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Orthologous mammalian A3A-mediated singlenucleotide resolution sequencing of DNA epigenetic modification 5-hydroxymethylcytosine†

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Epigenetic modifications in genomes play a crucial role in regulating gene expression in mammals. Among these modifications, 5-methylcytosine (5mC) and 5-hydroxymethylcytosine (5hmC) are recognized as the fifth and sixth nucleobases in genomes, respectively, and are the two most significant epigenetic marks in mammals. 5hmC serves as both an intermediate in active DNA demethylation and a stable epigenetic modification involved in various biological processes. Analyzing the location of 5hmC is essential for understanding its functions. In this study, we introduce an orthologous mammalian A3A-mediated sequencing (OMA-seg) method for the quantitative detection of 5hmC in genomic DNA at singlenucleotide resolution. OMA-seq relies on the deamination properties of two naturally occurring mammalian A3A proteins: green monkey A3A (gmA3A) and dog A3A (dogA3A). The combined use of gmA3A and dogA3A effectively deaminates cytosine (C) and 5mC, but not 5hmC. As a result, the original C and 5mC in DNA are deaminated and read as thymine (T) during sequencing, while the original 5hmC remains unchanged and is read as C. Consequently, the remaining C in the sequence indicates the presence of original 5hmC. Using OMA-seq, we successfully quantified 5hmC in genomic DNA from lung cancer tissue and corresponding normal tissue. OMA-seq enables accurate and quantitative mapping of 5hmC at single-nucleotide resolution, utilizing a pioneering single-step deamination protocol that leverages the high specificity of natural deaminases. This approach eliminates the need for bisulfite conversion, DNA glycosylation, chemical oxidation, or screening of engineered protein variants, thereby streamlining the analysis of 5hmC. The utilization of orthologous enzymes for 5hmC detection expands the toolkit for epigenetic research, enabling the precise mapping of modified nucleosides and uncovering new insights into epigenetic regulation.

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Introduction

Epigenetic modifications in nucleic acids are critical regulators of gene expression in mammals. 5-Methylcytosine (5mC) represents the first and most extensively studied modified nucleobase in DNA, playing a pivotal role in fundamental biological processes including gene expression regulation and normal developmental progression.2 The discovery of 5hydroxymethylcytosine (5hmC) in mammalian genomes marked a significant milestone in epigenomics research.3,4 5hmC is generated from 5mC through an active DNA demethylation process catalyzed by the ten-eleven translocation (TET) family of dioxygenases.5-8 Beyond its role as an intermediate in DNA demethylation, 5hmC is now recognized as a distinct epigenetic mark and functionally significant nucleobase within mammalian genomic landscapes.9 Emerging research increasingly highlights the multifaceted roles of 5hmC in diverse biological processes, including tumorigenesis and embryonic development.10

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5hmC is generally linked to active gene expression and may indicate transcription that is dynamically activated. The distribution of 5hmC is specific to different tissues, with a higher prevalence in genes that promote tissue differentiation and in tissue-specific transcription factors. Additionally, 5hmC plays a crucial role in sustaining a pluripotent state in cells as well as in cell differentiation. In the context of cancer, both a widespread reduction of 5hmC and localized increases in genes were observed during tumour progression. Therefore, analyzing 5hmC offers valuable insights into epigenetic activation during the course of tumour development.

Some methods have been developed to detect 5hmC in genomic DNA, including thin layer chromatography detection,3,4 and liquid chromatography coupled with mass spectrometry analysis.14-21 Typically, these approaches involve digesting genomic DNA into nucleosides or nucleotides for qualitative and quantitative 5hmC detection.22 Understanding the functional significance of 5hmC in DNA requires precise location analysis within genomic landscapes.23 Some techniques have emerged for mapping 5hmC in DNA. Hydroxymethylated DNA immunoprecipitation (hMeDIP) using specific antibodies and affinity-based purification and sequencing methods enable whole-genome 5hmC mapping.24,25 However, these approaches are limited by their resolution, which depends on the genomic DNA fragment size, preventing identification of exact 5hmC sites.26 Oxidative bisulfite sequencing (oxBS-seq)27 and TET-assisted bisulfite sequencing (TAB-seq)²⁸ were introduced to achieve single-nucleotide resolution 5hmC detection. A limitation of these methods is the use of bisulfite, which can cause significant DNA degradation, potentially destroying up to 99.9% of input genetic material.29 Single-molecule real-time (SMRT) sequencing offers another 5hmC detection avenue,30 though it currently suffers from a high false-positive rate in modification mapping.31 More recent innovations include TET-assisted pyridine borane sequencing (TAPS), β-glucosyltransferase blocking TET-assisted pyridine borane sequencing (TAPSB), and chemical-assisted pyridine borane sequencing (CAPS), which aim to map 5hmC at single-base resolution.32,33 Nevertheless, these methods face challenges, as the dihydrouracil (DHU)-containing DNA produced can compromise amplification efficiency, potentially introducing bias in genome-wide 5hmC mapping.34

More recently, enzyme-assisted methods in mapping modified nucleobases have been developed. The A3A (apolipoprotein B mRNA-editing catalytic polypeptide-like 3A) protein, present across most placental mammals, enables deamination of cytosines to uracil in DNA. That has been reported that human A3A (hA3A) protein can efficiently deaminate cytosine (C), 5mC, and 5hmC in DNA, but shows no deamination activity toward glycosylated 5hmC (β -glucosyl-5-hydroxymethyl-2′-deoxycytidine, 5gmC). That be using this property of hA3A, we and others have reported engineered deaminase-mediated sequencing (EDM-seq), single-step deamination sequencing (SSD-seq), and A3A-coupled epigenetic sequencing (ACE-seq) for mapping 5hmC in DNA. In ACE-seq, 5hmC is first glycosylated to produce 5gmC by using β -glucosyltransferase (β -GT). hA3A treatment leads to the conversion of C and 5mC in DNA to

uracil (U) and thymine (T), respectively. Both U and T are read as T, while 5gmC resists deamination and is read as C during sequencing. These methods, however, require the pre-treatment of DNA with β -GT to glycosylate 5hmC or demand laborious screening of engineered hA3A variants.

Previous studies have demonstrated that orthologous mammalian A3A cytidine deaminases exhibit varying degrees of deamination activity toward C and 5mC. 41,50,51 Apart from hA3A, the deaminase activity of other orthologous mammalian A3A proteins toward 5hmC remained largely unexplored. In this study, we comprehensively examined the deamination activities of six orthologous mammalian A3A proteins on C, 5mC, and 5hmC, revealing two distinct types of mammalian A3A proteins with specific sequence preferences. Green monkey A3A (gmA3A, from Chlorocebus aethiops) demonstrated complete deamination of C and 5mC but showed no deamination activity toward 5hmC at YC (Y = C/T) sites. Conversely, dog A3A (dog A3A, fromCanis lupus familiaris) could fully deaminate C and 5mC but exhibited no deamination activity toward 5hmC at RC (R = A/G) sites. Leveraging these unique deamination properties of gmA3A and dogA3A, we proposed an orthologous mammalian A3A-mediated sequencing (OMA-seq) method for singlenucleotide resolution and quantitative analysis of 5hmC. OMA-seq approach introduces novel, naturally occurring enzymes for precise 5hmC mapping and quantification.

Experimental section

Chemicals and reagents

The 24-mer cytosine-containing DNA, 24-mer 5mC-containing DNA, as well as the 24-mer 5hmC-containing DNA were synthesized by Takara Biotechnology Co., Ltd (Dalian, China). Other single-stranded DNAs were purchased from Tianyi Huayu Gene Technology Co., Ltd (Wuhan, China). The sequences of these oligonucleotides are provided in Table S1.† 2'-Deoxycytidine (dC), 2'-deoxyguanosine (dG), 2'-deoxyadenosine (dA), 2'-deoxythymidine (dT), and the 2'-deoxynucleoside 5'-triphosphates, including dATP, dCTP, dGTP, and TTP, as well as venom phosphodiesterase I, were purchased from Sigma-Aldrich (Bei-5-Methyl-2'-deoxycytidine-5'-triphosphate jing, China). (5mdCTP) and 5-hydroxymethyl-2'-deoxycytidine-5'-triphosphate (5hmdCTP) were obtained from TriLink BioTechnologies (San Diego, California, USA). DNase I, S1 nuclease, and alkaline phosphatase were purchased from Takara Biotechnology Co., Ltd. Lung cancer tissue and the matched adjacent normal tissue samples were collected from Zhongnan Hospital of Wuhan University. All experiments were conducted in compliance with the guidelines and regulations established by the Ethics Committee of Wuhan University.

In vitro expression and purification of orthologous mammalian A3A proteins

The sequences for orthologous mammalian A3A proteins are obtained from the NCBI database and are listed in Table S2.† For the expression and purification of the human A3A (hA3A) protein, the coding sequence was incorporated into the pET-

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41a(+) plasmid, which was synthesized *de novo* by TsingKe Co., Ltd (Beijing, China). This plasmid includes a human rhinovirus 3C protease (HRV 3C) site between the glutathione *S*-transferase (GST) tag and the hA3A protein sequence. The resulting plasmid, pET-41a(+)-hA3A, was transformed into the *E. coli* BL21(DE3) *pLysS* strain (Sangon). The detailed procedures for the *in vitro* expression and purification of hA3A could be found in the ESI.† The purity of the protein was assessed by SDS-PAGE, and was stored at -80 °C after the addition of 25% glycerol. The concentration of the purified protein was quantified using the BCA protein assay kit (Beyotime). The expression and purification of other mammalian A3A proteins, including cowA3A, pandaA3A, porpoiseA3A, gmA3A, and dogA3A, were conducted in a similar manner to that of hA3A.

Preparation of DNA with C, 5mC, and 5hmC modifications

The 216-bp double-stranded DNA (dsDNA) substrates, including DNA-C, DNA-5mC, and DNA-5hmC, were prepared for the sequencing method development. Detailed sequences are provided in Table S3.† Briefly, a 228-bp DNA (Table S3†) was synthesized and inserted into the pET-41a plasmid (pET41a-DNA-C) using SpeI and XhoI cloning sites (TsingKe). DNA-C was generated by PCR amplification with 2 ng of pET41a-DNA-C serving as the template. The PCR amplification was conducted in a 50 µL reaction mixture containing 2.5 U of Taq DNA polymerase (Accurate), 1× reaction buffer, 0.2 mM each of dATP, dGTP, TTP, and dCTP, 0.4 μM forward primer (5'-AGTGACGCT-GAGCTTGACGTCGCGC-3'), and 0.4 μM reverse primer (5'-CCAA-CATTCCACTAACAATTACTCTCT-3'). The PCR program included 95 °C for 5 min, 30 cycles of 95 °C for 30 s, 58 °C for 30 s, 72 °C for 30 s, followed by 72 °C for 5 min. DNA-5mC and DNA-5hmC were prepared similarly, with dCTP substituted by 5mdCTP and 5hmdCTP, respectively. The PCR products were verified by agarose gel electrophoresis and purified using KAPA Pure beads (Roche).

Deamination assay for dsDNA carrying various modifications

DNA-C, DNA-5mC, and DNA-5hmC were used to evaluate the deaminase activity of orthologous mammalian A3A proteins toward C, 5mC, and 5hmC. Typically, 60 ng of the dsDNA substrate was denatured by adding 2 µL of DMSO and heating at 95 °C for 10 min, followed by immediate cooling in ice water. The deamination reaction was conducted in a 20 µL solution containing the denatured DNA, 10 µM of the orthologous mammalian A3A protein, 20 mM MES (pH 6.5), and 0.1% Triton X-100. The reaction proceeded with a temperature gradient of 4 °C for 5 min, followed by a linear ramp from 4 to 37 °C at a rate of +0.1 °C every 13 s, then held at 37 °C for 2 h, and finally a linear ramp from 37 to 50 °C at a rate of +0.1 °C every 13 s. The reaction was terminated by heating at 95 °C for 15 min. Deaminase-treated DNA was subjected to PCR amplification. The PCR amplification was conducted in a 50 µL reaction mixture containing 1.25 U of EpiMark Hot Start Taq DNA polymerase (New England Biolabs), 1× reaction buffer, 0.2 mM each of dATP, dGTP, TTP, and dCTP, 0.4 µM forward primer (5'-AGTGATGTTGAGTTTGATGTTGT-3') and 0.4 μM reverse primer (5'-CCAACATTCCACTAACAATTACTCTCT-3'). The PCR program included 95 °C for 5 min, 25 cycles of 95 °C for 30 s, 55 °C for 30 s, 68 °C for 30 s, followed by 68 °C for 5 min. The resulting PCR products were subjected to Sanger sequencing and colony sequencing (TsingKe). Colony sequencing was performed according to previously reported method, 52,53 with detailed procedures available in the ESI.†

Deamination assay for ssDNA with various modifications

The 24-mer cytosine-containing DNA, 24-mer 5mC-containing DNA, and the 24-mer 5hmC-containing DNA were used to evaluate the deaminase activity of orthologous mammalian A3A proteins toward C, 5mC, and 5hmC. The deamination reaction was conducted in a 20 μ L solution containing 10 μ M of the orthologous mammalian A3A protein, 20 mM MES (pH 6.5), 0.1% Triton X-100, and 60 ng of the oligonucleotide at 37 °C for 2 h. The reaction was terminated by heating at 95 °C for 15 min. The deamination rates of the orthologous mammalian A3A proteins were determined using liquid chromatographytandem mass spectrometry (LC-MS/MS) analysis.

Enzymatic digestion of DNA

DNA was enzymatically digested according to the previously described method,⁵⁴ and the detailed procedures are available in the ESI.†

LC-MS/MS analysis

The analysis of nucleosides was conducted according to previous studies. 55-58 The measurements were performed using an LC-MS/MS system, which included a Shimadzu 8045 mass spectrometer (Kyoto, Japan) with a Turbo Ionspray electrospray ionization source, coupled with a Shimadzu LC-30AD UPLC system. The nucleosides were monitored using multiple reaction monitoring (MRM) in positive-ion mode. The optimized MRM parameters are provided in Table S4.† The detailed analytical procedures are available in the ESI.†

Quantitative analysis of 5hmC in specific loci of lung tissue DNA

Genomic DNA from lung cancer tissue and matched adjacent normal tissue were extracted using a tissue DNA kit (Omega Bio-Tek Inc., Norcross, GA, USA). The genomic DNA was fragmented to an average size of 300-500 bp using an Ultrasonic Homogenizer JY92-IIN (Scientz). For OMA-seq, 60 ng of fragmented DNA was denatured by heating and treated with 10 µM of orthologous mammalian A3A protein. For ACE-seq, 1 mg of fragmented DNA was treated with β-GT (New England Biolabs) at 37 °C for 2 h, followed by recovery using KAPA Pure beads (Roche). Subsequently, 60 ng of the treated DNA was denatured by heating and then treated with 10 µM of hA3A protein. The deaminase-treated DNA was amplified using EpiMark Hot Start Taq DNA polymerase (New England Biolabs) with site-specific primers (Table S5†) for the individual test sites. The PCR products were subjected to Sanger sequencing, and the quantification of 5hmC levels was achieved by calculating the peak heights of C and T at the test sites.

Results and discussion

Characteristics of different orthologous mammalian A3A proteins

A3A is a universal deaminase found in most placental mammals.⁴¹ However, only a few studies have reported the deamination activity of orthologous mammalian A3A proteins toward C and 5mC. Except for hA3A, the deamination characteristics of other orthologous mammalian A3A members toward 5mC and 5hmC remains inefficiently explored. In the current study, we expressed six orthologous mammalian A3A proteins *in vitro*, including human A3A (hA3A), cow A3A (cowA3A), panda A3A (pandaA3A), porpoise A3A (porpoiseA3A), green monkey A3A (gmA3A), and dog A3A (dogA3A). We carefully assessed their deamination activity toward C, 5mC, and 5hmC, as well as their substrate preferences. A schematic illustration of these six orthologous mammalian A3A proteins is shown in Fig. 1A. Alignment of these proteins revealed approximately 41% amino acid identity (Fig. S1†).

The deamination activity of hA3A toward cytosine has been reported to be influenced by the adjacent 5' nucleobase. 43,50 We designed three dsDNA substrates (DNA-C, DNA-5mC, and DNA-5hmC) to further evaluate the readouts of C, 5mC, and 5hmC in different sequence contexts (TC, CC, GC, and AC). Without deamination, C, 5mC, and 5hmC base pair with guanine (G) and are read as C in sequencing. However, if C, 5mC, and 5hmC are deaminated, they are converted into uracil (U), thymine (T), and 5-hydroxyuracil (5hmU), respectively, which base pair with adenine (A) and are read as T in sequencing. The dsDNA was denatured to ssDNA and then treated with hA3A. The resulting DNA was amplified by PCR and subjected to Sanger sequencing. The results showed that all cytosines in DNA-C and all 5mC in DNA-5mC were fully deaminated and read as thymine (T) after hA3A treatment (Fig. S2†). In contrast, the 5hmC in DNA-5hmC was completely read as T in YC (Y = C/T) sites but only partially read as T in RC (R = A/G) sites. This indicates that hA3A fully deaminates 5hmC at YC sites while partially deaminating 5hmC at RC sites (Fig. S2†).

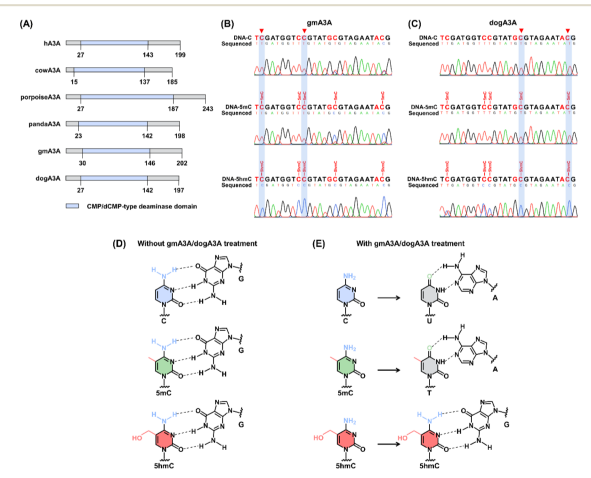


Fig. 1 Assessing the deaminase activity of orthologous mammalian A3A proteins. (A) Schematic representation of the constructs for six mammalian A3A proteins, each containing a CMP/dCMP-type deaminase domain (light blue box). (B) Sequencing results for DNA-C, DNA-5mC, and DNA-5hmC treated with gmA3A. Notably, gmA3A deaminated all C and 5mC at TC and CC sites, resulting in T reads, while 5hmC remained resistant to deamination and was still read as C. (C) Sequencing results for DNA-C, DNA-5mC, and DNA-5hmC treated with dogA3A. dogA3A deaminated all C and 5mC at GC and AC sites, resulting in T reads, while 5hmC remained unchanged and was read as C. (D) Base pairing configurations for C, 5mC, and 5hmC with deamination by gmA3A/dogA3A. (E) Base pairing configurations for C, 5mC, and 5hmC with deamination by gmA3A/dogA3A.

We then evaluated the characteristics of the other five orthologous mammalian A3A proteins. For cowA3A, the results showed that cytosines in DNA-C and 5mC in DNA-5mC were partially deaminated (Fig. S3†), indicating relatively low deamination activity. For porpoiseA3A, cytosines in DNA-C at RC sites and 5mC at CC sites were also partially deaminated (Fig. S4†), suggesting similarly low activity. In the case of pandaA3A, cytosines in DNA-C at RC sites were partially deaminated, while 5mC in DNA-5mC at RC sites was not deaminated, and at CC sites, it was partially deaminated (Fig. S5†). None of these three A3A proteins exhibited deamination activity toward 5hmC. These results suggest that hA3A, cowA3A, porpoiseA3A, and pandaA3A are unsuitable for method development.

For gmA3A, the results showed that all cytosines in DNA-C and all 5mC in DNA-5mC were fully deaminated and read as T, while all 5hmC in DNA-5hmC resisted deamination and was read as C at YC sites (Fig. 1B and S6†). Similarly, after treatment with dogA3A, all cytosines in DNA-C and all 5mC in DNA-5mC were read as T, while all 5hmC in DNA-5hmC was read as C at RC sites (Fig. 1C and S7†). The combined use of gmA3A and dogA3A may enable the distinction of 5hmC from C and 5mC at all sites, which encourages us to employ them in developing a method to detect 5hmC at single-base resolution (Fig. 1D and E).

OMA-seq

Leveraging the complementary deamination properties of gmA3A and dogA3A toward C, 5mC, and 5hmC in different sequence contexts, we developed an orthologous mammalian A3A-mediated sequencing (OMA-seq) method for singlenucleotide resolution and quantitative detection of 5hmC (Fig. 2). At YC (Y = C/T) sites, gmA3A fully deaminates C and 5mC to form U and T, which base pair with A and are read as T, while showing no deamination activity toward 5hmC, which remains paired with G and is read as C in sequencing (Fig. 2A). At RC (R = A/G) sites, dogA3A similarly fully deaminates C and 5mC to U and T, which base pair with A and are read as T, while

also showing no deamination activity toward 5hmC, which remains paired with G and is read as C (Fig. 2A). The detection of 5hmC in various sequence contexts, including TC, CC, GC, and AC sites, is achieved through the combined use of gmA3A and dogA3A. In OMA-seq, gmA3A detects 5hmC at TC and CC sites, while dogA3A detects it at GC and AC sites, with C and 5mC read as T, and 5hmC read as C (Fig. 2B).

To gain further insight into the deamination activity of gmA3A and dogA3A, we conducted LC-MS/MS analysis using various substrates. This analysis evaluated the deamination efficiency of gmA3A and dogA3A toward C, 5mC, and 5hmC. Based on the property of gmA3A for deaminating C, 5mC, and 5hmC, we treated three different DNA mixtures (TC-C and CC-C; TC-5mC and CC-5mC; TC-5hmC and CC-5hmC) with gmA3A, followed by LC-MS/MS analysis. The results showed that the signals for dC and 5mC were undetectable, while the signal intensity of 5hmC remained nearly intact after treatment with gmA3A (Fig. 3A). Similarly, we treated another set of DNA mixtures (GC-C and AC-C; GC-5mC and AC-5mC; GC-5hmC and AC-5hmC) with dogA3A and performed LC-MS/MS analysis. The results indicated that the signals for dC and 5mC were also undetectable, while the signal intensity of 5hmC was nearly unchanged following treatment with dogA3A (Fig. 3B). No significant changes were observed in the other normal nucleosides (dA, dG, and T) after treatment with either gmA3A or dogA3A (Fig. S8†). The LC-MS/MS results were consistent with those obtained from Sanger sequencing. Collectively, gmA3A demonstrated complete deamination of C and 5mC but showed no deamination activity toward 5hmC at TC and CC sites. In contrast, dogA3A also exhibited complete deamination of C and 5mC but did not deaminate 5hmC at GC and AC sites.

Quantitative evaluation of the deamination rates of 5mC and 5hmC in OMA-seq

We next assessed the quantitative capability of OMA-seq. Two types of dsDNA, DNA-5mC and DNA-5hmC, were utilized as

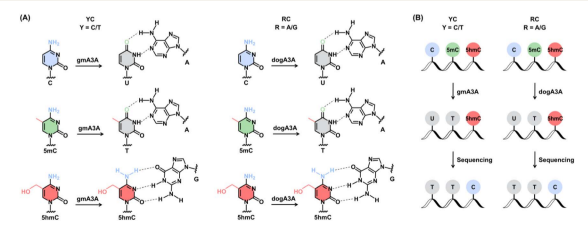


Fig. 2 Schematic illustration of the OMA-seg method. (A) Deamination patterns at YC and RC sites. C and 5mC are converted to U and T, respectively, by gmA3A (for YC sites) and dogA3A (for RC sites), while 5hmC remains resistant to deamination. As a result, U and T base pair with A, whereas 5hmC base pairs with G. (B) Sequencing results after treatment with gmA3A (for YC sites) and dogA3A (for RC sites). C and 5mC are deaminated and read as T, whereas 5hmC is not deaminated and retains its original base pairing. By combining gmA3A and dogA3A, OMA-seq enables site-specific and quantitative detection of 5hmC in diverse sequence contexts.

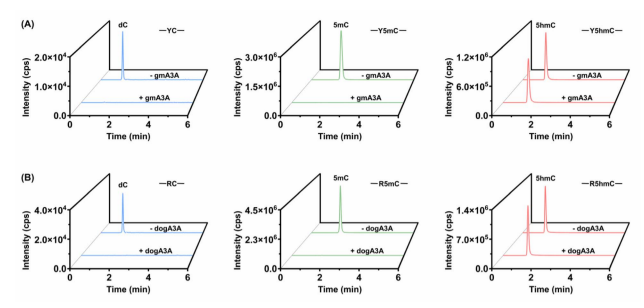


Fig. 3 Assessment of gmA3A and dogA3A deamination activity toward C, 5mC, and 5hmC using LC-MS/MS analysis. We prepared three DNA mixtures, each containing a different combination of nucleotides (TC-C and CC-C; TC-5mC and CC-5mC; TC-5hmC and CC-5hmC), and treated them with gmA3A. Similarly, we prepared another three DNA mixtures (GC-C and AC-C; GC-5mC and AC-5mC; GC-5hmC and AC-5hmC) and treated them with dogA3A. (A) Extracted-ion chromatograms show the deamination of dC, 5mC, and 5hmC in the presence or absence of gmA3A treatment. (B) Extracted-ion chromatograms demonstrate the deamination of dC, 5mC, and 5hmC in the presence or absence of dogA3A treatment.

substrates, and colony sequencing was employed to evaluate the readouts of 5mC and 5hmC in OMA-seq (Fig. 4A). The colony sequencing results showed that all 5mC was fully deaminated and read as T, while 5hmC exhibited significant resistance to deamination, with a zero deamination rate at TC sites and

a slightly 2% deamination rate at CC sites after treatment with gmA3A (Fig. 4B and S9†). Similarly, after treatment with dogA3A, all 5mC was fully deaminated and read as T, while 5hmC remained largely resistant to deamination, with a zero deamination rate at GC sites and a 2% deamination rate at AC

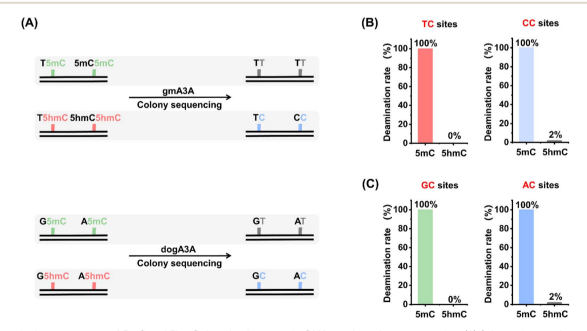


Fig. 4 Quantitative assessment of 5mC and 5hmC deamination rates in OMA-seq by colony sequencing. (A) Schematic overview of the evaluation process, which involves treating dsDNA samples (DNA-5mC and DNA-5hmC) with gmA3A or dogA3A, followed by colony sequencing. Fifty clones from each sample were randomly selected and sequenced. (B) Deamination rates at YC sites after gmA3A treatment. (C) Deamination rates at RC sites after dogA3A treatment.

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sites (Fig. 4C and S10†). Since cytosine is a more suitable substrate than 5mC and both gmA3A and dogA3A can fully deaminate 5mC, cytosine can also be fully deaminated by both enzymes. Taken together, the combined use of gmA3A and dogA3A enables the distinct detection of 5hmC in all sequence contexts.

Quantitative performance of OMA-seq

We quantitatively assessed the levels of 5hmC at specific sites using OMA-seq. Various mixtures of DNA-C and DNA-5hmC were prepared, with the percentage of DNA-5hmC ranging from 0% to 100%. These mixtures were analyzed using OMA-seq followed by Sanger sequencing. The measured ratio of C/(C+T) in Sanger sequencing reflects the 5hmC level in the mixture. The results indicated that the measured 5hmC levels increased proportionally to the theoretical percentages of 5hmC in the mixtures at the specified sites (Fig. 5). In conclusion, these results demonstrate that the OMA-seq method enables site-specific and quantitative detection of 5hmC in DNA.

Quantitative and site-specific detection of 5hmC in genomic DNA

We then applied OMA-seq to detect 5hmC in specific genomic regions. Dysregulation of 5hmC has been observed in various cancers. Previous studies have shown that 5hmC levels are significantly reduced in cancer cell lines and primary tissues. Additionally, 5hmC signature can serve as diagnostic biomarkers for certain human cancers. In this study, we utilized the OMA-seq method for site-specific and quantitative detection of 5hmC in lung cancer tissue and its adjacent normal tissue.

Since the deamination activities of gmA3A and dogA3A in the OMA-seq method can distinguish 5hmC at TC/CC sites and GC/ $^{\prime}$

AC sites, respectively, we selected different adjacent 5′ nucleobases of cytosine (TC, CC, GC, and AC sites) in genomic DNA for evaluation (Table S5†). 5hmC modification was reported to be present at these four sites. 65 Genomic DNA was extracted and fragmented from lung cancer tissue and its adjacent normal tissue. The fragmented DNA underwent deamination treatment with gmA3A (for TC and CC sites) or dogA3A (for GC and AC sites), followed by PCR amplification. Sanger sequencing results indicated that all four sites were partially read as C and partially read as T in normal tissue (Fig. 6 and S12–S15†), suggesting the presence of 5hmC. In contrast, all four sites were completely read as T in cancer tissue (Fig. 6 and S12–S15†), indicating the absence of 5hmC at these sites. These results demonstrate that 5hmC levels are significantly reduced in cancer tissue compared to adjacent normal tissue.

We also employed the previous ACE-seq method for the quantitative analysis of 5hmC at these four sites (Fig. S11†). The quantitative results indicated that, using the OMA-seq method, the 5hmC levels in normal tissues were 15.7% at the TC site, 18.9% at the CC site, 24.0% at the GC site, and 50.8% at the AC site (Fig. 6 and S12-S15†). Similarly, the ACE-seq method detected 5hmC levels of 16.4% at the TC site, 20.4% at the CC site, 22.9% at the GC site, and 48.2% at the AC site in normal tissues. Notably, the levels of 5hmC in lung cancer tissue measured by both methods were close to zero, highlighting a significant reduction in 5hmC in cancerous tissues compared to normal tissues. The consistency of results obtained from both OMA-seg and ACE-seg underscores the reliability of these methods for quantifying 5hmC. These results are particularly important as it validates the use of OMA-seq as a robust tool for site-specific and quantitative detection of 5hmC in genomic DNA.

The introduction of new and natural enzymes for detecting 5hmC within the OMA-seq framework enhances its applicability

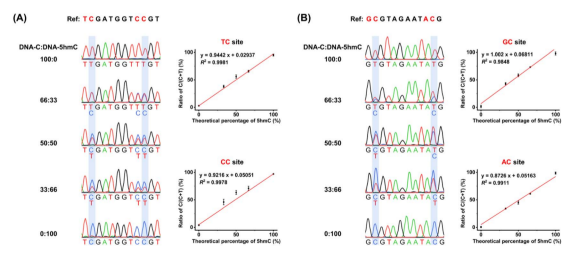


Fig. 5 Assessment of the quantitative capability of OMA-seq. The mixtures of DNA-C and DNA-5hmC with varying proportions of DNA-5hmC (0-100%) were employed. (A) Quantification of 5hmC levels at YC sites using OMA-seq. The mixtures were treated with gmA3A and analyzed by Sanger sequencing (left panel). A linear regression analysis was performed to correlate the measured C/(C + T) ratios at individual sites with the theoretical 5hmC percentages in the DNA-C and DNA-5hmC mixtures (right panel). (B) Quantification of 5hmC levels at RC sites using OMA-seq. The mixtures were treated with dogA3A and analyzed by Sanger sequencing (left panel). A linear regression analysis was performed to correlate the measured C/(C + T) ratios at individual sites with the theoretical 5hmC percentages in the DNA-C and DNA-5hmC mixtures (right panel).

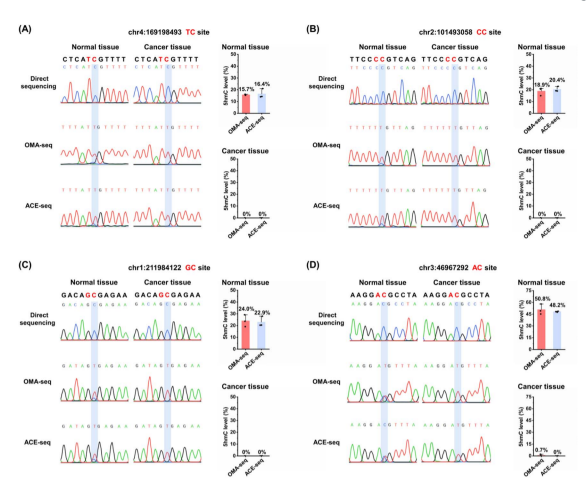


Fig. 6 Site-specific and quantitative analysis of 5hmC in lung cancer tissue and matched adjacent normal tissue using OMA-seq and ACE-seq. (A) 5hmC level at specific genomic loci: chr4:169198493 (TC site). (B) 5hmC level at chr2:101493058 (CC site). (C) 5hmC level at chr3:46967292 (AC site).

and effectiveness. The OMA-seq method is based on a straightforward principle and features a simple analytical procedure. In OMA-seq, the deamination reaction occurs under mild conditions that prevent DNA degradation, allowing for the use of only nanogram quantities of DNA for 5hmC analysis. This technique is particularly useful when there is a limited amount of input DNA, making it ideal for site-specific and quantitative analysis of 5hmC, including applications in single-cell and cell-free DNA studies. This innovative approach, when combined with highthroughput sequencing technologies, holds great promise for enabling the quantification and genome-wide mapping of 5hmC at single-nucleotide resolution in future studies. Such advancements could significantly contribute to our understanding of epigenetic modifications in various biological contexts, including cancer research, and may lead to the identification of novel biomarkers for early diagnosis and targeted therapies.

Conclusions

We conducted a systematic characterization of the deaminase activities of six orthologous mammalian A3A proteins on C, 5mC, and 5hmC in DNA. Notably, gmA3A and dogA3A

exhibited distinct deamination profiles compared to hA3A. Specifically, gmA3A completely deaminated C and 5mC at YC (Y = C/T) sites, but showed no activity towards 5hmC. In contrast, dogA3A effectively deaminated C and 5mC at RC (R = A/G) sites, with no activity on 5hmC. By combining gmA3A and dogA3A, we achieved differential deamination of C, 5mC, and 5hmC at diverse cytosine sites. Following enzyme treatment, C and 5mC were converted to T during sequencing, while 5hmC remained resistant to deamination and was still read as C. Building on these findings, we developed the OMA-seq method for quantitative and site-specific detection of 5hmC in DNA across various sequence contexts. This method enables quantitative analysis of 5hmC at individual sites in genomic DNA from lung cancer tissue and adjacent normal tissue. The OMA-seq approach is straightforward, requiring only a single-step deamination without the need for bisulfite treatment, DNA glycosylation, or chemical oxidation. This provides a valuable tool for the quantitative detection of 5hmC in DNA at single-nucleotide resolution. Furthermore, our strategy of utilizing orthologous enzymes for 5hmC detection expands the range of biological enzymes applicable in the localization analysis of modified nucleosides, offering new opportunities for epigenetic research.

Edge Article

Data availability

The data supporting this article have been included as part of the ESI.†

Author contributions

Bi-Feng Yuan: writing - review & editing, investigation, supervision, conceptualization, funding acquisition. Neng-Bin Xie: writing - review & editing, investigation, supervision, conceptualization. Yu Liu: writing - review & editing, investigation, supervision, conceptualization. Xia Guo: writing - original draft, methodology, formal analysis, data curation, conceptualization. Jianyuan Wu: methodology, formal analysis, data curation, conceptualization. Tong-Tong Ji: formal analysis, data curation, conceptualization. Min Wang: methodology, data curation. Shan Zhang: methodology, data curation. Jun Xiong: data curation. Fang-Yin Gang: data curation. Wei Liu: data curation. Yao-Hua Gu: formal analysis.

Conflicts of interest

The authors declare no competing financial interests.

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