Detection of preQ₀ deazaguanine modifications in bacteriophage CAjan DNA using Nanopore sequencing reveals same hypermodification at two distinct DNA motifs

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Received March 18, 2020; Revised August 19, 2020; Editorial Decision August 20, 2020; Accepted August 26, 2020

ABSTRACT

In the constant evolutionary battle against mobile genetic elements (MGEs), bacteria have developed several defense mechanisms, some of which target the incoming, foreign nucleic acids e.g. restrictionmodification (R-M) or CRISPR-Cas systems. Some of these MGEs, including bacteriophages, have in turn evolved different strategies to evade these hurdles. It was recently shown that the siphophage CAjan and 180 other viruses use 7-deazaguanine modifications in their DNA to evade bacterial R-M systems. Among others, phage CAjan genome contains a gene coding for a DNA-modifying homolog of a tRNAdeazapurine modification enzyme, together with four 7-cyano-7-deazaguanine synthesis genes. Using the CRISPR-Cas9 genome editing tool combined with the Nanopore Sequencing (ONT) we showed that the 7deazaguanine modification in the CAjan genome is dependent on phage-encoded genes. The modification is also site-specific and is found mainly in two separate DNA sequence contexts: GA and GGC. Homology modeling of the modifying enzyme DpdA pro-

vides insight into its probable DNA binding surface and general mode of DNA recognition.

INTRODUCTION

Bacteriophages are the most abundant biological entities in the biosphere (1). Phages have an immense genetic diversity, different lifestyle strategies and have major influence on the composition of the bacterial communities. The constant struggle for persistence between phages and their hosts is often referred to as the oldest 'arms race.' On one side, bacteria have developed strategies to eliminate phages, such as the use of restriction endonucleases (REs) (2), whereas phages evolved counter measures to circumvent the bacterial antiphage defense systems, which include modifying their nucleotides (3). Phage genomes are very diverse, having either single or double stranded RNA or DNA as nucleic material and even some of the viral nucleotides can deviate from the canonical A, T, C and G (4,5).

Phage CAjan belongs to the *Siphoviridae* family (dsDNA genome, long non-contractile tail) and infects the well-characterized strain *Escherichia coli* K-12 MG1655 (6,7). The genome of CAjan is resistant to digestion by several restriction enzymes (REs) (8). REs have been known for decades as a part of the innate bacterial resistance arsenal against invading exogenous DNA, such as phage genomes

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(9). Moreover, it was recently shown that the genome of phage CAjan encodes for several proteins with a high level of similarity to enzymes of the 7-deazaguanine biosynthesis pathway which is reported to modify certain tRNAs as well as DNA (10,11). In fact, we showed that CAjan hypermodifies its genome with a 7-deazaguanine modification, specifically 7-cyano-7-deazaguanine (also known as $preO_0$ (8). Remarkably, the 7-deazaguanine modification gene cluster in phage CAjan consists of twelve genes (Figure 1). Six of them can be assigned a function: four $preQ_0$ synthesis genes (folE, queD, queE, queC) as well as yhhQ gene which codes for a putative preQ₀ transporter and dpdAgene for a paralog of the archaeal guanine-tRNA translgycosylase (arcTGT) enzyme. The latter removes the G base at position 15 in target tRNA and exchanges it with $preQ_0$ (10). The rest of the genes in the 7-deazaguanine modification cluster remain without an assigned function (Figure 1). DNA modifications can be detected in genomes using various methodologies including restriction enzyme analysis, mass spectrometry or bisulphite sequencing (12). Recently, third generation sequencing technologies have been used to detect various modifications, mostly methylations, in DNA (13-15) and even in RNA (16). Nanopore-based sequencing applied by Oxford Nanopore Technologies (ONT) is a very promising method to detect a diverse set of modifications in DNA. Here, we investigate whether $preQ_0$ modifications in DNA can be detected by ONT sequencing and whether these substitutions occur at specific sites within phage CAjan genome. To achieve this, an in vitro synthetic DNA template identical to phage CAjan DNA was generated using the canonical nucleotides in order to provide a modification-free DNA reference for Nanopore sequencing. Moreover, using the CRISPR-Cas9 technology, we generated a series of phage mutants deleted in specific genes of the 7-deazaguanine biosynthesis pathway to examine their involvement and effect on the modification of CAjan's DNA (17). The results provide insights into the $preQ_0$ modifications in phage CAjan genome and offer a roadmap to detect additional 7-deazaguanine modifications in DNA and the sequence motifs in which they lie.

MATERIALS AND METHODS

Phage-host pair

The phage investigated is Enterobacteria phage CAjan (GenBank accession number NC_028776.1) and its host is *E. coli* K-12, MG1655 (U00096.3).

Unmodified whole genome amplification of DNA reference

An unmodified whole genome amplification (WGA) of phage CAjan was prepared using the illustraTM Ready-To-GoTM GenomiPhiTM V3 DNA amplification kit (GE Health-care, Pittsburgh, US), which uses Phi-29 DNA polymerase. This was followed by debranching with S1 nuclease. Briefly, 10 µl of 5× nuclease buffer and 2 µl (200U) of S1 nuclease (ThermoFischer Scientific) were mixed with 38 µl of WGA product and incubated at 25°C for 30 min followed by inactivation at 70°C for 10 min. Product was then purified using the Clean-up AX kit (A&A Biotechnology) according to the manufacturer's instructions.

pL2Cas9 construction

sgRNA targeting the phage genes of interest were cloned into pL2Cas9 (Addgene.org, plasmid #9884) (17). Pairs of designed oligonucleotides (50 μ m) were phosphorylated by mixing 2 μ l of each with 10 μ l of 5× T4 DNA ligase buffer (Thermo Scientific), 1 μ l of T4 kinase (Thermo Scientific) and 32 μ l of ddH₂O followed by incubating at 37°C for 30 min, and then at 65°C for 20 min. The phosphorylated oligos were then ligated into Bsal (NEB) digested pL2Cas9 (molar ratio 5:1) using T4 DNA ligase (Thermo Scientific). The ligation products were then dialyzed on membranes and transformed into the bacterial host by electroporation. Successful transformation was verified by PCR and sequencing. Oligonucleotide pairs for each deletion mutant are listed in Supplementary Table S1.

pNZ123 construction

The repair templates were constructed using Gibson assembly (18). Regions flanking the targeted gene were amplified by PCR with the DreamTaq polymerase (Thermo Scientific) according to manufacturer's protocol. PCR products were cleaned with the Clean & Concentrator-5 PCR cleanup kit (Zymo Research). The primer pairs are listed in Supplementary Table S1. Then, a one-step isothermal assembly with Xbal digested pNZ123 was performed as described elsewhere (18). Again, the resulting product was dialyzed on membranes and transformed into the host by electroporation. PCR-confirmed transformants were purified and the repair template plasmids were isolated from overnight cultures using the Plasmid Mini kit (A&A Biotechnology). The purified plasmids were then transformed into the bacterial recombinant host harboring the modified pL2Cas9, as described previously (19). Bacterial cultures containing both plasmids were used to generate the phage mutants (17).

CRISPR-Cas9 mutagenesis

The phage mutants deleted in *folE*, *queC*, *dpdA* or *yhhQ* were generated using the CRISPR–Cas9 technology as previously described (17). In short, the selected phage genes were targeted by specifically designed sgRNA encoded into pL2Cas9-derivative plasmids. Repair templates of the target genes designed to remove 150-350 bp within each gene, including the protospacers, were supplied on pNZ123-derivative plasmids. Hosts harboring both plasmids were then used for multiple cycles of phage infection, resulting in the generation of deletion mutants. Deletions were verified by PCR and confirmed by whole genome sequencing to also verify the absence of off-target mutations.

Amplification of wild-type and deletion-mutant phages and isolation of viral DNA

Wild-type phage CAjan (CAjan WT) and its phage mutants deleted in either *folE*, *queC*, *dpdA* or *yhhQ* were amplified to high titers by separately growing them in 500 mL of host cultures (w/wo modification plasmids) to an early log-phase and infecting half of the volume (125 ml) with a low multiplicity of infection (\sim 0.01). The culture medium was Brain–Heart Infusion (BHI, Oxoid), and for the deletion mutants,



Figure 1. (A) The 7-deazaguanine modification gene cluster in CAjan bacteriophage (NC_028776.1) located in the region 28 621–34 560 bp. Genes are colored by function: purple are genes involved in preQ₀ synthesis, orange is the gene coding for the preQ₀ transporter, blue is the dpdA gene and in white are genes coding for unknown functions. Numbers correspond to gene product numbers from NC_028776. On the right side, there is a 500 bp grey rectangle for size reference and rest of the Figure is drawn to this scale. (B) Biosynthesis pathway of preQ₀. The biosynthesis starts with GTP which is transformed to preQ₀ by enzymes FoIE, QueD, QueE and QueC. Alternatively, YhhQ can transport preQ₀ from the outside. The last step is DpdA enzyme that exchanges targeted G bases with the preQ₀.

BHI was supplemented with chloramphenicol (20 μ g/ml) and erythromycin (150 μ g/ml) to maintain the two plasmids. Both the infected and non-infected cultures were incubated at 37°C, with agitation (200 rpm). Phage lysates were treated essentially as described for lambda phage by Sambrook and Russell (20). Followed by the phenol–chloroform DNA extraction and ethanol precipitation. The DNA was dissolved in sterile deionized water.

Restriction endonuclease digestions

Restriction digestions were prepared in 10 μ l volume reaction containing ~500 ng of genomic DNA, 1 μ L of restriction enzyme: AanI, BcuI, BstEII, EcoRV, HaeIII or NsiI (ThermoFisher) and 1 μ l of a 10× corresponding reaction buffer. Reactions were performed at 37°C for 1 h. Afterwards, DNA fragments were visualized on a 1% agarose gel using GelRed as the DNA dye in a loading buffer.

3D protein modeling

A homology model of the CAjan dpdA gene product was generated using SWISS-MODEL (21). A template search in SWISS-MODEL returned the crystal structure of *Pyrococcus horikoshii* arcTGT as the top scoring template (PDB IDs 1IQ8 and 1IT8, (22)) with 15% sequence identity and 30% similarity to CAjan DpdA. The model quality is moderate as reflected by a mid-range GMQE (Global Model Quality Estimation) score of 0.43 and a QMEAN Z-score (an estimate of the degree of nativeness of the model) of -3.5 (21). A 10-base-pair dsDNA starting model containing an extrahelical G base was extracted from the crystal structure of the DNA repair enzyme AGT in complex with DNA (PDB ID 1T38, (23)), and its sequence changed to contain A, or GC 3' of the flipped G nucleotide. The resulting model was then docked onto the protein model using the HADDOCK server (version 2.2, (24)). In the docking protocol, active site residues Asp105, Asp63, Asp206, His132 and Phe189, and the flipped G nucleotide of DNA were designated as active residues to apply distance restraints.

Mass spectrometry of DNA

Quantitative analysis of DNA modifications was performed as described previously but with several modifications (8). Purified DNA was hydrolyzed in 10 mM Tris-HCl (pH 7.9) with 1 mM MgCl₂ with Benzonase (20 U), DNase I (4 U), calf intestine phosphatase (17 U) and phosphodiesterase (0.2 U) for 16 h at ambient temperature. Following passage through a 10 kDa filter to remove proteins, the filtrate was lyophilized and resuspended to a final concentration of 0.2 μ g/ μ l (based on initial DNA quantity). Quantification of the modified 2'-deoxynucleosides (dADG, dQ, $dPreQ_0$, $dPreQ_1$ and dG^+) and the four canonical 2'deoxyribonucleosides (dA, dT, dG and dC) was achieved by liquid chromatography-coupled triple quadrupole mass spectrometry (LC-MS/MS) and in-line diode array detector (LC-DAD; for quantifying canonical nucleosides). Aliquots of hydrolyzed DNA were injected onto a Phenomenex Luna Omega Polar C18 column ($2.1 \times 100 \text{ mm}$, 1.6 μ m particle size) equilibrated with 98% solvent A (0.1% v/v formic acid in water) and 2% solvent B (0.1% v/v formic acid in acetonitrile) at a flow rate of 0.25 mL/min and eluted with the following solvent gradient: 2-12% B in 10 min; 12-2% B in 1 min; hold at 2% B for 5 min. The HPLC column was coupled to an Agilent 1290 Infinity DAD and an Agilent 6490 triple quadruple mass spectrometer (Agilent). The column was kept at 40°C and the auto-sampler was cooled at 4°C. The UV wavelength of the DAD was set at 260 nm and the electrosprav ionization of the mass spectrometer was performed in positive ion mode with the following source parameters: drying gas temperature 200°C with a flow of 14 1/min, nebulizer gas pressure 30 psi, sheath gas temperature 400°C with a flow of 11 l/min, capillary voltage 3000 V and nozzle voltage 800 V. Compounds were quantified in multiple reaction monitoring (MRM) mode with the following m/z transitions: $310.1 \rightarrow 194.1, 310.1 \rightarrow 177.1, 310.1 \rightarrow 293.1$ for dADG, $394.1 \rightarrow 163.1, 394.1 \rightarrow 146.1, 394.1 \rightarrow 121.1$ for dQ, 292.1 \rightarrow 176.1, 176.1 \rightarrow 159.1, 176.1 \rightarrow 52.1 for dPreQ₀, 296.1 \rightarrow 163.1, 296.1 \rightarrow 121.1, 296.1 \rightarrow 279.1 for dPreQ₁ and $309.1 \rightarrow 193.1, 309.1 \rightarrow 176.1, 309.1 \rightarrow 159.1$ for dG⁺. External calibration curves were used for the quantification of the modified canonical 2'-deoxynucleosides. The calibration curves were constructed from replicate measurements of eight concentrations of each standard. A linear regression with $r^2 > 0.995$ was obtained in all relevant ranges. The limit of detection (LOD), defined by a signal-to-noise ratio (S/N) > 3, ranged from 0.1 to 1 fmol for the modified 2'-deoxynucleosides. Data acquisition and processing were performed using MassHunter software (Agilent).

Nanopore sequencing

Barcoded Nanopore sequencing libraries were built from phage DNAs and the WGA DNA serving as negative control without modifications. Libraries were built using the Rapid Barcoding Sequencing kit RBK-004 (ONT) and sequenced on the MinION platform using a single R9.4 flowcell. Sequencing was performed using MinKNOW (25). Base calling was performed with Guppy v.2.1.3 (26) using default parameters. Only barcoded reads were used for downstream analysis.

Nanopore data analysis

Nanopore sequencing data were analyzed with the Tombo v.1.5 software (15) for detection of modified bases. Briefly, fast5 files for each barcoded library were compared against the reference CAjan phage genome, using the Tombo 'resquiggle' command. Modified bases were detected in the wild-type and mutants by comparing reads of the respective samples with the phi29-amplified negative control containing no DNA modifications, using the Tombo function 'detect_modifications model_sample_compare'. Results were further parsed within the Tombo software suite and a DNA motif logo of the modified bases were made with the MEME software (27) using 'Zero or One Occurrence Per Sequence (zoops) approach, as recommended in the Tombo manual. From Tombo, the fraction of modified reads (Mod-Frac) were used for WT and all mutants for plotting the CAjan genome with modified bases in Circos (28). ModFrac is defined as a value ranging from 0 to 1 that reflects how big a fraction of individual reads has a potential modification in the specific position of the genome. For example, when the total coverage of a certain nucleotide position is 100 and the signal of 50 of these reads deviates between the two samples then the ModFrac for this position is 0.5. The same data was imported into R for further analyses and plotting using the ggplot2 package. For generating 'corrected ModFrac values' in Table 2, a custom Python script was used. In short, it found Gs with a high ModFrac values (>0.883) and replaced the ModFrac of two upstream nucleotides (NN) with 0, provided they were not GA. In cases where upstream nucleotide was GA only the ModFrac of A was replaced by 0.

RESULTS AND DISCUSSION

Validation of the presence of DNA modifications in CAjan phage

The quantitative landscape of 7-deazaguanine modifications in the CAjan phage genome was defined by LC-MS/MS analysis of the nucleosides dADG, dQ, dPreQ₀, dPreQ₁ and dG⁺ for WT and $\Delta queC$ phage mutant. dPreQ₀ was the only 7-deazaguanine detected and was present in both WT and $\Delta queC$ phage CAjan DNA, though $\Delta queC$ DNA contained 12-fold lower levels than WT (3.5 per 10^3 nt versus 44 per 10^3 nt, respectively). This was expected as, although the phage QueC does not contribute to the production of dPreQ₀ in the $\Delta queC$ mutant, the phage DpdA is still present and can insert a small amount of PreQ₀ synthetized by the host into the phage. Interestingly, N^6 methyl-dA was also detected in both $\Delta queC$ and WT, but with 10-fold higher levels in $\Delta queC$ (83 per 10⁶ nt versus 865 per 10^6 nt, respectively). This is similar to what was observed for the phage T4, where mutants unable to produce glucosyl 5-hydroxymethylcytosine (glc-HMC) modification had a higher amount of methylation present in the 5'-GATC-3' sequence (29) and suggests that the activity of adenine methylase is partially inhibited by glc-HMC or $dPreQ_0$.

Nanopore sequencing can detect preQ₀ modifications

Nanopore sequencing of the 59 670-bp genome (44.7 GC%) of the WT phage CAjan and its the deletion mutants, and the WGA DNA yielded more than adequate coverage for Tombo analyses (Table 1). In previous analysis, methylation sites were detected with a relatively low average coverage of $\sim 15 \times$ (U-test applied if at least five reads were present in both samples) (15). Here, we obtained much higher coverage (between $1207 \times$ and $42970 \times$), which significantly increases the statistical power, thus the relevance of this study (15). The control containing only canonical bases had the lowest average coverage of $1207 \times$, likely due to the DNA pooling. On the other hand, the presence of $preQ_0$ seems to partly impair the Nanopore sequencing process as manifested by a lower mean of Phred quality scores for samples with preQ₀ modification (WT and $\Delta yhhQ$) compared to the other samples (Table 1).

Comparison of the unmodified reads derived from the WGA genome to those of the wild-type CAjan phage and its mutants resulted in assigning the fraction of modified reads value (ModFrac) to each nucleotide position. The Tombo package does not suggest a fixed cut-off for ModFrac values to assess whether a base is modified or not. To find this cutoff, the density of ModFrac values for each of the four nucleotides were plotted for both strands (Figure 2).

In the WT phage CAjan genome, the density distribution clearly shows two groups of peaks corresponding to unmod-

Table 1.	Basic	characteris	stics of	genomic	DNA	preparations	used in	this study
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DNA	Genome size	Coverage	Genomic position of deletion	Sensitivity to RE digestion	Number of reads	Total bases	Mean Phred score
WT	59.670	31.007	None		149 692	1 3 Gbp	63
WGA	59 670	1207	None	n.a.	77 522	70.6 Mbp	8.0
$\Delta folE$	59 448	5 845	30 221	+	19 491	344.6 Mbp	7.5
$\Delta queC$	59 380	1 953	32 786	+	12 816	113.9 Mbp	7.5
$\Delta dp dA$	59 353	42 970	29 212	+	140 692	1.2 Gbp	7.6
$\Delta yhhQ$	59 120	29 334	31 850	_	122 460	1.0 Gbp	6.3



Figure 2. Density plot of the modified fraction for each nucleotide in the CAjan WT dataset. Continuous line shows plus strand, while dashed line shows minus strand. The vertical line indicates the ModFrac cutoff calculated as the mean of the rightmost peak for G (plus strand) subtracted by one standard deviation.

ified and modified bases. As previously shown, phage CAjan modifications occur only on G nucleotides (8). Accordingly, the peak with high ModFrac (mean for G = 0.93) values was extracted for G nucleotides (ModFrac > 0.75) and a cut-off was defined as the mean minus one standard deviation (0.883). On the reverse strand these numbers were similar and were calculated to be 0.921 and 0.871, respectively, suggesting an asymmetric motif. While the G nucleotide had the most prominent peak in the high Mod-Frac range, the A, C and T bases also showed smaller peaks in the same range. As discussed below, this is likely due to a neighboring effect of modified bases (15). Nanopore sequencing of the DNA genome of phage mutant $\Delta yhhQ$ led to density plots comparable to the WT. Conversely, density plots of ModFrac values were similar for both DNA strands of phage mutants $\Delta folE$, $\Delta queC$ and $\Delta dpdA$ (Figure S1), indicating that their deletions eliminated the modified peaks. Although $\Delta dpdA$ is completely lacking the modifications while $\Delta folE$ and $\Delta queC$ mutants have a secondary modification (ModFrac ~ 0.3) peak which is likely caused by the 12-fold lower incorporation of preQ₀ derived from the host metabolism. Because the ModFrac values for each base from the forward and reverse strands, are very similar (Figure 2), only the forward strand was used for further analysis. Plotting the fraction of modified read values (cut-off 0.883) of the phage variants against the CA-jan WT genome confirmed that the WT and $\Delta yhhQ$ mutant have modified positions throughout the genome (Figure 3). This was an expected outcome because the products of *folE*, *queC* and *dpdA* are necessary for the synthesis of



Figure 3. Circular representation of phage CAjan and the fraction of modified reads per nucleotide positions for WT and phage mutants. Starting from the inside, rings display, functional genetic modules (various colors), open reading frames (black: plus strand, grey: minus strand), genome track (yellow), GC-content. The five outermost rings show ModFrac in order from inside: mutants Δ yhhQ, Δ dpdA, Δ queC, Δ folE, and WT ModFrac. Each of the deletion mutants lacks a part of the WT genome and the comparison artificially shows high ModFrac values at the deletion sites.

preQ₀, whereas YhhQ is a preQ₀ transporter dispensable for *de novo* formation of preQ₀ in the host (30). Additionally, all phage mutants showed high ModFrac values in the genomic region corresponding to their respective deletion. This is a consequence of comparing the mutant reads to the WT genome, which is required for comparative analyses in Tombo (Figure 3). Because each deletion mutants lacks a part of its genome, the comparison artificially shows high ModFrac values at the deletion sites. The respective deleted regions for each mutant were removed from the datasets for subsequent analyses. According to LC–MS/MS data a small amount of dPreQ₀ was incorporated to the $\Delta queC$ phage mutant genome, however the signal derived from 12-fold lower level of dPreQ₀ was below the set ModFrac cut-off (0.883).

Nanopore modification calling of $preQ_0$ is convoluted by the neighboring effect

The identified cut-off (0.883, Figure 2) was used to calculate the percentage of modified positions for each of the four nucleotides on the positive strand (Table 2, top). The

 Table 2. Percentage of modified bases using the identified ModFrac cutoff of 0.883

Modified sites bas	ed on ModFrac			
Strain	А	С	G	Т
WT	9.2	9.4	34.9	6.7
$\Delta folE$	0.0	0.0	0.0	0.0
$\Delta queC$	0.0	0.0	0.0	0.0
$\Delta dp dA$	0.0	0.0	0.0	0.0
$\Delta y hhQ$	9.9	10.0	36.0	7.8
Corrected ModFr	ac values			
WT	1.91	0.1	23.11	0.20

WT strain and $\Delta yhhQ$ mutant have similar percentages of modified positions for each of the four nucleotides, while the $\Delta folE$, $\Delta queC$ and $\Delta dpdA$ mutants are below the set ModFrac cutoff (0.883).

While all other evidences indicated that only G's are modified in phage CAjan genome, Table 2 suggests that other nucleotides are also modified in WT and $\Delta yhhQ$. It is documented that nucleotides neighboring a modified base can also appear modified due to the nature of Nanopore sensing (15,31). This is because the nanopore signal comes from multiply contiguous bases occupying the pore (5-6 bases) with 3 bases mostly influencing the signal in newer pores (R9, R9.5) (32). In our dataset, the two bases immediately prior to (5') and to a much lesser extent one subsequent base (3') of the modified base had increased ModFrac values. This pattern is not fully consistent and is muddled if several modified bases are found in close proximity (Figure 4). This neighboring effect is responsible for false-positive results of bases other than G (Table 2) in the genomes of WT CAjan and its mutant $\Delta yhhQ$. Using a custom Python script (see Materials and Methods), we corrected for this neighboring effect in the WT genome (Table 2, bottom part). After the correction, the percentage of modified Gs dropped to 23.11% (52 per 10^3 nt), which is lower than what we reported previously (8) but slightly higher compared to LC-MS/MS results obtained in this study (44 per 10^3 nt). Other factors could affect the percentage, as discussed below, secondary motifs (GGT, GGTT) may have somewhat elevated ModFrac values and could contribute to a higher fraction of modified Gs in the mass spectrometry data. On the other hand, LC-MS/MS data could be affected by residual, unmodified host DNA, while nanopore results are specific to the modified phage DNA.

DNA is modified at two separate sites with different lengths

The cut-off defined above was used to extract significantly modified regions of 15 bases centered on the base with highest ModFrac value, and then motifs of the aligned regions were generated as described in Materials and Methods. A single significant motif (*E*-value 10E–1220) was produced for both strands that are exactly complementary to each other (Figure 5). The motif 5'-GRH-3' was identified, in which the initial G is always appearing as the most conserved base in modified regions, followed by an A or G (R) and then a less conserved A, C or T (H). To further investigate the neighboring effect, all 16 possible combinations of dinucleotides were extracted from the dataset together with their corresponding ModFrac values, and the average Mod-

Frac value for the first of the two nucleotides was plotted (Figure 6A). In support of the 5'-GRH-3' motif identified using MEME (Figure 5), the dinucleotides GA followed by GG showed the highest average ModFrac values of all the dinucleotide combinations (Figure 6A). The dinucleotides ending with a guanine (AG, TG and CG) had average Mod-Frac values elevated above the background level, which is likely due to the neighboring effect discussed above. The density plots of the ModFrac values of all dinucleotide combinations (Figure 7) clearly displayed two peaks for the NG motifs, corresponding to the unmodified (ModFrac ~ 0.15) and modified (ModFrac ~ 0.9) bases, whereas only one peak at ~ 0.9 was present for the GA motif (Figure 7C). This is the expected outcome if the increased average ModFrac values are a result of the neighboring effect, as sequences ending with a G could be modified. The GG motif shows a considerably higher average ModFrac value and larger peak around the ModFrac value 0.9 than the other dinucleotides ending with a guanine (AG, TG and CG). To investigate this pattern, we looked at the average ModFrac value of all trinucleotides (Supplementary Table S2 and Figure 6B). All trinucleotide motifs containing GA (NGA and GAN) had high ModFrac averages (>0.8). Interestingly, the GGC motif also displayed a high ModFrac value (~ 0.9) on par with the GA motifs, indicating that nearly all GGC motifs are modified. This is corroborated by the density plot of the GGC motif (Figure 8A). The ModFrac values of GG dinucleotides (unlike AG, TG and CG) are affected not only by the neighboring effect from modified GA motif but also secondary GGC motif.

In the average ModFrac values of all possible 256 tetranucleotide combinations (Supplementary Table S2), all four base combinations where the first nucleotide has a Mod-Frac average above 0.45 can be explained by the presence of either a GGC or a GA motif.

Taken together, our data strongly indicate that G bases modified with $preQ_0$ in the CAjan phage genome are located within GA and GGC motifs. The vast majority of these motifs were modified in the genome, however some of the sites do not cross the ModFrac threshold (Figure 4). The erroneous estimation of the motif GRH by the MEME software was caused by the fact that the modifications takes place at sites with two different motifs of different lengths. Identification of the correct motifs were further complicated by the large amounts of false positives resulting from the neighboring effect.

Restriction digestion confirms the prediction

We previously reported that $preQ_0$ modification of phage DNA prevents some restriction enzymes from cutting viral DNA (8). In order to biologically confirm the double motif that was predicted, we selected several restriction enzymes and digested both the wild-type and the $\Delta queC$ phage mutant genomic DNAs (Figure 9). Lower levels (12-fold lower) of dPreQ₀ are insufficient to protect the majority of DNA from restriction digestion. As demonstrated in Figure 9, it is clear that an endonuclease (AanI), which does not have a G in its recognition site, is able to cut both DNA samples (Figure 9). When the recognition site contains a GA or GGC, the restriction enzymes (EcoRV and HaeIII) were



Figure 4. Neighbouring effect. In the top part random fragment of plus strain of CAjan phage genome is represented alongside ModFrac values of Δ queC mutant (Δ queC ModFrac +) and WT phage (WT ModFrac +). The bottom part shows the same region for minus strand (WT and Δ queC ModFrac -, respectively). The modified Gs are highlighted in red. High ModFrac values of two preceding bases (the neighbouring effect) can be seen in orange. Gs that are not modified are marked in green. In the dashed square a position where G in GA motif is below the set threshold of modification.



Figure 5. Motif predicted by MEME from significantly modified regions of the WT genome (ModFrac > 0.883). Five positions on either side of the most modified base per region were included for motif discovery.

able to digest the DNA from the $\Delta queC$ phage mutant but not from the wild-type phage (Figure 9). The GC, GT and GGT containing-sites in both phage genomes were cut by restriction enzymes BcuI, NsiI and BstEII. The recognition site of BstEII contains a ubiquitous nucleotide (N) followed by an A, which could result in a GA motif in particular positions in the genome, thereby preventing the restriction enzyme activity, however confirmation of this requires further investigation.

Minor deviations from the discovered motifs

The TGA motif displayed both a slightly lower average ModFrac value than AGA, CGA and GGA (Supplementary Table S2) as well as a slightly distorted peak in the density plot (Figure 8B) compared to the other trinucleotide combinations containing the GA motif. This could indicate that this motif is slightly less likely to be modified than the other GA motifs, even though the difference is quite small. Another possibility is that not all bases are affected equally by the neighboring effect of a $preQ_0$ modification and that the lower ModFrac values of TGA sites compared to AGA, GGA or CGA is a result of lower impacts of the neighboring effects on the thymine at the TGA sites.

The GGT motif has an average ModFrac value of 0.28 (Supplementary Table S2) however, if we exclude sites that are elevated as a result of the neighboring effects, the GGT motif showed a higher average ModFrac value than the background (Supplementary Table S2). To investigate whether the slightly higher ModFrac average was a result of a few GGT sites being completely modified or was due to a fraction of the reads for each site being modified, we inspected the density plot of the GGT motif (Figure 8A). The GGT motif showed no modified peak around the Mod-Frac value 0.9. Instead the GGT motif has the background peak around 0.15 shifted to the right, indicating that only a fraction of the reads for each GGT position was modified. However, when considering the 256 tetranucleotide combinations (Supplementary Table S2), not all possible GGT sites were equally modified. The GGTT site is responsible for a disproportionate amount of the increased ModFrac average of the GGT sites, particularly when the neighboring effect is considered. However, even GGTT sites did not appear to be completely modified, as none of the GGTT sites showing a ModFrac value >0.7 (Supplementary Table S2).

Structural insights into DNA recognition by CAjan DpdA

DpdA was initially identified as a paralog of the archaeal tRNA-guanine translgycosylase (arcTGT) enzyme, which removes the G base at position 15 in the D-loop of archaeal tRNA and exchanges it with preQ₀ (11). ArcTGT utilizes a $(\beta/\alpha)_8$ TIM barrel catalytic domain to bind the D-loop of tRNA and places the G15 substrate in a deeply buried active site at the center of the barrel. The base ex-



Figure 6. Mean ModFrac values of all combinations of dinucleotides (A) and selected trinucleotides (B) with standard errors. Only the ModFrac values of the first nucleotides are plotted.

change reaction starts with nucleophilic attack by a conserved active site Asp side chain on the C1' of the substrate nucleotide to detach the guanine base, forming a covalent intermediate with tRNA. The incoming $preQ_0$ replaces the detached guanine in the same pocket where its N9 atom, following deprotonation by another Asp residue, performs a second nucleophilic attack on C1' to form the product (22,33).

Homology modeling of CAjan phage DpdA in SWISS-MODEL more closely reveals its structural similarity to the catalytic (N-terminal) domain of *Pyrococcus horikoshii* arcTGT (residues 1–350, PDB IDs 1IQ8 and 1IT8 (22), 1.14 Å over 276 C_{α} atoms, Figure 10), despites sequence identity and similarity of only 15% and 30%, respectively. 3D superposition and resulting structure-based sequence alignment show that the preQ₀/guanine nucleotide binding pocket in arcTGT as well as the two catalytic aspartate residues are present in CAjan phage DpdA at the same positions (Figures 10A, B), suggesting a similar catalytic mechanism, even though the two enzymes act on different nucleic acid substrates. Based on this alignment, residues of the CAjan phage DpdA preQ₀/guanine binding pocket would be Ser64, Phe67, Asp105, Gly153, Gly154, His132 and Phe189, the catalytic nucleophile for the first step of the transglycosylation reaction would be Asp206, and the second aspartate for deprotonation of incoming $preQ_0$ is Asp63.

Further, the model suggests that the active site is at the center of a positively charged groove that could accommodate double-stranded DNA (Figure 10C). Like the TGTs and DNA glycosylases involved in DNA repair (23,34), DpdA would be expected to access the substrate G by flipping the base out of the helix into the active site. However, docking of dsDNA harboring a flipped G onto that surface failed to place the extrahelical G deep enough into its binding pocket without significant clashes with the flanking nucleotides, suggesting that a conformational change in the protein and/or partial melting of the DNA duplex is needed to maximize surface complementarity with the enzyme. It is also possible that recognition of the nucleotides 3' of the substrate G may occur via direct readout of flipped out bases as seen in DNA repair endonucleases (35). Overall, a detailed understanding of how DpdA enzymes recognize their target DNA to catalyze sequence-context-specific preQ₀ insertion awaits further biochemical and structural characterization.



Figure 7. Density plots of fraction of modified dinucleotides for the WT phage CAjan dataset staring with A (panel A), C (panel B), G (panel C) and T (panel D).

CONCLUSION

We have demonstrated that Nanopore sequencing can be used for detection of a novel hypermodifications ($preQ_0$) in phage genomic DNA. Phages exhibit a large reservoir of DNA modifications and the possibility of detecting them can lead to the identification of novel phage defense and counter-defense systems. The identification of new enzymes involved in these pathways may also lead to new biotechnology applications. The workflow described in this study can also be applied to detection in other organisms, e.g. bacteria. Additionally, the Nanopore data provide information on the genomic localization of the nucleotide modification (motif), which is not the case when using only a mass spectrometry approach. Yet, our dataset also clearly demonstrated that extra care must be taken when using alignment-based software (e.g. MEME) for motif discovery. This is dictated by two main factors: (a) Nanopore sequencing is affected by the neighboring effect and (b) motifs can have different lengths, thereby complicating the analysis. Therefore, we recommend supplementing the detection process using alignment-based software with



Figure 8. Density plots showing fraction of modified first bases for selected trinucleotide combinations. (A) All trinucleotide combinations starting in GG (GGN). (B) All trinucleotide combinations containing GA (NGA and GAN) with CTA included as a negative control, showing no modification nor neighbouring effect. (C) All trinucleotide combinations ending in guanine (NNG).



Figure 9. Restriction patterns for genomic DNA of WT CAjan and Δ queC mutant. Six different restriction enzymes were used. Enzymes that recognize G in modified motif (GA and GGC) are indicated with red. Enzymes that recognize G in a motif not predicted to be modified are indicated with green (GT, GC and GGT). AanI enzyme does not have G in its recognition site. On each side of the gel is the 2-log ladder for size reference. On the right side there is a graphic representation of the expected restriction pattern. The number of predicted restriction sites is presented on the bottom.



Figure 10. Structural insights into DNA recognition by CAjan phage DpdA. (A) Homology model of CAjan phage DpdA (cyan) superposed on the catalytic domain of *P. horikoshii* arcTGT (brown, PDB ID 11T8) viewed down the TIM barrel. (B) Close up of the superposed active sites showing $preQ_0$ as seen bound to arcTGT. DpdA residues are labeled. (C) Docking model of dsDNA harboring a flipped G onto CAjan phage DpdA showing the putative DNA binding mode to the positively charged groove on the protein surface. The protein is shown in electrostatic surface potential representation. The active site in (A) and (C) is indicated with an asterisk.

biological cross-verification, detection with an additional method (e.g. mass spectrometry), manual verification of the obtained results and in-depth evaluation of phage gene content.

DATA AVAILABILITY

Data were deposited in SRA under project PRJNA633876.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

ACKNOWLEDGEMENTS

We thank Marie-Laurence Lemay for her help with the genome editing of phage CAjan.

FUNDING

Villum Foundation [17595 to W.K.]; Human Frontier Science Program [RGP0024 to L.H., V.dC-L., S.M.]; National Research Foundation of Singapore through the Singapore-MIT Alliance for Research and Technology Antimicrobial Resistance IRG (to C.L. and P.C.D.); National Institute of General Medical Sciences [GM110588 to M.A.S.]; California Metabolic Research Foundation; S.M. holds the Tier 1 Canada Research Chair in Bacteriophages. Funding for open access charge: Villum Foundation [17595]. *Conflict of interest statement*. None declared.

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