Processive DNA Demethylation via DNA Deaminase-Induced Lesion Resolution

Don-Marc Franchini\(^1\,\,2\), Chun-Fung Chan\(^3\,\,3\text{a}\), Hugh Morgan\(^3\,\,3\text{b}\), Elisabetta Incorvaia\(^1\), Gopinath Rangam\(^1\,\,2\), Wendy Dean\(^3\), Fatima Santos\(^3\), Wolf Reik\(^3\,\,4\), Svend K. Petersen-Mahrt\(^1\,\,2\)

\(^1\) DNA Editing in Immunity and Epigenetics, IFOM-Fondazione Istituto FIRC di Oncologia Molecolare, Milano, Italy, \(^2\) 2DNA Editing Lab, Clare Hall Laboratories, London Research Institute, South Mimms, United Kingdom, \(^3\) Laboratory of Developmental Genetics and Imprinting, The Babraham Institute, Cambridge, United Kingdom, \(^4\) Centre for Trophoblast Research, University of Cambridge, Cambridge, United Kingdom

Abstract

Base modifications of cytosine are an important aspect of chromatin biology, as they can directly regulate gene expression, while DNA repair ensures that those modifications retain genome integrity. Here we characterize how cytosine DNA deaminase AID can initiate DNA demethylation. In vitro, AID initiated targeted DNA demethylation of methyl CpGs when in combination with DNA repair competent extracts. Mechanistically, this is achieved by inducing base alterations at or near methyl-cytosine, with the lesion being resolved either via single base substitution or a more efficient processive polymerase dependent repair. The biochemical findings are recapitulated in an in vivo transgenic targeting assay, and provide the genetic support of the molecular insight into DNA demethylation. This targeting approach supports the hypothesis that mCpG DNA demethylation can proceed via various pathways and mCpGs do not have to be targeted to be demethylated.

Introduction

Cytosine methylation and its oxidative variants are important mammalian DNA modifications, playing key roles for the maintenance of genomic stability and cellular identity by controlling gene expression, genomic imprinting, X chromosome inactivation, and silencing of transposable elements \cite{1,2}. Concurrently, DNA demethylation is required during early development, in somatic cells during differentiation, and for cellular reprogramming, stressing the reversible nature of DNA methylation \cite{2,3}. Yet, while the process of establishing and maintenance of DNA methylation by DNA methyltransferases (DNMTs) is well characterized, understanding the molecular mechanism underlying active DNA demethylation is only in its beginnings \cite{3}. In mammals, proposed mechanisms for DNA demethylation involve modification of the methylcytosine (5mC), followed by DNA repair dependent cytosine (dC) substitution \cite{1,2,3,4,5,6}. 5mC-modifying enzymes include hydroxylases and deaminases. The Ten-eleven translocation (TET) family proteins hydroxylate 5mC \cite{7,8}, with the resulting hydroxymethylcytosine (5hmC) either oxidized \cite{8,9} and/or replaced with cytosine - possibly via base excision repair (BER) \cite{10}. Proteins of the activation-induced deaminase (AID)/apolipoprotein B mRNA-editing enzyme complex (APOBEC) family can deaminate 5mC to thymine (dT) \cite{6}, creating a dT:dG mismatch, which can be repaired back to a cytosine via the BER pathway \cite{11,12}. Although there is a debate on the extent of AID’s involvement in DNA demethylation, a number of publications have identified important genetic links. These include DNA methylation alterations in zebrafish after addition and removal of AID \cite{12}, loss of global DNA demethylation in AID deficient mice \cite{13}, lack of complete reprogramming from AID deficient cells during heterokaryon fusions \cite{14}, and inefficient iPSC formation from AID -/- cells \cite{15}. On the other hand, from the recent literature it is clear that other enzymes or pathways (e.g. GADD45 or Nucleotide Excision Repair - NER) are also involved in DNA demethylation \cite{4,5}, hence it is unlikely for AID (or family members) to be responsible for all observed DNA demethylation events.

AID was initially described as being essential for the diversification of immunoglobulin (Ig) genes. In activated B cells AID deaminates, on single stranded DNA (ssDNA), cytosine residues to uracil (dU) \cite{16}. Depending on multiple factors, such as the cell...
state, chromatin state, and location, the dU lesions will lead to repair, or point mutations and DNA recombination. In the Ig locus, dUs are necessary to induce antibody affinity maturation via somatic hypermutation (SHM), and change of antibody effector functions via class switch recombination (CSR) [17,18]. The exact molecular mechanisms are beginning to be revealed, with proteins from DNA repair pathways [including BER and mismatch repair (MMR)] playing a necessary role during SHM [19,20,21].

Here we begin to delineate the molecular mechanisms of AID-induced lesion processing - in vivo and in vitro - leading to repair and DNA demethylation. We utilized our recently developed GAL4 targeting in an in vitro Xenopus egg extract system [22] as well as a transgenic mouse approach. DNA demethylation was analyzed either from controlled methylated plasmids (in vitro) or the differentially methylated region (DMR) of the imprinted H19 locus, dUs are necessary to induce antibody affinity maturation via SHM [19,20,21].

Materials and Methods

Generation of transgenic mice

All experimental procedures were conducted under licenses by the Home Office (UK) in accordance with the Animals (Scientific Procedures) Act 1986. The AID coding region up to 510 bp (excluding the nuclear export sequence at 550 to 596 bp at the C-terminus) was amplified from oocyte cDNA using primers BamHI-AIDF (GGATCCATGACAGCGTTCGATGAAAGCAA) and EcoRI-AIDR (GAATTCGAGATCTGACGGCCCTTCC-CAG) to generate BamHI-AID-EcoRI PCR product. This was inserted into a vector containing the CMV promoter driving the GAL4 DNA binding domain [23] (a kind gift from François Faks, Free University of Brussels, Belgium) to generate CMV GAL4-AID plasmid. Plasmids CMV GAL4-AID1 and CMV GAL4-AID2 were derived from CMV GAL4-AID by in vitro mutagenesis. Inserts were excised from the plasmids by NcoI and DdeI restriction enzyme digestion, purified (Qiagen), and used for microinjection into F1 (C57Bl/6J x CBA/CA) x F1 pseudopregnant mothers. Genotyping of transgenic mice was carried out by PCR on DNAs from tail biopsies, using primers GAL4AID (s) GTCCAGTGAGCAGGAGGTG and GAL4AID (as) CCAAGGAAACCCGAACTGCA. PCR reactions without reverse transcriptase were always done in reverse transcriptions with 0.75 U UNG inhibitor (UGI - New England Biolabs) to prevent amplification of endogenous aid gene.

Expression Vectors and protein expression

Human His-Tagged GAL4-AID coding vector was constructed by inserting the DNA-binding domain of GAL4 into the Nol restriction site of the pET30 derived vector encoding for AID with a C-terminal His tag (described in Morgan et al. [6]). Site-directed mutagenesis was used to create catalytic inactive AID C89R, Wt and mutant GAL4-AID proteins were prepared as described for untagged AID [25]. Mouse GAL4-AIDmC was cloned the same way and its mutation activity tested in a standard rifampicin mutation assay [25].

DNA Methylation Assays

Bisulfite sequencing of genomic DNA. gDNA from tissues were digested with Nol restriction enzyme, alkaline denatured, and treated with bisulfite as described [24]. All three regions [Bi-2, -3, -4] were amplified by the use of a nested PCR strategy. Sequences of primers are in Table S1. The PCR products were cloned (TA Cloning Kit, Invitrogen) and sequenced using the Applied Biosytems sequencing system.

Expression Vectors and protein expression

Human His-Tagged GAL4-AID coding vector was constructed by inserting the DNA-binding domain of GAL4 into the Nol restriction site of the pET30 derived vector encoding for AID with a C-terminal His tag (described in Morgan et al. [6]). Site-directed mutagenesis was used to create catalytic inactive AID C89R, Wt and mutant GAL4-AID proteins were prepared as described for untagged AID [25]. Mouse GAL4-AIDmC was cloned the same way and its mutation activity tested in a standard rifampicin mutation assay [25].

In vitro resolution assay (IVR)

The details of this assay have been published [22]. Briefly: o. 5 µg of GAL4-AID (wt or mutant) were incubated with 0.1 pmol of the plasmid at 37°C for 30 min in buffer IVR. The target plasmid was either unmethylated or in vitro methylated with the M.SssI methyltransferase (New England Biolabs). The repair reaction was performed by adding 150 µg of Xenopus laevis egg extract (FE) [26] complemented with 5 µg aphidicolin (Sigma-Aldrich), 0.05 mM dNTP (without dCTP or dATP), and 0.05 mM biotinylated-dCTP or biotinylated-dATP, and incubated at 23°C for 30 min. When specified, 0.75 U UNG inhibitor (UGI - New England Biolabs) was added to the FE prior to plasmid addition.
Treated DNA was purified via Qiagen Mini-prep (saved as input), isolated on streptavidin magnetic beads (Invitrogen-Dynal), collected in 100 μl TE, and 2 μl of this bead mixture subjected to real-time PCR. PCR reactions (20 μl) contained 10 μl of the LightCycler 480 SYBR Green I master mix (Roche Applied Sciences) and primers (1512 GGCCTAATCCTGCGGTTAC Rev -1518 GTCCACCTGCGATATGTC). The reaction was monitored in a LightCycler 480 Real-Time PCR System (Roche Applied Sciences); with the ‘input’ DNA analyzed in parallel as reference. Ct values for the biotinylated-DNA were correlated to the Ct values for the input DNA. Results were presented either as relative (fold-change) or absolute (% of input) quantification [22]. For fold-change, all samples were correlated to their input and then the FE alone sample (or another specified sample) was used as reference and set to one. Alternatively, in the % of input analysis the Ct, qPCR values of input and output were converted to an absolute amount of DNA based on a standard curve, with the amount of isolated biotinylated-plasmid being expressed as a percentage of the initial amount of plasmid (input).

Statistical analysis

We performed a paired two-tailed Student T-test on the values for the bisulfite analysis and ΔΔCt analysis of AID-induced m6A-demethylation. Significance was indicated for p<0.05. We also performed chi-squared analysis of the bisulfite data. Here, the untreated (G-AID + FE) demethylation were set as the expected and the treated (G-AID + FE plus Ugi) as the observed. Chi-squared values were obtained using the Yates correction for using only one degree of freedom. Significance was indicated for p<0.01.

Results

In vitro AID-induced lesion repair

Although it is not fully clear to what extent AID is involved in DNA demethylation, we want to delineate the molecular mechanism of DNA demethylation from DNA deaminase-induced lesions. For this we are utilizing both an in vitro as well as an in vivo approach. The in vitro assay allowed us to obtain qualitative and quantitative readouts, while at the same time providing a means to control all aspects of the AID-induced DNA demethylation. While the transgenic mouse system recapitulated the in vitro results and provided important genetic insight into AID-induced demethylation in vivo.

To develop an in vitro DNA demethylation system, we modified our in vitro resolution (IVR) assay [Figure S1 and [22]]. Briefly, a bacterially produced GAL4-AID (G-AID) fusion protein was targeted to GAL4 DNA-binding sites (UAS) on a methylated (SsoI) supercoiled plasmid. Once bound, AID deaminated dCs to dUs in our in vitro reaction time of AID (no GAL4) and plasmid was extended AID had reduced biotin incorporation, which was alleviated once the incubation time of AID (no GAL4) and plasmid was extended [22]. This indicated that the GAL4 domain simply enhanced the molecular crowding aspect of the reaction by allowing AID to find its DNA target more efficiently. Since our assay was designed to understand the mechanisms downstream of AID-induced lesions, the GAL4 DNA binding domain only served as a means to enhance the speed and accuracy of the reaction but not the targeting. The in vivo mechanism of AID-targeting had been addressed previously by us [28], and we did not want to add this extra complexity to a biochemical and mechanistic analysis of post-AID lesion events.

From our work on unmethylated substrates for AID-induced damage resolution [22], we were able to demonstrate that the lesions stimulate various DNA repair pathways. They included BER (BP-BER and LP-BER) and MMR (non-canonical), which directly recapitulated usage of those DNA repair pathways that arise from AID-induced lesions during immunoglobulin diversification in vivo [19].

IVR development for AID-induced demethylation

A current model for AID-induced DNA demethylation suggests that the T-G mismatch, created by AID upon deamination of 5mC, is processed via the BER pathway [6,11,12]. To better understand which DNA repair pathway is involved, we modified our IVR assay using a methylated substrate (Figure S1), where the plasmid DNA was fully methylated at 5mCpG with M.SsoI (Figure S2). Because M.SsoI treatment can introduce DNA damage itself, stemming from lack of the co-factor SAM or buffer composition, utilization of various mock treatments (Figure S2A) demonstrated that the observed IVR result (Figure S1) was only dependent on the AID activity acting on the methylated substrates. The M.SsoI treatment did not change plasmid topology significantly (Figure S2B), nor did it alter the efficacy of the AID-IVR assay (Figure S2C). As seen in Figure 1A, just as for the unmethylated substrate [22], the methylated plasmid was acted upon by catalytic G-AID and repaired in the IVR. Neither the catalytic dead AID (G-AIDm C87R - [29]), the GAL4 DNA binding domain (G-DBD), nor the untargeted AID (AID) were able to induce DNA repair on the methylated plasmid.

BER and other DNA repair pathways act during DNA demethylation

In our related work on unmethylated substrates [22], we were able to demonstrate that the AID-induced DNA damage can be resolved via various DNA repair pathways using the IVR system. Applying the same approach here, we demonstrate that with a methylated plasmid different types of DNA repair pathways are acting on the AID-induced lesions. When the BER inhibitor Ugi...
was added to the IVR, we observed a reduction without complete inhibition (Figure 1B). Although the use of Ugi could have minor side effects, the peptide has been extensively studied and characterized and shown remarkable specificity for inhibiting UNG2 [the predominant BER protein acting on dUs] [30,31] without affecting other uracil DNA glycosylases (e.g. TDG, MBD4, SMUG). This indicated that although UNG-dependent BER is important in AID-induced lesions resolution of methylated plasmids, other DNA repair pathways are also playing a role. During SP-BER the dU:dG (or dT:dG) lesion is repaired with the incorporation of a single dC, resulting in a dC:dG base pair. Processive polymerase-dependent repair pathways (e.g. LP-BER, MMR), not only resynthesize the lesion, but also incorporate nucleotides that are downstream of the initial dU. These pathways can be detected in the IVR by addition of biotinylated-dA (Figure 1B), where a significant amount of biotin incorporation can be seen after treatment with G-AID and FE. Importantly, by adding Ugi, we are able to discern between UNG-dependent LP-BER and other processive polymerase-dependent DNA repair pathways, such as non-classical MMR [21].

AID-induced single and processive DNA demethylation

Aside from identifying the various DNA repair pathways acting on the AID-induced lesions of a methylated substrate, we also determined the extent of local DNA demethylation. Using bisulfite analysis of a region downstream of the GAL4 DNA-binding site, we identified both single site demethylation as well as consecutive (processive) demethylation events (Figure 2A), with AID activity leading to 43% cytosine demethylation (Figure 2B). One should note that although the DNA is CpG methylated, unmethylated dCs outside a 5mCpG context are still substrates for AID-induced deamination. When we treated the FE with Ugi, to inhibit UNG2 dependent BER, bisulfite analysis showed a significant decrease in the efficiency of the extract to induce DNA demethylation (Figure 2A & B). This strongly suggested that AID can induce DNA demethylation by acting on dCs, since dUs are the only substrate for UNG2. Furthermore, if dCs are deaminated to dUs and UNG2 lesion processing leads to 5mCpG demethylation, then LP-BER plays a role in local DNA demethylation.

Local 5mCpG context prescribes DNA repair pathway choice for DNA demethylation

Given the multitudes of DNA repair pathways acting on AID-induced lesions, we wanted to determine if there is a preference of DNA repair pathway choice for demethylation of individual 5mCpGs. To this end, we statistically analyzed the difference of demethylation frequency of each CpG in the G-AID + FE and G-AID + FE (Ugi) samples. Using chi-squared analysis we were able to identify 3 regions (CpG 4, 14–15, and 23–24) that where, upon treatment of the FE with Ugi, significantly precluded to undergo DNA demethylation. Furthermore, some regions (CpG 7, 24, 26) had less significant inhibition of DNA demethylation, while others (CpG 1–2, 8–9, 11, 18, and 29) were refractory for the Ugi inhibition.

Overall, these results indicate that AID-induced DNA demethylation can be mediated via different DNA repair pathways, which are either UNG-dependent (5mC independent), UNG-independent (e.g. TDG), and/or are repaired beyond the initial lesion via a processive DNA polymerase.

Induced demethylation does not require CpG recognition

We have previously determined that AID deaminates unmethylated cytosines about 5–10 times more efficiently than methylated ones [6]. Furthermore, the IVR target plasmid contains 844 dCs in the context of WRC (A/T,A/G,C - the preferred AID target), yet only 161 are in the context of WRCpG. Hence given the random targeting of G-AID, it is more likely that both dU:dG and dT:dG mismatches are present on an AID treated methylated-substrate, rather than just dT:dG. This mixture also precluded the identification of a strict requirement for only 5mC deamination leading to DNA demethylation. Also, the experiments with bio-dC or/and Ugi inhibition (Figure 1B) suggested that DNA repair processed and incorporated bio-dC/bio-dA beyond the initial AID-induced lesion. All of this suggests that if a 5mC was in proximity to a deaminated dC (dU) and processive DNA polymerase repair replaced the 5mC with dC, then DNA demethylation proceeded without having initially targeted the 5mCpG.

To prove this non-targeting aspect of DNA demethylation we demonstrated that G-AID can induce methyl-adenosine demethylation on the plasmid via targeting dCs. Plasmid DNA isolated from common molecular biology grade E. coli hosts is methylated on adenosine in the context of Gm6ATC (dam methylation - m6A). In the IVR, dam methylation did not alter G-AID-induced activity on cytosines (comparing plasmids from dam (-) or (+) hosts - data not shown), indicating that the FE did not recognize m6A DNA base as a lesion. We analyzed 4 different Gm6ATC sites in the plasmid (position 468, 1287, 2001, and 3230) for the effect of AID-induced m6A demethylation (Figure 3A), with the distribution of each site on the plasmid not biasing the IVR results [22]. After G-AID and FE treatment and prior to qPCR, the plasmids were restriction-digested by either of three isochizomers recognizing GATC: Mbol (sensitive to m6A), Sau3AI (resistant to m6A), or DpnI (requiring m6A); cutting the GATC site prevents PCR amplification and reduces the signal. Therefore, m6A containing plasmids are digested (less recovery) by Sau3AI and DpnI, while they are resistant (more recovery) to Mbol cleavage. A decrease in resistance after Mbol cutting and enhanced resistance after DpnI cutting is a reflection of the incorporation of dA during AID-induced repair, and therefore demonstrates m6A demethylation.

We previously described that FE alone can induce minute amounts of DNA repair [22], and hence induce 6mA demethylation on its own. As seen in Figure S3, FE alone (no G-AID) treated substrates showed some difference between Mbol and

![Figure 1. AID-induced lesions on a methylated substrate are resolved by DNA repair.](image-url)
uncut, or DpnI and Sau3A I digestion. The ratios of the differences were then used as a correction factor to determine AID-induced m6A demethylation. In Figure 3B and 3C, uncut G-AID treated samples were set to 100% and G-AID induced demethylation represented as % recovery to uncut.

MboI reduced the efficacy of PCR amplification when compared to the uncut sample, indicating that the methylation of the four analyzed m6A sites had been lost. This AID-dependent demethylation of m6A sites was also observed when we restricted the plasmid with DpnI and Sau3A I. DpnI restriction was not as complete as Sau3A I (enhanced recovery), which also indicated a loss of m6A sites (Figure 3C).

These results clearly demonstrate that AID-induced lesions (dU) can be repaired with a processive polymerase-dependent repair system, leading to substitution of methylated adenosine by unmethylated adenosine. Therefore, DNA repair from AID-induced lesions can be sufficient to induce demethylation without directly targeting the methylated base. TET modified 5hmC is not a target for direct AID induced deamination, as its bulky side-chain on C5 does not fit the active site of AID [32], nevertheless 5hmC can be removed from DNA by an AID-dependent mechanism, if AID-induced uracils are present near 5hmC prior to processive polymerase dependent DNA repair.

In vivo targeting of AID induces local demethylation

The in vitro data suggest that lesions induced by targeting of AID to a specific locus can induce DNA demethylation. In order to determine whether the in vitro observations also hold true in vivo, we used the GAL4-AID fusion and targeting strategy in a transgenic mouse approach. Here, a female transgenic GAL4-AID fusion protein mouse (Figure 4) was bred to a male mouse having the GAL4 binding sites (UAS) introduced into the first (of four) methylated H19-DMR (H19 DMR-UAS), using a previously developed transgenic strategy [33] (Figure 5A). This DMR (differential methylated region) was chosen because the paternal allele remains stably methylated in all embryonic and somatic tissues of the offspring. For the various GAL4-AID fusions (Figure 4A), transgenes were driven by the CMV promoter and the C-terminal region of AID, including the nuclear export signal, was deleted. Although deletion of the C-terminal region can alter AID turnover [34,35], overall it can enhance nuclear localization without diminishing deaminase activity [36,37]. We also generated two mutants of the same transgene: CMV GAL4-DAD1 carries two amino acid changes (D89G & C147R) and CMV GAL4-ΔAID2 has a single amino acid change (E58G - [29]) in the catalytic domain (Figure 4A). Both of the mutant AID proteins are severely reduced in their catalytic activity (Figure S4). In all mouse lines the transgene was expressed in various embryonic and somatic tissues of the offspring.

Figure 2. AID-induced lesion repair results in demethylation. (A) AID induces demethylation of 5mCpG in vitro. Methylated plasmids from IVR reactions in Figure 1 were subjected to bisulfite sequence analysis of regions BS, 3’ of the GAL4 binding site. Methylation was monitored from IVR-samples performed with GAL4-AID (G-AID + FE) or without (FE), and in the presence of Ugi (G-AID + FE (Ugi)). White and black circles represent unmethylated and methylated cytosines, respectively. (B) Quantification of 5mC demethylated after the IVR assay from (A). Error bars indicate ± SD (n = 3); a t-test showed a significant difference between the two groups (p<0.05). (C) Graphical representation of those CpGs that were significantly inhibited by Ugi treatment for resolving AID-induced demethylation in the IVR. Each CpG (1 - 29) was analyzed using chi-squared analysis, and red circles indicate p<0.01.

doi:10.1371/journal.pone.0097754.g002
Figure 3. AID induces demethylation independent of the nucleotide context. (A) The table summarizes whether methylation of the adenosine (m6A) in GATC sites influences cutting by the restriction enzymes Mbol, Sau3AI or DpnI. On the left is the preference of cutting depending on the state of adenosine-methylation. In the middle the absolute PCR recovery after digestion is indicated. The outcome of the PCR recovery of digested DNA after AID-induced demethylating m6A is indicated on the right. (B) (C) AID induces demethylation of methyl-adenosine. Supercoiled plasmids (m6A(+)) as isolated from standard E. coli were subjected to IVR assays containing GAL4-AID as in Figure 1. Streptavidin-isolated plasmids were either undigested (mock) or incubated with the restriction enzymes Mbol, DpnI, and Sau3AI (NEB, USA) prior to qPCR amplification. The different GATC sites (positions: 468, 1287, 2001 and 3230) are indicated by colors and marked on the plasmid map; there are 37 GATC sites within the plasmid substrate. IVR results were quantitated by setting the uncut G-AID treated samples to 100%. Correction factors based on FE alone activity (for Mbol) or Fe + Sau3AI activity (for DpnI) was determined from Figure S3 and applied to the G-AID induced recoveries. Loss of m6A is shown in (B) by a decrease in recovery after Mbol cutting versus uncut, and by an increase in recovery after DpnI digest (C). A standard t-test indicated that after G-AID treatment all sites showed a significant difference between Sau3AI and DpnI cutting; (p<0.05 - indicated as *), (n=3). doi:10.1371/journal.pone.0097754.g003
postnatal tissues, as tested by RT-PCR (Figure 4B) and by immunofluorescence in zygotes (Figure 4C).

We bred males harboring the H19 DMR-UAS locus with females carrying the GAL4-AID, GAL4-DΔAID1, and GAL4-DΔAID2, which were excised from the plasmid backbone with NruI and DraIII and microinjected into zygotes, resulting in transgenic strains TG 4 and 5, TG 7, and TG 8. Lower panel: The GAL4 DNA binding domain was fused to the AID cDNA lacking the C terminal nuclear export signal (NES). In addition to wild-type AID two mutant forms of AID cDNA, harbouring amino acid exchanges D89G and C147R, and E58G, respectively, were fused. Numbers refer to amino acids position in AID. (B) GAL4-AID expression in transgenic strains TG 5 (GAL4-AID), TG 7 (GAL4-DΔAID1), and TG 8 (CMV GAL4-DΔAID2) was monitored by RT-PCR. RNA samples from embryo, placenta and various adult tissues were analyzed. Hprt mRNA served as loading control. (C) Transgenic AID localizes in pronuclei of zygotes in TG 5 (GAL4-AID) and TG 7 (GAL4-DΔAID1). GAL4-AID expression in zygotes was analyzed by immunofluorescence using an anti-GAL4 antibody. DNA was stained with DAPI. TG 5 and TG 7 resulted from crossing of transgenic mother with H19 DMR-UAS father, as a control crosses between C57Bl6 mother and H19 DMR-UAS father was used.

doi:10.1371/journal.pone.0097754.g004

Figure 4. Structure of GAL4-AID transgenes and their expression. (A) GAL4-AID fusion cDNAs were inserted into a CMV promoter containing vector resulting in three transgene constructs (GAL4-AID, GAL4-DΔAID1, and GAL4-DΔAID2), which were excised from the plasmid backbone with NruI and DraIII and microinjected into zygotes, resulting in transgenic strains TG 4 and 5, TG 7, and TG 8. Lower panel: The GAL4 DNA binding domain was fused to the AID cDNA lacking the C terminal nuclear export signal (NES). In addition to wild-type AID two mutant forms of AID cDNA, harbouring amino acid exchanges D89G and C147R, and E58G, respectively, were fused. Numbers refer to amino acids position in AID. (B) GAL4-AID expression in transgenic strains TG 5 (GAL4-AID), TG 7 (GAL4-DΔAID1), and TG 8 (CMV GAL4-DΔAID2) was monitored by RT-PCR. RNA samples from embryo, placenta and various adult tissues were analyzed. Hprt mRNA served as loading control. (C) Transgenic AID localizes in pronuclei of zygotes in TG 5 (GAL4-AID) and TG 7 (GAL4-DΔAID1). GAL4-AID expression in zygotes was analyzed by immunofluorescence using an anti-GAL4 antibody. DNA was stained with DAPI. TG 5 and TG 7 resulted from crossing of transgenic mother with H19 DMR-UAS father, as a control crosses between C57Bl6 mother and H19 DMR-UAS father was used.

doi:10.1371/journal.pone.0097754.g004

postnatal tissues, as tested by RT-PCR (Figure 4B) and by immunofluorescence in zygotes (Figure 4C).

We bred males harboring the H19 DMR-UAS locus with females carrying the GAL4-AID, GAL4-DΔAID mutants, or the previously described CMV GAL4-Myc [33] expressing transgenes, and determined the extent of DNA methylation in F1 offspring. Due to technical limitation of obtaining enough material from fertilized oocytes we could not perform bisulfite analysis on homogenous tissues right after fertilization (zygote). Hence, we choose to analyze tissue samples for methylation analysis of various regions surrounding the UAS (Bi-2, -3, -4) from neonatal liver (Figure 5). Since adult liver did not express the transgene itself (Figure 4B), it was likely that any observed demethylation had to occur in earlier stages of development. The H19 locus is an imprinted locus, with the paternal allele being methylated and the maternal allele unmethylated. Due to the genetic manipulations of the system regions Bi-2 & 3 can be amplified from the paternal allele, while region Bi-4 can only be amplified from the maternal allele (Figure 5B). As shown in Figure 5C, Bi-2 was significantly more demethylated in GAL4-AID than in GAL4-Myc mice. More importantly, the demethylation required AID catalytic activity, as transgenic mouse 7 (TG 7 - harboring a catalytic inactive GAL4-AID) did not show extensive DNA demethylation in this region. Bisulfite analysis of the Bi-3 region confirmed the results for the demethylation capacity of a catalytic active AID (Figure 5D and summarized in Figure S5A), where TG 4 and TG 5 induced over 95% demethylation. Loss of the methylation on the paternal allele may induce methylation on the maternal allele – possibly via dosage compensation [38]. Yet targeting of AID to the paternal allele did not influence the DNA methylation status on the maternal allele, since bisulfite analysis of Bi-4 showed no change in any of the mice analyzed (Figure 5E and Figure S5B). The paternal DMR DNA methylation status (Bi-2 and Bi-3) was also analyzed from embryos and placenta (Figure 6 and Figure S5C),
Figure 5. Targeting of GAL4-AID to the H19 DMR-UAS leads to demethylation. (A) Schematic of the paternal knock-in H19 DMR-UAS mouse and the maternal transgene GAL4-AID (where GAL4 is in yellow and AID in salmon) mouse. GAL4-AID transgenic females were crossed with H19 DMR-UAS homozygous male. (B) Methylation of the loci surrounding the UAS was analyzed by bisulfite sequencing. The regions analyzed by bisulfite sequencing (Bi-2, Bi-3, Bi-4), the CTCF binding sites (blue lines), the G repeats sequence, the transcription start site (black arrow) and the position of the UAS within the H19 DMR are depicted. (C–E) AID-induced methylation alterations at the Bi-2 (C) and Bi-3 (D), but not at Bi-4 (E) regions. DNA methylation of H19 DMR alleles was analyzed by bisulfite sequencing of DNA from neonatal liver (L) of transgenic offspring of GAL4-AID wt (TG 4 and TG 5), mutant GAL4-ΔAID (TG 7), or GAL4-myc (TG 1) crossed with H19 DMR-UAS. The top of the figure (C) represents the schematic of the context of the CpGs within the Bi-2 sequence: CpG with an AID hotspot motif WRC (in red, WRCpG) or in the cold spot motif SYC (in green, SYCpG). Filled circles represent methylated CpGs, open circles unmethylated ones. GAL4 is represented by a yellow triangle, AID wt by a salmon circle, AID mutant by a misformed salmon circle, and Myc by a green rectangle. Summary of the results are shown in Figure S5.

doi:10.1371/journal.pone.0097754.g005
and analogous to the results from the liver tissue, catalytic AID induced local DNA demethylation.

AID-induced demethylation outside its target motif

We estimated the extent of AID-induced DNA demethylation of at least 1,000 bases at the H19 locus, as the paternal specific Bi-2 and Bi-3 are each about 500 bases long and were substantially demethylated. The upstream border of DNA demethylation could be situated near the 5' part of the Bi-2 (further upstream are no polymorphisms to distinguish between maternal and paternal alleles); the downstream border is likely the G-repeats, inhibiting the DNA demethylation.

AID favors the hot spot motif WRC (A/T,A/G,C) and strongly disfavors SYC (cold spot - G/C, C/T, C) sequences for deamination [27], which is even more pronounced on a 5mC containing substrate [6]. Sequence context analysis of the Bi-2 demethylated CpGs showed that there was no difference in targeting cold spots or hot spots (Figure 5C; cold spot - green, hot spot - red), indicating that it is unlikely for AID to have targeted each and every CpG. During immune diversification, AID can be linked to the SP-BER pathway [16], while we [6] and others [11,12,39] have implicated SP-BER with AID-induced demethylation. Yet, this single 5mCpG targeting pathway has its limitations. The AID-induced demethylated Bi-2 & Bi-3 regions contained 88 5mCs among 582 dCs. In order for AID to target each CpG separately (followed by faithful DNA repair), 5279 independent deaminations would have had to occur (Figure S6), with each lesion being repaired by SP-BER. It is more likely that in vivo DNA deamination induced DNA demethylation proceeds via a processive DNA polymerase repair pathway (e.g. LP-BER or MMR), allowing for multiple CpG demethylations to occur from a single DNA lesion. Overall, these data indicated that the GAL4-AID-induced demethylation occurred early in mouse development and involves both processive (initiated from a single AID-induced lesion but leading to multiple demethylation events) and non-processive (one lesion one demethylation) DNA demethylation activity.

Discussion

DNA instability plays a pivotal role in survival and evolution, with physical and chemical DNA damage being mutagenic, while at the same time controlled DNA alterations provide adaptation to environmental stress, either during meiosis or the development of the adaptive immune system. DNA methylation is the sum of DNA methylation and active/passive demethylation [2,3,40], with active removal of the methyl-mark requiring DNA base modification or removal of the modified-cytosine. The molecular processes activated to induce DNA demethylation will determine the extent and efficiency of the epigenetic change. Recent data have shown that single base demethylation can be achieved via different pathways [5,6,8,11], while our current work proposes that a more efficient way for DNA demethylation can occur. By utilizing different DNA repair pathways, either UNG dependent or independent (coupled to processive DNA polymerizations), a single DNA lesion could induce multiple DNA demethylation events (Figure 7). At a targeted locus AID activity on cytosine (regardless of its methylation status) would induce DNA mismatches (dU:dG or dT:dG). dT:dG processing via BER would lead to a single DNA demethylation, while dU:dG processing via BER leads to the status quo. On the other hand, DNA repair utilizing a processive DNA polymerase (e.g. MMR or long-path BER) would lead to extensive removal of 5mC marks from a single lesion. MMR is known to replace up to 2 kb of ssDNA away from the lesion itself [41,42], and if taking place near 5mCpGs then each 5mC would be replaced by cytosine within this stretch. This model would reduce the number of DNA damages required to induce DNA demethylation (Figure S6), increase efficiency and accuracy, and be in line with known AID-induced DNA repair pathways. Recent analysis of DNA methylation status immediately post-fertilization suggested processive DNA demethylation (possi-
bly via LP-BER) to occur in the second phase of active DNA demethylation [43], mimicking our in vitro and in vivo findings and supporting our insights into the mechanisms of active DNA demethylation. Future work will determine if the findings form our model system and its DNA repair pathway choice will also be observed as pathways for AID induced demethylation in vivo.

The design of the IVR assay and in vivo targeting precludes analysis of how AID reaches its target as the GAL4 binding capacity vastly exceeds that of AID. In the B cell community it is well known that targeting of AID to the Ig locus is insufficient to explains AID’s ability to induce a mutation rate that is 10^6 fold above background mutations [19,28,44]. Even within the same cell an AID lesion can be repaired or lead to mutations depending on the chromatin context [45]. Hence the cellular milieu and chromatin context can be instructive for AID’s effectiveness. Both of these aspects can be carefully controlled in our IVR assay and we are currently developing new tools and readouts to gain more insight. We are also looking at the mechanistic differences of how AID’s activity in early developmental stages can lead to DNA demethylation [13,43] or how AID can alter epithelial-mesenchymal transition (EMT) [46]. Interestingly, in B cells where AID is to induce DNA mutations AID seems to have less of an influence on the dynamics of DNA methylation [47].

Insights from novel DNA damage resolution assays

Our systems allowed us to dissect the molecular mechanisms of DNA lesion resolution though genetics and biochemistry. Unlike previous in vitro systems for studying AID lesion resolution [48,49], the IVR system utilizes the physiological DNA damaging activity of AID, thereby providing the first biochemical approach to study multiple aspects of AID-induced demethylation. Addition of Ugi and bio-dA to the IVR allowed DNA damage resolution to proceed either through the UNG-dependent BER pathway (Ugi sensitive) or through a processive DNA polymerase dependent repair pathway (MMR-like pathway - dA incorporation). This lesion channelling (BER vs MMR-like) is analogous to the in vivo DNA repair pathway choice of Ugi addition during Ig diversification [19]. The IVR systems results were recapitulated in the in vivo AID targeting work and provided additional evidence that DNA demethylation can proceed via various DNA repair pathways.

Aphidicolin is predominately used to inhibit replicative polymerases, but can also inhibit DNA pol delta, a polymerase associated with MMR. Therefore, the use of aphidicolin during the resolution phase (in the presence of Ugi) of the IVR suggests that the AID-induced dU:dG mismatch is not resolved via classical MMR. Interestingly, it has been suggested that AID-linked MMR activity during Ig diversification proceeds via a noncanonical MMR pathway [21], raising the possibility that AID-induced DNA demethylation can also involve non-classical DNA repair pathways. This possible ‘high-jacking’ of classical DNA repair factors for non-classical functions also occurs during AID-induced SHM [19]. A link of this interplay has been observed in vivo, as the CpG methylation status within the Ig locus can alter SHM.

Figure 7. Model for lesion-induced DNA demethylation. Active DNA demethylation dependent on 5mC targeting. 5mC (filled circle) can be hydroxymethylated (5hmC - small grey circle) and further processed by the TET protein family (top right panel - grey). 5mC can be deaminated by the cytosine deaminase AID (bottom left panel - red). Deamination of 5mC results in thymine (dT - in red) and creates a dT:G mismatch that can be recognized by multiple DNA repair pathways, including BER or processive polymerase dependent repair. The processing of the dT:G mismatch by BER produces a single cytosine demethylation event, while processive polymerase dependent repair, in either direction, replaces long stretches of all bases, including 5mC, leading to multiple cytosine-demethylation events. Active DNA demethylation independent of 5mC targeting (central panel - green). AID targets and deaminates cytosine (white C:G), forming uracil (dU – in green) and leading to a dU:G mismatch. BER processing leads to the status quo (intact methylated DNA), while activation of a processive polymerase dependent repair, in either direction, replaces long stretches of all bases, including 5mC, leading to multiple cytosine-demethylation events. After AID and DNA repair induced demethylation on one strand, the complementary strand can be targeted as well and/or DNA replication can take place in the absence of DNMT1 activity. Active DNA demethylation of TET enzyme induced ShmC can be independent of targeting (right panel - blue). AID targets and deaminates cytosine (white C:G), forming uracil (dU – in green) and leading to a dU:G mismatch. Processing as for untargeted 5mC demethylation with the outcome of replacing ShmC with dC.

doi:10.1371/journal.pone.0097754.g007
outcome [50,51]. Future work using the IVR and AID will also allow for the uncovering of the precise molecular mechanisms of how AID-induced demethylation can proceed during primordial germ cell formation [13], during pluripotency reprogramming [14,15], and zebrafish development [12].

It is of course possible that other DNA damaging events (aside from AID-induced lesions) away from the 5mCpG, which can be repaired with processive DNA repair, such as DNA topoisomerase lesions, could serve as substrates for DNA demethylation. It is interesting to note that in vivo treatment of cells with topoisomerase inhibitors lead to DNA demethylation, rather than hypermethylation as observed for other DNA synthesis inhibitors [32].

AID and TET dependent DNA demethylation

We and others have previously shown that AID is unable to deaminate 5hmC containing substrates [32,53], and hence the proposed direct genetic links [54,55] between AID and 5hmC were only speculative. Our discovery that AID-induced lesions at non-methylated cytosines can lead to active DNA demethylation provides an alternative indirect pathway for AID to resolve TET induced 5hmC modifications. As shown in Figure 7 (far right pathway), if the AID-induced uracil is present near a 5hmC, then upon subsequent polymerase dependent DNA repair 5hmC would be replaced with unmodified cytosine. If and to what extent this pathway has an in vivo physiological role is yet to be discovered.

Conclusion

We have shown that local DNA demethylation, induced from a single DNA lesion (e.g. deamination of dG or 5mC), proceeded via at least two different efficient DNA repair pathways. We propose that this will also hold true for some aspects of global DNA demethylation, with pathway choice influencing the extent and efficiency of this mechanism. Depending on the lesion (e.g. base modification, mismatch, ssDNA nick), the genetic loci (intragenic, intergenic), the chromatin state (e.g. DNA methylation, histone modification, polycomb complex association), or cellular milieu (e.g. B cell, germ cell) different DNA repair pathways will induce lesion resolution and DNA demethylation. Therefore, a single lesion may lead to single (via BER) or multiple (possibly via processive DNA polymerase dependent repair) 5mCpG demethylation. As epigenetics, including DNA (de)methylation, is becoming more relevant for understanding oncology, this work could have a direct impact on patient care. Clinical evidence already suggest a link between DNA methylation and DNA repair for drug efficacy, therefore the IVR and in vivo targeting could provide a means to identify new drug targets.

Supporting Information

Figure S1 Schematic of IVR on methylated substrate.

Schematic description of the in vitro assay using a methylated substrate, modified from [1]. Prior to the reaction the supercoiled plasmid is in vitro methylated by using the CpG DNA methyltransferase M.SssI. The methylated (filled lollipops) DNA plasmid containing GAL4 binding sites is incubated with a recombinant Gal4-AID fusion protein creating a dU lesion (green star). The supercoiling provides a region of dsDNA for GAL4 binding and a region of ssDNA for AID activity. Addition of frog egg extract (FE), containing topoisomerase, relaxes the substrate plasmid forming a dU:dG mismatch. The repair phase in the FE is carried out in the presence of biotinylated dCTP (bio-dC) or dATP (bio-dA) - (blue arrow), along with normal dNTPs. Repaired and biotinylated DNA is isolated via magnetic streptavidin beads. Prior to streptavidin isolation a small sample (input) is removed from the reaction. Eluted products and input are then subject to quantitative real-time PCR (red bar).

Figure S2 In vitro methylation does not trigger IVR activity. (A) The plasmid was in vitro methylated with the CpG methyltransferase M.SssI. The unmethylated plasmid (UM) was incubated with the buffer ingredients only (UM Buffer), mock methylated (Mock M) with the M.SssI only (-SAM) or the cofactor SAM only (-SssI), or methylated with all the components (M). The methylation status was monitored by digestion with the methyl sensitive enzyme BstUI and analysis on a 0.8% agarose gel, post-stained with SYBR safe. (B) Topology of an in vitro methylated plasmid. 0.5 μg of all of the different plasmids obtained in (A) were electrophoresed on 0.8% agarose gel for 10 h at 5 V/cm at 4°C. After migration the gel was soaked in 1x TBE containing 0.3 mg/ml ethidium bromide for 1 h and visualized with a Gel Doc (Bio-Rad). (C) Mock methylated plasmids are equivalent to unmethylated plasmids in the IVR. Unmethylated plasmid (UM), unmethylated plasmid containing methylation buffer (UM Buffer), and mock methylated plasmids (Mock M,-SAM or -SssI) were used in the IVR assay. The bars represent the ratio of the amount of recovered plasmids from reactions carried out in the presence of G-AID versus absence of G-AID (FE alone was set to 1). Error bars indicate ± standard deviation (SD, n = 3).

Figure S3 FE activity on dam methylated adenine substrates. IVR was performed with plasmids isolated from Dam+ bacteria. Briefly, the dam-methylated plasmid was incubated with the FE and after treatment the plasmid was either mock digested (/-) or digested with MboI, DpnI and Sau3AI prior to isolation and qPCR amplification. The bars show ratio of cut versus uncut DNA (set to 1) after FE treatment.

Figure S4 Assessment of mAID protein activity. Wild type but not mutant mouse AID protein induces mutation in E. coli. The control empty vector (vector), mouse AID wild type (mAID), mouse AID mutant D89 - C147R (mutations in TG 8 line) and mouse AID mutant E58G (mutations in TG 8 line) were transformed into bacteria (BW310 - ung line), and mouse AID mutant E58G (mutations in TG 8 line) were transformed into bacteria (BW310 - ungA - [2]). Protein expression was induced with 0.4 mM IPTG for 14 h at 37°C. The bacteria were plated on low salt LB agar plates to assess viability or on rifampicin plates to determine the mutation frequency in the rpoB gene. For each sample the number of RifK clones per 10^5 viable cells is plotted [25].

Figure S5 Table summary of G-AID transgenic demethylation. (A) Table showing the summary of the DNA methylation status at the Bi-2 and Bi-3 regions after bisulfite treatment of DNA from fetal liver. Data summarized from Figure 5. The number in bracket is the number of transgenic mice analyzed. (B) Table showing the summary of the DNA methylation status at the Bi-4 region after bisulfite treatment of DNA from fetal liver. Data summarized from Figure 5. (C) Table showing the summary of the DNA methylation status at the Bi-4 region after bisulfite treatment of DNA from fetal liver. Data summarized from Figure 5. (D) Table showing the summary of the DNA methylation status at the Bi-4 region after bisulfite treatment of DNA from Embryos and Placentas (E12.5). Data summarized from Figure 6. The number in bracket is the number of transgenic mice analyzed.

Figure S6 Theoretical approximation of the number of deaminations required for demethylation based on single BER events. The transgenic mouse H19 locus is schematically drawn with the GAL4 binding sites (UAS) in the centre, surrounded by the bisulfite sequenced flanking (filled circle
- 5mC) regions. GAL4-AID (circle-triangle) is bound at the UAS with arrows representing individual DNA demethylations, leading to complete demethylation (open circle). Number of total dCs (582) and 5mC (88) in this region is indicated, followed by a set of assumptions for the calculation. After n deaminations, the probability for a single position of being never hit is (581/582) and its probability of being hit at least once is 1-(581/582)^n. Deriving a formula representing the number of deaminations that have to occur in order to have ‘hit’ 88 5mC in 582 dC with 99% confidence.

**References S1**

Primers used for bisulfite analysis. List of primers used

target of being never hit is (581/582) for the calculation. After n deaminations, the probability for a single

5mC (88) in this region is indicated, followed by a set of assumptions complete demethylation (open circle). Number of total dCs (582) and

with arrows representing individual DNA deaminations, leading to

- 5mC) regions. GAL4-AID (circle-triangle) is bound at the UAS

and is expressed in pluripotent tissues: implications for epigenetic reprogram-

ming. J Biol Chem 279: 5253–5260.

- Ko M, Huang Y, Jankowska AM, Pape UJ, Tahiliani M, et al. (2010) Impaired

hydroxylation of 5-methylcytosine in myeloid cancers with mutant TET2.

Nature 468: 839–847.

- Tahiliani M, Koh KP, Shen Y, Pastor WA, Bandukwala H, et al. (2009) Conversion of 3-methylcytosine to 5-hydroxymethylcytosine in mammalian DNA by MLL partner TET1. Science 324: 930–935.

- Ito S, D’Alessio AC, Tarazona OV, Hong K, Sowers LC, et al. (2010) Role of Tet proteins in 5mC to 5hmC conversion, E-cadherin and inner cell mass specification. Nature 466: 1129–1133.

- He Y-F, Li B-Z, Li Z, Liu P, Wang Y, et al. (2011) Tet-Mediated Formation of 5-Carboxylcytosine and Its Excision by TDG in Mammalian DNA Science.

- C佐川 D, Kono K, Seldrige J, Lettieri T, Naito Y, et al. (2013) Embryonic lethal phenotype reveals a function of TDG in maintaining epigenetic stability. Nature 470: 419–423.

- Rai K, Huggins JJ, James SR, Karpf AR, Jones IA, et al. (2000) DNA demethylation in zebrafish involves the coupling of a deaminase, a glycosylase, and a NAD. Cell 135: 1201–1212.

- Pop C, Dean W, Feng S, Cokus SJ, Andrews S, et al. (2010) Genome-wide erasure of DNA methylation in mouse primordial germ cells is affected by AID deficiency. Nature 463: 1101–1105.

- Bhutani N, Brady JJ, Damian M, Sacco A, Corbel SY, et al. (2010) Reprogramming towards pluripotency requires AID-dependent DNA demethylation. Nature 463: 1042–1047.

- Kumar R, DMello L, Schoede N, Liu TC, Franck P, et al. (2013) AID stabilizes stem-cell phenotype by removing epigenetic memory of pluripotency genes. Nature 508: 89–92.

- Petersen-Mahrt S (2005) DNA deamination in immunity. Immunol Rev 203: 90–97.

- Muramatsu M, Kinosita K, Fagasan A, Yamada S, Shinkai Y, et al. (2000) Class switch recombination and hypermutation require activation-induced cytidine deaminase (AID), a potential RNA editing enzyme. Cell 102: 553–563.

- Revy P, Muto T, Levy Y, Geissmann F, Plebani A, et al. (2000) Activation-induced cytidine deaminase (AID) deficiency causes the autosomal recessive form of the Hyper-IgM syndrome (HIGM2). Cell 102: 553–563.

- Schmitz K-M, Petersen-Mahrt SK (2012) AIDing the immune system—DIAbolic in cancer. Semin Immunol 24: 241–245.

- Liu M, Duke JL, Richter DJ, Vinuesa CG, Goodnow CC, et al. (2008) Two levels of protection for the B cell genome during somatic hypermutation. Nature 451: 841–845.

**Acknowledgments**

We thank Kristina Trenz, Claudia Cosentino and Vincenzo Costanzo for introducing us to the *Xenopus* system, as well as continued fruitful discussions; Hiro Mahbubani for technical support of *Xenopus* management; Andrea Cocito for mathematical analysis of demethylations per base-pairs; Kerstin-Maike Schmitz for critical insights, discussions, lab support, and manuscript preparation.

**Author Contributions**

Conceived and designed the experiments: DMF HM WR SKPM. Performed the experiments: DMF CFC HM EI GR WD FS SKPM. Analyzed the data: DMF CFC WD WR SKPM. Wrote the paper: DMF WR SKPM.
Molecular Mechanisms of Demethylation

46. Munoz DP, Lee EL, Takayama S, Coppe JP, Heo SJ, et al. (2013) Activation-induced cytidine deaminase (AID) is necessary for the epithelial-mesenchymal transition in mammary epithelial cells. Proc Natl Acad Sci U S A 110: E2977–2986.

47. Fritz EL, Rosenberg BR, Lay K, Mihailovic A, Tuschl T, et al. (2013) A comprehensive analysis of the effects of the deaminase AID on the transcriptome and methylome of activated B cells. Nature immunology 14: 749–755.

48. Pham P, Zhang K, Goodman MF (2006) Hypermutation at A/T sites during G/U mismatch repair in vitro by human B-cell lysates. J Biol Chem 281: 31754–31762.

49. Schanz S, Castor D, Fischer F, Jiricny J (2009) Interference of mismatch and base excision repair during the processing of adjacent U/G mispairs may play a key role in somatic hypermutation. Proc Natl Acad Sci U S A 106: 5593–5598.

50. Fraenkel S, Mostoslavsky R, Novobrantseva TI, Pelanda R, Chaudhuri J, et al. (2007) Allelic 'choice' governs somatic hypermutation in vivo at the immunoglobulin kappa-chain locus. Nat Immunol 8: 715–722.

51. Jolly CJ, Neuberger MS (2001) Somatic hypermutation of immunoglobulin kappa transgenes: association of mutability with demethylation. Immunol Cell Biol 79: 18–22.

52. Nye J, Liu L, Jones PA (1986) Variable effects of DNA-synthesis inhibitors upon DNA methylation in mammalian cells. Nucleic Acids Research 14: 4353–4367.

53. Nabel CS, Jia H, Ye Y, Shen L, Goldschmidt HL, et al. (2012) AID/APOBEC deaminases disfavor modified cytosines implicated in DNA demethylation. Nature chemical biology 8: 751–758.

54. Cortellino S, Xu J, Sannai M, Moore R, Caretti E, et al. (2011) Thymine DNA glycosylase is essential for active DNA demethylation by linked deamination-base excision repair. Cell 146: 67–79.

55. Guo JU, Su Y, Zhong C, Ming GL, Song H (2011) Hydroxylation of 5-methylcytosine by TET1 promotes active DNA demethylation in the adult brain. Cell 145: 423–434.