lysine 20 on histone H4. However, when we incubated dG9α with H4 carrying a mutation of lysine 20 to alanine we observed no reduction of activity (Figure 4G). To further investigate the specificity, we tested whether dG9α was able to methylate H4 molecules carrying different N-terminal deletions (29). We observed that dG9α could methylate the H4 N-terminus when the first five amino acids were deleted, excluding K5 as a possible substrate. However, the activity was lost when we used the H4 A10 mutant (Figure 4G). This suggests that the substrate is K8, but considering that the minimal substrate specificity for mG9α surrounding K9 contains seven amino acids (TARK-STG) (49) we cannot exclude that another downstream lysine can serve as a substrate. MALDI-TOF analysis of H4 methylated by dG9α, showed that only the peptide containing amino acids 4–17 was methylated in vitro (Figure 4H). We conclude from these experiments that dG9α can methylate lysine 8, 12 or 16 of H4 in vitro. It remains to be seen to what level these lysines are methylated in vivo and what the function of this methylation is.

dG9α is required for normal development

To investigate the in vivo function of dG9α, transgenic flies with an inverted-repeat UAS-dG9α.IR construct were crossed to different GAL4-driver lines (22). The vector used for making the inverted repeats has an independent UAS-GFP marker so that a tissue exposed to RNAi will simultaneously show GFP expression [pUds-GFP; (21)] as an internal control to RNAi expression. In addition, down regulation of dG9α was confirmed by RT–PCR (Figure 5A).

Using the ubiquitously expressed da-GAL4 driver with the UAS-dG9α.IR flies, the progeny developed normally until the end of the third larval instar. However, these RNAi larvae did not form their puparium and crawling larvae were found after 7–8 days (Figure 5B). The majority of these larvae developed melanotic tumors, either one or two larger ones or several smaller. The larvae finally stopped moving, and in the few cases where ‘pseudo-prepupae’ were formed,
these maintained the elongated larval form and failed to evert the anterior spiracles; there was only slight melanization of these ‘pseudo-prepupae’ (Figure 5C).

The defects in puparium formation seen in dG9a RNAi animals could result from either a decrease in the ecdysone titer or a decrease in the ability of the ecdysone signal to be transduced. To distinguish between these possibilities we examined the effects of feeding ecdysone to dG9a RNAi larvae. This method has been shown to effectively rescue phenotypes associated with ecdysone-deficient mutations (36). Mid- and late-third instar larvae were transferred to food either with or without 20-hydroxyecdysone (20E) for 6–8 h and scored on a 12-h basis. Feeding 20E to dG9a RNAi larvae did not rescue them to puparium formation. Therefore, we conclude that ecdysone is not limiting in the da-GAL4/UAS-dG9a;IR animals and that dG9a functions downstream of ecdysone biosynthesis and release.

When the ubiquitously expressed, but weaker, Act5C-GAL4 driver was used, development of progeny of genotype Act5C-GAL4/UAS-dG9a;IR proceeded up to and through puparium formation (data not shown). However, most of the pupal cases were only lightly tanned and no flies eclosed from these cases. Upon dissection, dead and partially differentiated pharates with no eye pigmentation or legs were found (data not shown). Development appeared to have proceeded further in the posterior part of fly. Of all pupae formed (N = 133), ~17% developed normally, and the eclosed flies were without exception females.

As immunostaining indicated a role for dG9a in neuroectoderm and CNS, a driver which expresses GAL4 in the brain and throughout CNS, but not in discs, of third instar larvae, P[GawB]c698a, was used. The phenotypes observed were similar to those observed with the Act5C-GAL4 driver. Here, however, most of the pupae were partly more melanized (Figure 5D), without differentiation proceeding any further than that described for the Act5C-GAL4 driver (Figure 5E). A small portion (~5%) of the pupae actually developed normally but of these almost all failed to escape from the pupal case. The very few that succeeded were all normal females and lived for at least 2 weeks.

The results described above strongly suggest a defect in ecdysone responses at puparium formation, similar to those reported earlier for mutants in the ecdysone pathway (50). Also, using RNAi very similar results were obtained with the H3–K36 HKMTase dSet2 gene (M. Stabell, unpublished data).

Ecdysone controls wing morphogenesis and cell adhesion by regulating integrin expression during metamorphosis (51). Therefore, to test further the possible involvement of dG9a in the ecdysone regulation hierarchy, we triggered dG9a RNAi in the wing disc by the ap-GAL4 driver. This resulted in slightly held-out/up wings with an anterior–posterior compression (Figure 5F), and occasionally in blister formation (Figure 5G). We next generated flies of genotype ap-GAL4,UAS-dG9a;IR/ExR^M554Q. ExR^M554Q is a loss of function mutation where only half the amount of the ecdysone receptor is present in mutant flies. As shown in Figure 5I, these animals show a wing phenotype of a more extreme character, with both wings having a blister and being clearly smaller than normal, most likely because the wings never completely unfold. This phenotype has 100% penetrance. Taken together, these results support the notion that dG9a functions in ecdysone signaling pathways during development.

**DISCUSSION**

dG9a is an euchromatic histone methyltransferase

In this study we show that CG2995 is the *Drosophila* homolog of the HKMTase G9a, and that it specifically mono-, di- and trimethylates H3–K9, H3–K27 and K8, 12 or 16 in H4. This methylation pattern is mainly correlated with silencing (11) suggesting that dG9a is involved in transcriptional repression. Further, we showed that dG9a methylates free histones but has no detectable activity on nucleosomal arrays. As revealed by the staining of polytene chromosomes, the centromeric region, where Su(var)3–9 predominantly stains (52), is devoid of dG9a. We therefore conclude that dG9a is a euchromatic histone methyltransferase that acts on loosely packed DNA and that methylation by dG9a may occur on pre-assembled histones.

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**Figure 4.** Characterization of recombinant dG9a. (A) Eluted FLAG tagged dG9a (789–1637 amino acid) was separated on a 12% SDS–PAGE and stained with Coomassie blue G250. (B) In vitro methylation reactions using dG9a (lanes 1–6), no enzyme (lane 7) and DsUav( var)3–9 (lane 8). In the reaction we used 1 μg of different histones: recombinant histone H3 (lane 1), recombinant histone H4 (lane 2), recombinant (lane 3) and native histone octamer (lane 5) and recombinant and native nucleosomes (lanes 4 and 6) reconstituted on circular pBS(KS) from equimolar amounts of histones. The upper panel shows Coomassie stained gel and the lower panel the autoradiograph. (C) Activity of recombinant mouse G9a expressed in baculovirus infected cells (a kind gift from S. Pradhan). HKMTase activity on 1 μg of different histone substrates: recombinant histone H3 (lane 1), recombinant histone H4 (lane 2), recombinant histone H4 (lane 3) and recombinant and native histone octamers (lanes 4 and 6). Mock control (lane 7) is incubation of recombinant octamer without enzyme. The Coomassie gel is shown at the top and the corresponding autoradiograph at the bottom. (D) FLAG dG9a wild type versus H1536K mutation of the conserved region of the SET domain. The upper panels shows a western blot of the two proteins. Recombinant octamer (2 μg) was used as substrate for 25, 50 and 100 ng of wt (lanes 1–3) and H1536K mutant (lanes 4–6). The corresponding autoradiograph is shown in the lower panel. (E) In vitro methylation of 2 μg of recombinant H3 (lane 1), H3 mutated at lysine 9 (lane 2), H3 mutated at lysine 27 (lane 3) or both (lane 4) using dG9a and a mock purification. Coomassie stained gel is shown in the upper panel and a corresponding autoradiograph in lower panel. A corresponding filter binding assay is shown to the right. The y-axis displays the percent radioactivity incorporated on 2 μg of histone H3 and H3 mutants K9A, K27A and K9/K27A with radioactivity incorporated on wt H3 set to 100% and the background is subtracted. (F) HKMTase activity of mG9a on histone H3 molecules and H3 K9A, H3 K27A and the double mutant K9A/K27A. A gel of histone H3 and H3 mutants K9A, K27A and K9/K27A with radioactivity incorporated on wt H3 set to 100% and the background is subtracted. (G) Amino acid sequence of the H4 N-terminus is shown at the top. The N-terminus of the histone H4 primary structure is shaded. The corresponding autoradiograph is shown in the lower panel. A corresponding filter binding assay is shown to the right. The y-axis displays the percent radioactivity incorporated with activity on wt H3 set to 100% and the background is subtracted. (H) Quantitative MALDI-TOF analysis of 500 ng of H3 and H4 methylated by 100 ng of dG9a. Peptides spanning amino acids 9–17 and 27–40 of H3 and 4–17 of H4 is represented by graphs. No signals were observed in other peptides. Mono-, di- and trimethylation are shown as percent of total H3 or H4. This figure is representative for at least three different methyltransferase assays.
dG9a is expressed throughout development

Immunostaining revealed that the dG9a protein is found throughout oogenesis, embryogenesis, and larval development. During these stages, large cells (like nurse cells and salivary gland cells) are metabolically very active, and having multiple copies of genes (polyteny) permits a high level of gene expression; that is, abundant transcription and translation to produce the gene products. (53). In adult flies the dG9a transcript and protein are solely found in the gonads, where cells are undergoing extensive endo- and mitotic replication. One can assume that is important that certain genes are kept silent in these cells, and one possible function of dG9a could be to maintain repression of a subset of genes in cells that otherwise have a high gene expression level.

In the RNAi knock down studies, no lethality was observed during embryogenesis, but this can be ascribed to the fact that the RNAi construct was made using a pUAST based vector that is defect in the germline during oogenesis (22). Conditional knock down of dG9a in the female germline was therefore not possible in this study, but should be subject for future investigation. An interesting observation is that the escapers from the RNAi studies are exclusively females. This result suggests that dG9a may have different roles in males versus females. It is possible that conditional depletion of dG9a in transgenic flies may affect the expression of genes that are required for chromatin stability, chromosome segregation and proper histone modifications resulting in a preferential lethality in male flies. This has recently been reported for Su(var)205 (also called HP1) (54), and bonus (bon), encoding a homolog of the vertebrate TIF1 transcriptional cofactors and required for male viability (55). Interestingly, bon is associated with genes that are implicated in the ecdysone pathway.

dG9a is involved in ecdysone mediated signaling

Next, we provide evidence that dG9a is required for important transitions during Drosophila development. Our results suggest a role for dG9a in regulation of genes, especially during the onset of metamorphosis, and wing development, processes tightly correlated to ecdysone responsive signaling (56). Additional evidence for dG9a being involved in the ecdysone hierarchy is the formation of melanotic tumors in the larvae that do not form their puparium. Several chromatin-modifying or chromatin-associated complexes (57) as well as ecdysone have been implicated in hemocyte development and melanotic tumor formation (58,59). Furthermore, our genetic studies revealed that the EcR<sup>554fs</sup> is able to dominantly affect the wing phenotype in ap-GAL4 UAS-dG9a/IREcR<sup>554fs</sup> flies. Genetic interactions between mutant alleles of different genes are indicative of these genes belonging to the same functional pathway. Thus, the genetic studies support the results from the RNAi experiments, and together provide strong evidence of dG9a being involved in the ecdysone signaling hierarchy. As the RNAi mutants are not rescued by hormone feeding, dG9a must exert its effect downstream of ecdysone biosynthesis and metabolism. One possible scenario is that dG9a acts as a co-regulator for the ecdysone receptor mediating downstream gene regulation as a response to ecdysone pulses. A similar scenario has been reported for mammalian G9a, where a reduction of endogenous G9a reduced hormonal activation of an endogenous target gene by the androgen receptor (60).
The genetic interaction between EcR<sup>Δ554fs</sup> and ap-GAL4,UAS-dG9a-IR on wing development may suggest a molecular interaction between the EcR receptor and dG9a. Activation and repression of transcription involve the recruitment of many co-regulator (co-repressor or co-activator) proteins to the regulated gene promoter by sequence-specific DNA binding transcription factors. As dG9a contains an AT-hook, it could tether the ecdysone receptor to the DNA, or, more plausible, the DNA binding activity of EcR by could bring dG9a to the promoter. Two models could explain the EcR-dG9a relationship observed:

(i) dG9a act as a co-repressor of the early puffs according to the Ashburner model for the hormonal control of polytene chromosome puffing (61). Briefly, this model proposed that ecdysone, bound to its specific receptor, directly induces the expression of a small set of early regulatory genes. The protein products of these genes, in turn, repress their own expression and induce a much larger set of late target genes. dG9a could be involved in this repression.

(ii) dG9a act as a co-activator coupled to the transcription apparatus during activation of ecdysone regulated genes.

Vakoc and co-workers (62) reported recently that H3–K9 methylation was found at high levels in the transcribed region of four genes while they were transcribed. This observation is rather remarkable in that it implies a coupling of the traditionally accounted H3–K9 silencing mark to active transcription. Therefore, the possibility that dG9a plays a role in maintaining transcription should be further investigated. In addition, there are observations that murine G9a acts both as a co-repressor (63–65), and a co-activator (60), depending on promoter context and/or regulatory environment, along with the observation that the zinc finger protein wiz links G9a/GLP histone methyltransferases to the co-repressor molecule CtBP (66). Furthermore, NSD1, which methylates both H3–K36 and H4–K20 by sequence-specific DNA binding transcription factors. As this repression.

As a complement to the RNAi approach, we have tried to generate null mutants (deletions) by re-mobilization of the P-element inserted in the 5′-untranslated region (5′-UTR) of dG9a in the dG9a<sup>13414</sup> stock. Whereas several independent lines with precise excision of the P-element were obtained we failed to find any imprecise excision (deletion) events (M. Stabell, unpublished data). During the course of the preparation of this manuscript, Mis et al. (68) also identified CG2995 as being the Drosophila homolog of mammalian G9a. This group also reported unsuccessful mobilization of the P-element, and suggested that this may be due to a defective P-element. Instead, they investigated the dG9a<sup>13414</sup>/dG9a<sup>13414</sup> mutant and report only a minor phenotype without characterizing the nature of the mutant. On the other hand, they showed that this dG9a<sup>13414</sup> mutant suppresses position effect variegation (PEV) and that it interacts genetically with Su(var)3-9, suggesting that the two proteins have an overlapping role in heterochromatic gene silencing and may be members of protein complexes involved gene silencing. In contrast to Mis et al. (68) who concluded that dG9a is a H3–K9 HKMTase, we provide evidence that dG9a (i) methylates H4 as well as H3, (ii) is able to add three methyl groups, (iii) methylates K9 and K27 on histone H3 with a preference for K9 and (iv) has a specificity towards K8, K12 or K16 on the H4 N-terminus. In polytene chromosomes dG9a is excluded from the chromocenter (Figure 3), indicating a euchromatic role for dG9a. But as the majority of full-length GFP-mG9a fusion proteins has been found in pericentric heterochromatin (16), we cannot rule out a conceivable function for dG9a during facultative heterochromatinization in other tissues and/or stages of development.

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