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ABSTRACT

Antibody-mediated humoral immunity is thought to play a central role in mediating the immunopathogenesis of acute DENV infection, but limited data are available on the diversity, specificity, and functionality of the antibody response at the molecular level elicited by primary or secondary DENV infection. In order to close this functional gap in our understanding of DENV-specific humoral immunity, we utilized high-throughput single cell RNA sequencing to investigate B cells circulating in both primary and secondary natural DENV infections. We captured full-length paired immunoglobulin receptor sequence data from 9,027 B cells from a total of 6 subjects, including 2,717 plasmablasts. In addition to IgG and IgM class-switched cells, we unexpectedly found a high proportion of the DENV-elicited plasmablasts expressing IgA, principally in individuals with primary DENV infections. These IgA class-switched cells were extensively hypermutated even in individuals with a serologically confirmed primary DENV infection. Utilizing a combination of conventional biochemical assays and high-throughput shotgun mutagenesis, we determined that DENV-reactive IgA class-switched antibodies represent a significant fraction of DENV-reactive IgGs generated in response to DENV infection, and that they exhibit a comparable epitope specificity to DENV-reactive IgG antibodies. These results provide insight into the molecular-level diversity of DENV-elicited humoral immunity and identify a heretofore unappreciated IgA plasmablast response to DENV infection.

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1. Introduction

DENV is a positive-sense single-stranded RNA virus of the family Flaviviridae, genus Flavivirus, that is maintained in a primarily anthropo-ponic cycle between the Aedes aegypti mosquito and humans [1]. Consisting of four genetically and immunologically distinct serotypes (DENV-1, –2, –3, and –4), DENV is thought to infect between 280 and 550 million people worldwide every year, with as many as 100 million infections resulting in some degree of clinical presentation [2,3]. While DENV infection is subclinical in the majority of cases, in susceptible individuals it can cause a debilitating flu-like illness known as dengue fever. The majority of individuals suffering from dengue fever recover without the need for extensive medical intervention, but approxi-mately 500,000 individuals per year develop severe dengue, dengue hemorrhagic fever/dengue shock syndrome (DHF/DSS), which has a mortality rate of up to 20% [4–7].

A unique epidemiological feature of DENV infection is that severe immunopathological symptoms are more likely to occur in individuals previously infected with a heterologous viral serotype compared to individuals without any preexisting DENV immunity [8]. While the
Research in context

Evidence before this study

Antibody-mediated humoral immunity is thought to play a central role in mediating the immunopathogenesis of acute DENV infection. However, limited data are available on the molecular-level diversity, specificity, and functionality of the antibody response elicited by primary or secondary DENV infection. At the onset of this study in 2018 we queried PubMed for articles either directly comparing/contrasting the plasmablast immunoglobulin repertoire elicited by either primary or secondary DENV infection, or for articles that assessed the DENV-elicited plasmablast repertoire in an unbiased fashion without upfront immunoglobulin isotype selection. No previously published report satisfied either of these criteria.

Added value of this study

Our study is the first to examine the molecular level diversity and antigen specificity of the plasmablast antibody repertoire elicited by both primary and secondary dengue in an unbiased and high-throughput fashion. This approach revealed a previously unappreciated role for DENV-reactive IgA in both primary and secondary dengue, with IgA the dominant plasmablast-restricted isotype observed following primary DENV infection.

Implications of all the available evidence

These results identify a heretofore unappreciated role for DENV-reactive IgA in the humoral response to DENV infection, especially in the setting of primary DENV infection.

demonstrated for plasmablasts and plasma cells in animal studies, and the frequency of plasmablast-phenotype B cells circulating 5–10 days after vaccination has been shown to positively correlate with serum antibody titers achieved several weeks after DENV antigen exposure [30–34].

The functional relationship between DENV-elicited plasmablasts and the durable humoral immune profile present after the resolution of infection is further reinforced by work assessing the clonal diversity and antigen specificity of the plasmablast population generated in response to DENV infection. Previous work has demonstrated that the plasmablasts elicited by a secondary DENV infection primarily express broadly cross-reactive, moderately neutralizing, and extensively hypermutated immunoglobulins [32,35–37]. This antigen specificity is reminiscent of the humoral immune profile observed in most individuals following resolution of a secondary DENV infection [33]. In all of these studies, 70%–90% of the immunoglobulins expressed by circulating plasmablasts were observed to bind DENV, and the overwhelming majority of the plasmablasts appear to be derived from memory B cells, as indicated by both the relatively restricted clonal diversity of the population, as well as the extensive pre-existing SHM burden of the antibodies [32,35–37].

In contrast to the abundance of data available on the plasmablast phenotype associated with secondary DENV infection, relatively little data are available on the immunoglobulin receptor specificity and functionality of plasmablasts circulating in response to a primary DENV infection. The most well characterized dataset addressing this question is derived from a human primary dengue serotype 2 infection model [38]. As in secondary DENV infection, abundant plasmablast expansion was observed following infection. However, in contrast to the specificity and functional profile of immunoglobulins generated in response to secondary DENV infection, only ~40% of plasmablast-derived antibodies reacted with DENV, with the majority of antibodies exhibiting serotype-specific reactivity and generally poor neutralization potential. Again, this profile closely mirrors the functional profile of serum following primary DENV infection [39,40].

Although these previous studies provide valuable insight into the molecular complexity of the humoral immune response elicited by DENV infection, there are several significant limitations inherent within the current publication record. Firstly, no data are available on the plasmablast-derived immunoglobulin diversity and functionality following natural primary DENV infection. Secondly, the aforementioned studies restricted their immunoglobulin receptor analysis to cells that expressed IgG or IgM, either by up-front selection during the cell isolation procedure or by virtue of the primers used to amplify the immunoglobulin receptor sequences [32,35–37]. This lack of an unbiased approach to the analysis of plasmablast and immunoglobulin receptor sequence diversity leaves open the possibility that significant features of the humoral immune profile associated with either primary or secondary infection may have been overlooked. Filling this potential knowledge gap may generate new tools for the rapid clinical assessment of previous DENV exposure, and may also provide some insight into the divergent clinical outcomes observed following either primary or secondary DENV infection.

To fill this knowledge gap, we utilized single cell RNA sequencing technology to assess the clonal and transcriptional diversity of plasmablasts elicited in response to primary and secondary DENV infections in an unbiased and high-throughput fashion. Using this approach, we were able to capture full-length paired immunoglobulin receptor sequence data from 9,027 B cells circulating in response to both primary and secondary DENV infection, including 2,717 DENV-elicited plasmablasts. We found an unexpectedly high proportion of plasmablasts expressing IgA in PBMC from patients with primary infection, in addition to the expected IgG and IgM class-switched plasmablasts. These IgA class-switched cells were extensively hypermutated, even in individuals with serologically confirmed primary DENV infection. IgA class-switched IgG
xponents of the plasmablast immunoglobulin repertoire elicited by both primary and secondary dengue in an unbiased and high-throughput fashion. This approach revealed a previously unappreciated role for DENV-reactive IgA in both primary and secondary dengue, with IgA the dominant plasmablast-restricted isotype observed following primary DENV infection.

The primary source of circulating DENV-reactive antibodies persisting after the resolution of infection are non-dividing, terminally differentiated, bone-marrow resident, plasma cells [23–25]. However, while plasma cell-derived antibodies take several weeks to peak and stabilize after initial antigen exposure, B cell plasmablasts can be found in circulation just days after an initial pathogen exposure [23,24,26]. The majority of plasmablasts generated in response to infection rapidly undergo apoptosis following the resolution of inflammation, but a fraction of these cells terminally differentiate into long-lived plasma cells and take-up residency in the bone marrow [25,27–29]. A direct precursor/progeny relationship has been...
secondary DENV infections neutralized DENV and reacted with canonical epitopes on the DENV E protein. Our data demonstrate a previously unappreciated role for IgA class-switched B cells during acute DENV infection - especially during a primary DENV infection - and highlight the utility of unbiased scRNAseq analysis in identifying novel immunological consequences of viral infection.

2. Materials and methods

2.1. Sample collection

Peripheral blood mononuclear cells (PBMC) and plasma were isolated from whole blood specimens obtained from children enrolled in a hospital-based acute illness study in Bangkok, Thailand, the design of which has been previously described [41,42]. Blood samples were obtained serially during acute infection and at early and late convalescent time points; the term ‘fever day’ is used to report acute illness time points relative to Day 0, defined as the day of defervescence. The infecting virus type (DENV-1-4) was determined by RT-PCR and/or virus isolation as previously described [43], and serology (EIA and HAI assays) was used to distinguish primary and secondary DENV infections [44]. Written informed consent was obtained from each subject and/or his/her parent or guardian. The study protocol was approved by the Institutional Review Boards of the Thai Ministry of Public Health, the Office of the U.S. Army Surgeon General, the University of Massachusetts Medical School, and the University of Rhode Island. PBMC and plasma samples were cryopreserved for later analysis.

2.2. Flow cytometry and cell sorting

Cryopreserved PBMC were thawed and placed in RPMI 1640 medium supplemented with 10% heat-inactivated normal human serum (100–318, Gemini Bio-Products), L-glutamine, penicillin, and streptomycin prior to analysis. Cell viability was assessed using CTL-LDC Dye (Cellular Technology Limited [CTL], Shaker Heights, OH) and a CTL-ImmunoSpot S6 Ultimate-V Analyzer (CTL). Surface staining for flow cytometry analysis and cell sorting was performed in PBS supplemented with 2% FBS at room temperature. Aqua Live/Dead (ThermoFisher, L34957) was used to exclude dead cells in all experiments. Antibodies and dilutions used for flow cytometry analysis are listed in Supplemental Table 7. Cell sorting was performed on a BD FACSAria Fusion instrument, and data analyzed using FlowJo v10.2 software (Treestar).

2.3. Single-cell RNA sequencing library generation

Flow-sorted viable B and T cell suspensions were prepared for single-cell RNA sequencing using the Chromium Single-Cell 5’ Reagent version 2 kit and Chromium Single-Cell Controller (10x Genomics, CA) [45]. Approximately 2000–8000 cells per reaction suspended at a density of 50–500 cells/μL in PBS plus 0.5% FBS were loaded for gel bead-in-emulsion (GEM) generation and barcoding. Reverse transcription, RT-cleanup, and cDNA amplification were performed to isolate and amplify cDNA for downstream 5’ gene or enriched V(D)J library construction according to the manufacturer’s protocol. Libraries were constructed using the Chromium Single-Cell 5’ reagent kit, V(D)J Human B Cell Enrichment Kit, 3’/5’ Library Construction Kit, and 17 Multiplex Kit (10x Genomics, CA) according to the manufacturer’s protocol.

2.4. Sequencing

scRNAseq 5’ gene expression libraries and BCR V(D)J enriched libraries were sequenced on an Illumina NovaSeq 6000 instrument using the S1, S2, or S4 reagent kits (300 cycles). Libraries were balanced to allow for ~150,000 reads/cell for 5’ gene expression libraries, and ~20,000 reads/cell for BCR V(D)J enriched libraries. Sequencing parameters were set for 150 cycles for Read1, 8 cycles for Index1, and 150 cycles for Read2. Prior to sequencing, library quality and concentration were assessed using an Agilent 4200 TapeStation with High Sensitivity D5000 ScreenTape Assay and Qubit Fluorometer (Thermo Fisher Scientific) with dsDNA BR assay kit according to the manufacturer’s recommendations.

2.5. 5’ gene expression analysis/visualization

The 5’ gene expression alignment from sorted PBMC was performed using the 10x Genomics Cell Ranger pipeline [45]. Sample demultiplexing, alignment, barcode/UMI filtering, and duplicate compression was performed using the Cell Ranger software package (10x Genomics, CA, v2.1.0) and bc12fastq2 (Illumina, CA, v2.20) according to the manufacturer’s recommendations, using the default settings and mkfastq/count commands, respectively. Transcript alignment was performed against a human reference library generated using the Cell Ranger mkref command and the Ensembl GRCh38 v87 top-level genome FASTA and the corresponding Ensembl v87 gene GTF.

Multi-sample integration, data normalization, visualization, and differential gene expression was performed using the R package Seurat (v3.0) [46,47]. Non-viable cells and doublets were filtered from the dataset by removing cells with >10% mitochondrial RNA content, and those cells expressing fewer than 200 or more than 6,000 unique genes. Differential gene expression analysis was performed using a Wilcoxon rank sum test with Bonferroni correction to control for False Discovery Rate (FDR).

2.6. Immunoglobulin sequence analysis

Sorted B cell immunoglobulin clonotype identification, alignment, and annotation was performed using the 10x Genomics Cell Ranger pipeline. Sample demultiplexing and clonotype alignment was performed using the Cell Ranger software package (10x Genomics, CA, v2.1.0) and bc12fastq2 (Illumina, CA, v2.20) according to the manufacturer’s recommendations, using the default settings and mkfastq/vdj commands, respectively. Immunoglobulin clonotype alignment was performed against a filtered human V(D)J reference library generated using the Cell Ranger mkvdef command and the Ensembl GRCh38 v87 top-level genome FASTA and the corresponding Ensembl v87 gene GTF. Immunoglobulin clonotype visualization, diversity assessment, and analysis were performed using either the Loupe VDJ Browser (10x Genomics, CA, v2.0.0) or custom R code. Paired immunoglobulin clonotype identity, hypermutation burden, and clonal lineage was assessed using the software package BRILA, with a 15% sequence similarity threshold for clonal lineage assignment [48]. Heavy chain restricted framework and CDR hypermutation burdens were calculated using the IMGT/HighV-QUEST server [49].

2.7. Recombinant monoclonal antibody synthesis

The variable regions from the heavy and light chains of targeted immunoglobulin sequences were codon optimized, synthesized in vitro and subcloned into a pcDNA3.4 vector containing the human IgG1 Fc region by a commercial partner (Genscript). Transfection grade plasmids were purified by maxiprep and transfected into a 293-6E expression system. Cells were grown in serum-free FreeStyle 293 Expression Medium (Thermo Fisher), and the cell supernatants collected on day 6 for antibody purification. Following centrifugation and filtration, the cell culture supernatant was loaded onto an affinity purification column, washed, eluted, and buffer exchanged to the final formulation buffer (PBS). Antibody lot purity was assessed by SDS-PAGE, and the final concentration determined by 280 nm absorption. The clonotype information for all monoclonal antibodies generated as part of this study is listed in Supplemental Table 2.
2.8. Viruses

DENV-1-4 (strains Nauru/West Pac/1974, S16803, CH53489, and TVP-360, respectively) propagated in C6/36 mosquito cells were utilized for ELISA, FlowNT50, and ADE assays. Virus for ELISA was purified by ultracentrifugation through a 30% sucrose solution and the virus pellet was resuspended in PBS.

2.9. Monoclonal antibody DENV-capture ELISA

Monoclonal antibody DENV reactivity was assessed using a 4G2 DENV capture ELISA protocol. In short, 96 well NUNC MaxSorb flat-bottom plates were coated with 2 µg/ml flavivirus group-reactive mouse monoclonal antibody 4G2 (Envigo Bioproducts, Inc.) diluted in borate saline buffer. Plates were washed and blocked with 0.25% BSA + 1% Normal Goat Serum in PBS after overnight incubation. DENV-1, -2, -3 or -4 (strains Nauru/West Pac/1974, S16803, CH53489, and TVP-360) were captured for 2 h in the appropriate wells, followed by extensive washing. Serially diluted monoclonal antibody samples were incubated for 1 h at RT on the captured virus, and DENV-specific antibody binding quantified using anti-human IgG HRP (Sigma-Aldrich, SAB3701362). Secondary antibody binding was quantified using the TMB Microwell Peroxidase Substrate System (KPL, cat. #50-76-00) and Synergy HT plate reader. Single-dilution ELISA was performed using a 1/500 serum dilution for IgM and IgA assays, and a 1/20,000 serum dilution for IgG assays.

2.10. Flow cytometry-based 50% neutralization (FlowNT50) assay

Neutralizing antibody titers of monoclonal antibodies and heat-inactivated sera were determined using a flow cytometry-based neutralization assay in U937 cells expressing DC-SIGN as previously described [50,51]. Serial dilutions of antibody or sera were mixed with an equal volume of virus, diluted to achieve 10%–15% infection of cells/well, and incubated for 1 h at 37 °C. After 1 h of incubation, an equal volume of medium (RPMI-1640 supplemented with 10% FBS, 1% penicillin/streptomycin, 1% L-glutamine (200 mM), and 1% nonessential amino acids (10 mM)) containing $5 \times 10^5$ U937-DC-SIGN cells were added to each serum-virus mixture and incubated 18–20 h overnight in a 37 °C, 5% CO$_2$, humidified incubator. Following overnight incubation, the cells were fixed, permeabilized and immunostained with flavivirus group-reactive mouse monoclonal antibody 4G2 (Envigo Bioproducts, Inc.), and secondary polyclonal goat anti-mouse IgG PE-conjugated antibody (#550589, BD Biosciences). The percentage of infected cells were quantified on a BD Accuri C6 Plus flow cytometer (BD Biosciences). Data were analyzed by nonlinear regression to determine 50% neutralization titers in GraphPad Prism 8 (GraphPad Software, La Jolla, CA).

2.11. Antibody-dependent enhancement (ADE) assay

In vitro antibody-dependent enhancement of DENV infection was quantified as previously described [52]. Two-fold serial dilutions of antibody or heat-inactivated sera were incubated with virus (in sufficient amounts to infect 10%–15% of U937-DC-SIGN cells) at 1:1 for 1 h at 37 °C. This mixture was then added to a 96-well plate containing 5 $\times 10^4$ cells (K562) per well in duplicate. Cells were infected 18–20 h overnight in a 37 °C, 5% CO$_2$, humidified incubator. Processing and quantification continued as outlined in the FlowNT50 methods. Fold-infection relative to control serum was reported.

2.12. Serum IgG/IgM/IgA ELISA

DENV-reactive serum IgM/IgG/IgA levels were assessed using a modified 4G2 DENV capture ELISA protocol. 96 well NUNC MaxSorb flat-bottom plates were coated with 2 µg/ml flavivirus group-reactive mouse monoclonal antibody 4G2 (Envigo Bioproducts, Inc.) diluted in borate saline buffer. Plates were washed and blocked with 0.25% BSA + 1% Normal Goat Serum in PBS after overnight incubation. DENV-1 (strain Nauru/West Pac/1974) was captured for 2 h in separate wells, followed by extensive washing. Serially diluted serum samples were plated in duplicate and incubated for 1 h at RT on the captured virus. DENV-specific IgM/IgG/IgA levels were quantified using anti-human IgM HRP (Seracare, 5220-0328), anti-human IgG HRP (Sigma-Aldrich, SAB3701362), and anti-human IgA HRP (Biolegend, 411,002). Secondary antibody binding was quantified using the TMB Microwell Peroxidase Substrate System (KPL, cat. #50-76-00) and Synergy HT plate reader. Single-dilution ELISA was performed using a 1/500 serum dilution for IgM and IgA assays, and a 1/20,000 serum dilution for IgG assays.

2.13. Shotgun mutagenesis epitope mapping

Epitope mapping was performed as outlined previously [53]. Expression constructs for prM/E from DENV-1 (strain Nauru/West Pac/1974) and DENV-2 (strain 16681) were subjected to high-throughput ‘Shotgun Mutagenesis’ to generate comprehensive mutation libraries, with each prM/E polyprotein residue mutated to alanine (and alanine residues to serine), which were arrayed into 384-well plates (one mutation per well). For MAb library screening, plasmids encoding the DENV protein variants were transfected individually into human HEK-293T cells and allowed to express for 22 h before fixing cells in 4% paraformaldehyde (Electron Microscopy Sciences), and permeabilizing with 0.1% (w/v) saponin (Sigma-Aldrich) in PBS plus calcium and magnesium (PBS++). Cells were incubated with purified mAbs (0.1–2.0 µg/mL) diluted in 10% NGS (Sigma)/0.1% saponin, pH 9.0. Before screening, the optimal concentration was determined for each antibody, using an independent immunofluorescence titration curve against wild-type prM/E to ensure that signals were within the linear range of detection and that signal exceeded background by at least 5-fold. Antibodies were detected using 3.75 µg/mL Alexa Fluor 488-conjugated secondary antibody (Jackson ImmunoResearch Laboratories) in 10% NGS/0.1% saponin. Cells were washed three times with PBS++. Cells were washed three times with PBS++. Cells were washed two times in PBSwash. Mean cellular fluorescence was detected using an Intellicyt high throughput flow cytometer (HTFC, Intellicyt). Antibody reactivity against each mutant protein clone was calculated relative to reactivity with wild-type prM/E, by subtracting the signal from mock-transfected controls and normalizing to the signal from wild-type protein-transfected controls. The entire library data for each MAb were compared to the equivalent data from control mAbs. Mutations within clones were identified as critical to the MAB epitope if they did not support reactivity of the test MAB, but supported reactivity of other, control antibodies. This counter-screen strategy facilitates the exclusion of DENV prM/E protein mutants that are misfolded or have an expression defect.

2.14. Statistical analysis

All statistical analysis was performed using GraphPad Prism 8 Software (GraphPad Software, La Jolla, CA). A P-value < 0.05 was considered significant.

3. Results

3.1. Robust B cell plasmablast circulation occurs following primary and secondary DENV infection

Following introduction into a human host by an infected mosquito during blood meal acquisition, DENV asymptomatically replicates for 3–14 days prior to the onset of measurable viremia or any clinical manifestation of infection (Fig. 1a) [54]. As the precise interval
between when an individual is infected with DENV and when viremia/symptoms manifest is highly variable, the day of defervescence (referred here as “fever day 0”) is used as a reference point to standardize the kinetics of infection-associated immunopathology. While viremia generally resolves by fever day 0, lymphocyte activation in the periphery is generally not detectable until just before fever abatement, with maximal T and B cell activation occurring in the subsequent 3–5 days (“early convalescence”; Fig. 1a) [34,55–57]. While this kinetic profile has been described in previous studies of secondary DENV infection, limited data are available regarding the functional diversity of plasmablasts activated in response to both primary and secondary DENV infection. Therefore, to determine if there are any differences in the phenotype and immunoglobulin repertoire of plasmablasts activated in response to either primary or secondary DENV infection, we analyzed PBMC from 6 pediatric patients with serologically confirmed primary or secondary DENV infection.

We first analyzed the frequency of circulating plasmablasts during the acute (febrile) and early convalescent period by flow cytometry. Consistent with other previously published reports [34,55–57], robust plasmablast expansion was observed in PBMC of subjects with primary and secondary DENV infections. For individuals with serologically confirmed primary DENV infections, a significant population of plasmablasts was observed at two time-points per subject (1 acute, early convalescent) (Fig. 1b and Supplementary Fig. 1), and individuals with serologically confirmed secondary DENV infections exhibited significant plasmablast expansion over three time points (2 acute, 1 early convalescent) (Fig. 1c and Supplemental Fig. 1).

3.2. Identification and transcriptional characterization of DENV-elicited B cell plasmablasts by scRNAseq following primary or secondary DENV infection

To better understand the clonal and transcriptional diversity of the plasmablast repertoire elicited by primary and secondary DENV infections, we subjected sorted total CD3+ T cells and CD19+ B cells from the samples highlighted in Fig. 1 to single cell RNA sequencing (scRNAseq) analysis using the 10x Genomics platform (Supplemental Fig. 2). Cells were processed so as to recover both the unbiased 5’ gene expression profile from each cell, as well as the paired full-length immunoglobulin sequences utilizing targeted nested PCR. B cells were identified within the parental dataset and extracted for subsequent targeted analysis. Using this approach, we recovered a total of 9,027 B cells with high-quality transcriptomic and immunoreceptor data from the 6 subjects included in our analysis (range, 260–1281 per subject-time point). Four transcriptionally distinct populations of B cells were identified within the total captured B cell pool by unsupervised hierarchical clustering of the cells based on their complete gene expression profile (Fig. 2a). Differential gene expression analysis of these four statistically distinct B cell populations determined that these clusters correspond to canonical naïve B cells, memory B cells, plasmablasts, and pre-plasmablasts (Fig. 2b and c, Supplemental Table 1). Consistent with their ascribed functional potential, transcriptional imputation of the cell-cycle stage of the B cells captured in this dataset revealed that the naïve B cells, memory B cells, and plasmablasts were primarily in the G1 stage of the cell cycle, while pre-plasmablasts were exclusively in G2M/S (Fig. 2d). All
Fig. 2. Identification, quantification, and transcriptional characterization of B cell subsets by scRNAseq. Conventional B cell subset identification by unsupervised hieratical clustering of scRNAseq data from individuals with primary and secondary DENV infections. (A) UMAP projection of transcriptionally defined B cell populations from all subjects/time points included in this analysis with key populations demarcated. (B) Expression of canonical key B cell lineage genes across all cells in the dataset. (C) Heat map of most differentially expressed gene products as defined by Wilcoxon Log Rank test distinguishing naïve B cells, memory B cells, plasmablasts, and pre-plasmablasts across all subjects and time points. Data are down-sampled to 200 cells/population for visualization. (D) Imputation of cell cycle stage based on the expression of canonical cell-cycle stage associated gene products within naïve B cells, memory B cells, plasmablasts and pre-plasmablasts in the integrated dataset. (E) Abundance of naïve B cells, memory B cells, plasmablasts, and pre-plasmablasts longitudinally within each subject in the study as defined by scRNAseq. (F) Correlation analysis of the frequency of B cell plasmablasts as defined by flow cytometry and scRNAseq for all subjects and time points within the study. r² and p value calculated by linear regression.
Table 1
Subject, timepoint, and cell-type information.

| Subject      | Age / Sex | Infection | Serotype | Severity | Time point (fever day) | Total B cells | Cell type breakdown |
|--------------|-----------|-----------|----------|----------|------------------------|---------------|---------------------|
| Primary #1   | 9/M       | Primary   | DENV1    | DF       | –1                     | 430           | Naive 146 (33.9%)   |
|              |           |           |          |          |                        |               | Memory 246 (57.2%)  |
| Primary #2   | 9/M       | Primary   | DENV1    | DF       | 3                      | 525           | Plasmablast 28 (6.5%) |
|              |           |           |          |          |                        |               | Pre-plasmablast 10 (2.3%) |
| Primary #3   | 5/M       | Primary   | DENV1    | DF       | 1                      | 554           | Naive 168 (32.5%)   |
|              |           |           |          |          |                        |               | Memory 278 (52.9%)  |
|              |           |           |          |          |                        |               | Plasmablast 71 (13.5%) |
| Secondary #1 | 12/M      | Secondary | DENV3    | DHF      | 1                      | 521           | Naive 90 (16.2%)    |
|              |           |           |          |          |                        |               | Memory 228 (41.1%)  |
|              |           |           |          |          |                        |               | Plasmablast 185 (33.4%) |
| Secondary #2 | 13/F      | Secondary | DENV1    | DHF      | –1                     | 434           | Naive 215 (41.2%)   |
|              |           |           |          |          |                        |               | Memory 266 (51.1%)  |
|              |           |           |          |          |                        |               | Plasmablast 33 (6.3%) |
| Secondary #3 | 11/F      | Secondary | DENV1    | DF       | 2                      | 432           | Naive 112 (25.9%)   |
|              |           |           |          |          |                        |               | Memory 166 (34.8%)  |
|              |           |           |          |          |                        |               | Plasmablast 112 (25.9%) |
|              |           |           |          |          |                        |               | Pre-plasmablast 42 (9.7%) |

four distinct populations of B cells were identified within each subject at each timepoint, with significant population expansion contraction observed in each subject (Table 1, Fig. 2e). The frequency of plasmablasts as assessed by scRNAseq analysis of sorted CD19+ B cells correlated with the results obtained from flow-cytometry analysis of the same samples (Fig. 2f).

3.3. Isotype distribution and somatic hypermutation burden of immunoglobulins expressed by circulating B cells during primary and secondary natural DENV infection

Using the population identifiers assigned by the transcriptional profile of the B cells captured in our scRNAseq analysis, we assessed the antibody isotype diversity and immunoglobulin somatic hypermutation (SHM) burden of naïve B cells, memory B cells, and plasmablasts/pre-plasmablasts present in individuals experiencing primary or secondary DENV infections. As pre-plasmablasts and plasmablasts are direct clonal relatives and only differ in their stage of the cell cycle, (Fig. 1c and d, Supplemental Table 1) [24,58], these two populations were merged in all subsequent immunoglobulin receptor analysis. To maximize data quality in downstream evaluation, immunoglobulin sequence analysis was restricted to those cells from which a single paired full-length heavy chain and full-length light chain sequence was obtained.

Consistent with their canonical profile, the Ig sequences expressed by transcriptionally-demarcated naïve B cells were observed to be IgD or IgM in isotype with little or no SHM away from their germline configuration (Fig. 3a and d). No difference in the isotype distribution or SHM burden was observed within the naïve B cell compartment between individuals with primary versus secondary DENV infection. In contrast, transcriptionally-defined memory B cells were observed to express lgs with a diverse array of isotypes, including IgA1, IgA2, IgG1, IgG2, IgG3, IgM, and to a lesser extent IgD (Fig. 3b). However, no difference in the isotype distribution of the lgs expressed by memory B cells was observed in PBMC from individuals with primary versus secondary DENV infection. The SHM burden of the IgM, IgG, and IgA immunoglobulins expressed by memory B cells were higher than those observed in naïve B cells, with IgG and IgA class-switched memory B cells having a significantly higher SHM burden than their IgM expressing counterparts (Fig. 3e). No difference in the SHM burden was observed in the IgM, IgG, and IgA expressing memory B cell compartment between individuals with primary versus secondary DENV infection.

In contrast to the modest differences in isotype distribution and SHM burden observed in naïve and memory B cells identified in individuals with primary versus secondary DENV infections, significant differences were observed within the corresponding plasmablast compartments. Plasmablasts identified in individuals with primary DENV infections had nearly equal ratios of IgA, IgG and IgM expressing cells, while individuals experiencing a secondary DENV infection exhibited a significant IgG1 bias within the plasmablast compartment, with only modest IgA or IgM representation (Fig. 3c and Supplemental Fig. 3, Table 2). No difference was observed in the SHM burden of the IgM expressing plasmablasts between individuals with either primary or secondary DENV infections, while IgG expressing plasmablasts from individuals with a secondary DENV infection exhibited a significantly higher SHM burden than their counterparts from individuals with primary DENV infections (Fig. 3f and Supplemental Fig. 3). Surprisingly, the IgA class-switched plasmablasts in both primary and secondary DENV infections exhibited an equally high SHM burden, with the accumulation of mutations following similar patterns in both the framework regions and CDRs of the antibodies (Fig. 3f and Supplemental Figs. 3 and 4). This is despite the fact that by all other metrics the individuals classified as experiencing a primary DENV infection have never encountered DENV antigen previously. Therefore, in addition to the anticipated differences in IgM and IgG expressing plasmablasts, we contend that primary DENV infection can be characterized by a high relative proportion of extensively hypermutated IgA expressing plasmablasts. Clonal expansion was evident in the both the IgG and IgA expressing plasmablast compartments from subjects with either primary or secondary DENV infection (Supplemental Fig. 5), but no discernable trend or enrichment in Ig gene segment usage was evident (Supplemental Fig. 6).

3.4. Functional characterization of DENV-elicited plasmablast immunoglobulins

To determine the antigen specificity and functional potential of the lgs expressed by DENV-elicited plasmablasts captured in our scRNAseq dataset – especially IgA class-switched cells - we synthesized 5–20 monoclonal antibodies (mAb) from each subject included in our analysis. We selected lg sequences based on the clonal abundance of the sequence and the amount of the lg transcript expressed within the corresponding plasmablast (> 1000 copies/cell). The only exception to these criteria were VDB-5, -6, -7, and -8, which were selected based on their clonal relationship to VDB-2. Using these benchmarks, we selected and synthesized a total of 56 mAbs from DENV-elicited plasmablasts: 21 from individuals with primary DENV infection and 35 from individuals with secondary DENV infection (Supplemental Table 2). Within the selected pool of mAb candidates were antibodies with parental IgM, IgG1, IgG3, and IgA1 isotypes. The mAbs were screened for the ability to bind DENV-1, -2, -3 and -4 by...
virus-capture ELISA. In agreement with previous results obtained from individuals with experimental primary DENV infection and subjects with natural secondary infections, 38% (8/21) of the mAbs generated from individuals with a primary DENV infection exhibited DENV reactivity, while 71% (25/35) of mAbs from secondary infection plasmablasts exhibited DENV reactivity (Fig. 4a, Supplemental Table 3, Supplemental Fig. 7). The mAbs from secondary DENV infection exhibited significant serotype cross-reactivity, while moderate bias towards the infecting serotype was observed in mAbs from primary infections (Fig. 4a). Notably, 47% (8/17) of the mAbs synthesized from antibodies with a parental IgA isotype were DENV reactive. This is a slightly lower rate than what was observed for IgG1 mAbs (76.6%, 23/30), but indicates that a significant fraction of the IgA expressing plasmablasts found in circulation following acute DENV infection are indeed virus-reactive.

Any mAb exhibiting DENV binding ability was subsequently tested for DENV neutralization potential using a flow-cytometry based microneutralization assay (Fig. 4b, Supplemental Table 3, Fig. 3.

Isotype distribution and somatic hypermutation burden of immunoglobulins expressed by circulating B cells during primary and secondary natural DENV infections. Immunoglobulin sequence analysis was restricted to cells from which a full-length heavy/light chain pair was successfully isolated and annotated by scRNAseq. (A) Isotype distribution of immunoglobulins expressed by transcriptionally defined naïve B cells within all subjects at all time points captured in the dataset. (B) Isotype distribution of immunoglobulins expressed by transcriptionally defined memory B cells within all subjects in all time points captured in this analysis. Analysis split by isotype. (C) Isotype distribution of immunoglobulins expressed by transcriptionally defined plasmablast phenotype B cells from all subjects at all time points captured in this analysis. Analysis split by isotype. Error bars +/- SEM. * p < 0.05, **** p < 0.0001, one-way ANOVA.

| Subject | Time point (fever day) | Total number of plasmablast | # paired heavy/light | % recovery | Plasmablast isotype distribution (fully annotated) |
|---------|------------------------|-----------------------------|---------------------|------------|--------------------------------------------------|
|         |                        |                             |                     |            | IgM IgG IgA                                      |
| Primary #1 | −1                     | 38                          | 35                  | 92.11%     | 9 (26.5%) 4 (11.8%) 21 (61.8%)                  |
|          | 3                      | 79                          | 73                  | 94.07%     | 6 (8.3%) 55 (24.8%) 85 (38.3%)                  |
|          | 5                      | 40                          | 36                  | 90.12%     | 10 (2.6%) 24 (66.7%) 7 (19.4%)                  |
| Primary #2 | −1                     | 26                          | 21                  | 80.77%     | 2 (10.5%) 4 (21.1%) 13 (66.4%)                  |
|          | 2                      | 154                         | 133                 | 86.36%     | 39 (29.8%) 53 (40.4%) 39 (29.8%)                |
|          | 5                      | 40                          | 36                  | 90.12%     | 10 (2.6%) 24 (66.7%) 7 (19.4%)                  |
| Primary #3 | −1                     | 9                           | 8                   | 88.89%     | 3 (37.5%) 5 (62.5%) 0 (0%)                       |
|          | 1                      | 597                         | 642                 | 92.11%     | 17 (2.7%) 560 (87.4%) 64 (10%)                  |
|          | 5                      | 37                          | 33                  | 89.19%     | 6 (18.8%) 6 (18.8%) 20 (62.5%)                  |
| Secondary #1 | 0                     | 435                         | 396                 | 91.03%     | 12 (3%) 352 (89.1%) 31 (7.8%)                   |
|          | 1                      | 549                         | 494                 | 89.98%     | 10 (2%) 428 (86.8%) 55 (11.2%)                  |
|          | 7                      | 118                         | 104                 | 88.14%     | 3 (2.9%) 61 (59.2%) 39 (37.9%)                  |
| Secondary #2 | −1                     | 697                         | 642                 | 92.11%     | 17 (2.7%) 560 (87.4%) 64 (10%)                  |
|          | 1                      | 61                          | 54                  | 88.89%     | 3 (37.5%) 5 (62.5%) 0 (0%)                       |
|          | 5                      | 155                         | 141                 | 90.97%     | 18 (12.8%) 82 (58.2%) 41 (28.1%)                |
| Secondary #3 | 0                     | 83                          | 75                  | 90.36%     | 11 (14.7%) 40 (53.3%) 24 (32%)                  |
|          | 1                      | 155                         | 141                 | 90.97%     | 18 (12.8%) 82 (58.2%) 41 (28.1%)                |
Fig. 4. DENV-elicited plasmablast-derived monoclonal antibody specificity and functional activity. (A) Heat map summary of the specificity and avidity of 61 mAbs derived from immunoglobulin sequences extracted from the scRNAseq analysis of plasmablast-phenotype B cells from individuals experiencing primary or secondary natural DENV infections. (B) Heat map summary of the neutralization potential (IC$_{50}$) of 33 mAbs derived from immunoglobulin sequences extracted from the scRNAseq analysis of plasmablast-phenotype B cells from individuals experiencing primary or secondary natural DENV infections. Only mAbs exhibiting DENV specificity by ELISA (EC$_{50}$ < 5 μg/ml) were assessed by FlowNT. (C) Scatter plot of reciprocal DENV-1, -2, -3, and -4 IC$_{50}$/EC$_{50}$ values calculated for all plasmablast-derived mAbs generated in this study which exhibited an EC$_{50}$ value of <5 μg/ml for at least one viral serotype. Symbol color indicates the virus serotype tested (DENV-1 in blue, DENV-2 in red, DENV-3 in green, DENV-4 in purple). Symbol fill indicates if the antibody was isolated from an individual with a primary (open) or secondary (closed) DENV infection. (D) ADE potential of mAb VDB33 against DENV-1, -2, -3, and -4. All values are shown as fold-increased infection compared to no antibody. Dashed line indicates the mAb IC$_{50}$ as determined by FlowNT using the same viral preparation as used in the ADE assay. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
Supplemental Fig. 7), mAbs from individuals experiencing primary DENV infections exhibited relatively poor neutralization potential, with only 37.5% (3/8) DENV reactive mAbs exhibiting any neutralization activity. In contrast, 92% (23/25) of the DENV-reactive mAbs generated from individuals experiencing a secondary DENV infection exhibited viral neutralization, with 60% (15/25) capable of neutralizing all 4 DENV serotypes to some degree. Of note, mAbs generated from individuals experiencing secondary DENV infections often exhibited better binding and neutralization activity towards serotypes other than the infecting virus at the time of sample collection, potentially providing insight into the virologic exposure history of the individual.

While the neutralization potential of a DENV-reactive mAb provides some indication of its theoretical in vivo efficacy, the ratio between neutralization potential and avidity is an important secondary consideration. Antibodies requiring high viral occupancy in order to neutralize viral infectivity – reflected in a high EC$_{50}$/IC$_{50}$ ratio – are potentially capable of facilitating ADE in situations where they are present at a concentration where they can bind a virus but not neutralize it if they possess an Fc domain capable of interacting with Fc receptors expressed by DENV-susceptible leukocytes. Indeed, while some relationship is apparent between the EC$_{50}$ and IC$_{50}$ values of the mAbs analyzed (Fig. 4c), many mAbs sit above a theoretical linear line of identity for some DENV serotypes, and therefore may be capable of facilitating ADE if they possessed an appropriate Fc domain. Of particular note, mAb VDB33 demonstrated a uniquely high EC$_{50}$/IC$_{50}$ ratio, as it potently binds all four DENV serotypes (EC$_{50}$ 2.32–5.07 ng/ml) while showing poor neutralization potential (IC$_{50}$ 1181 to >10,000 ng/ml). Shogun mutagenesis epitope mapping revealed that this mAb bound the fusion loop of the DENV E protein, the canonical binding site for antibodies thought to facilitate ADE (Fig. 5a). Indeed, when tested in an in vitro ADE assay, VDB33 exhibited dramatic ADE potential, especially against DENV3 (Fig. 4d). This is notable, as the individual from which VDB33 was obtained was experiencing a secondary DENV-3 infection (resulting in DHF) at the time of sample collection.

A contemporaneous serum sample from this individual exhibited a very similar ADE profile as VDB33 (Supplemental Fig. 8). While it is not possible to draw a direct causal connection between this plasmablast-derived antibody (or its relatives) and the clinical manifestation of DENV infection observed in the individual from which it was isolated using these data, these findings underscore the utility of assessing the molecular level detail of DENV-elicited humoral immunity in future studies.

3.5. DENV-reactive serum immunoglobulin isotype diversity in primary and secondary DENV infection

To determine if the DENV-elicited immunoglobulin isotype diversity observed in the plasmablast compartment was reflected in the contemporaneous antibodies in circulation, we performed DENV-capture ELISA on serum samples from individuals with either primary or secondary DENV infection and quantified the relative proportions of DENV-reactive IgM, IgG, and IgA present therein. To provide temporal resolution to the analysis, four time-points per subject were analyzed: two samples during acute viral infection (acute 1, acute 2), one sample obtained 3–7 days following fever abatement (early-convalescent), and one sample collected 6 months post infection (Fig. 5, Supplemental Table 4).

Relatively modest DENV-reactive Ig levels were observed in acute serum samples obtained from individuals with either primary (Fig. 5a) or secondary (Fig. 5b) DENV-1 infections. However, a dramatic increase in circulating DENV-reactive IgS was observed in early-convalescent serum samples from both primary and secondary DENV infections (Fig. 5a and b). Notably, while DENV-reactive IgA was detectable in all subjects at this time point, IgA represented a much higher fraction of the total DENV-reactive Ig pool in individuals experiencing a primary DENV infection, while IgG production dominated in secondary DENV-1 infections (Fig. 5a and b). Total DENV-reactive Ig titers returned to more modest levels in all subjects by 6 months post infection, with little DENV-reactive IgA observed (Fig. 5a and b). Therefore, despite the fact that DENV-reactive IgA represent an appreciable fraction of the total DENV-reactive Ig in circulation just following the resolution of infection – especially in individuals experiencing a primary DENV infection - DENV-reactive IgA does not appear to persist in circulation.

To further validate these observations, single-dilution ELISA analysis was performed on acute and convalescent serum samples from an additional 9 individuals experiencing either a primary or secondary DENV-1 infection (Supplemental Table 5). As was observed in the limiting-dilution ELISA analysis, relatively little DENV-reactive IgG or IgA was observed in acute infection time points from subjects with either primary or secondary infection (Fig. 5c). However, the DENV-reactive IgG and IgA signals increased dramatically in the corresponding early convalescent serum samples, with IgA comprising a much higher fraction of the total DENV-reactive Ig signal in samples from primary DENV-1 infections than from samples from secondary infections (Fig. 5d). These data confirm that DENV-reactive IgA represents a significant fraction of the total DENV-reactive Ig pool following the resolution of acute infection, especially in individuals experiencing a primary DENV infection.

3.6. Epitope specificity of plasmablast-derived/DENV-reactive mAbs

In light of the unexpectedly high frequency of DENV-reactive IgA clones observed in our dataset and the pervasiveness of DENV-reactive IgA in contemporaneous serum samples—especially in individuals experiencing a primary DENV infection - we endeavored to determine if DENV-reactive IgA lgs recognize an unconventional epitope on DENV or if their antigen specificity mirrored that of other “conventional” DENV-reactive antibody isotypes. To this end, we utilized shotgun mutagenesis epitope mapping of plasmablast-derived mAbs.

To further validate these observations, single-dilution ELISA analysis was performed on acute and convalescent serum samples from individuals with either primary or secondary DENV infections. Of note, the DENV-reactive IgA signals increased dramatically in the corresponding early convalescent serum samples, with IgA comprising a much higher fraction of the total DENV-reactive Ig signal in samples from primary DENV-1 infections than from samples from secondary infections (Fig. 5d). These data confirm that DENV-reactive IgA represents a significant fraction of the total DENV-reactive Ig pool following the resolution of acute infection, especially in individuals experiencing a primary DENV infection.

The three parentally IgG1 DENV-reactive mAbs picked for epitope mapping were found to exclusively interact with the fusion loop region of the DENV E protein (Fig. 6a, Supplemental Table 6). Three parentally IgM antisera generated from subjects with primary or secondary DENV infections were observed to either bind the DENV E protein fusion loop (VDB31/VDB50) or the DENV E protein domain II hinge region (VDB37) (Fig. 6b, Supplemental Table 6). These results indicate that DENV-reactive IgA antibodies are capable of recognizing multiple distinct epitopes on the DENV E protein even in individuals who have not previously encountered DENV, and that DENV-reactive IgG and IgA isotype antibodies can overlap in their epitope specificity.

4. Discussion

In this study, we utilized single cell RNA sequencing technology to assess the clonal and transcriptional diversity of plasmablasts elicited in response to primary and secondary DENV infections in an unbiased and high-throughput fashion. We found that an unexpectedly high proportion of DENV-elicited plasmablasts expressed IgA lgs, notably in individuals experiencing a primary DENV infection. These IgA immunoglobulins were extensively hypermutated, even in individuals with serologically confirmed primary DENV infection. Utilizing a combination of conventional biochemical assays and high-throughput shotgun mutagenesis, we determined that DENV-reactive IgA antibodies represented a significant fraction of DENV-reactive lgs generated in response to DENV infection, and that they have a overlapping epitope specificity to “conventional” DENV-reactive IgM and IgG class-switched antibodies reported in the literature. These results provide insight into the molecular-level diversity of DENV-elicited humoral immunity, and suggest a hitherto unknown mechanism of DENV antigen presentation.
Fig. 5. Kinetics and magnitude of DENV-reactive IgM/IgG/IgA in circulation following primary or secondary DENV infection. (A) Limiting dilution DENV-1 capture ELISA analysis of relative virus-reactive IgG, IgA, IgM levels present in serum obtained from three subjects with serologically- and PCR-confirmed primary DENV-1 infections. (B) Limiting dilution DENV-1 capture ELISA analysis of relative virus-reactive IgG, IgA, IgM levels present in serum obtained from three subjects with serologically- and PCR-confirmed secondary DENV-1 infections. (C) Single-dilution DENV-1 capture ELISA analysis of the relative virus-reactive IgG, IgA, IgM levels present in acute infection serum samples obtained from subjects with serologically and PCR confirmed primary (n = 5) and secondary (n = 4) DENV-1 infections. (D) Single-dilution DENV-1 capture ELISA analysis of the relative virus-reactive IgG, IgA, IgM levels present in early-convalescent serum samples obtained from subjects with serologically and PCR confirmed primary and secondary DENV-1 infections. Error bars indicate +/- SEM. ** p < 0.01, one-way ANOVA.
unappreciated role for DENV-reactive IgA in the humoral response to DENV infection, especially in the setting of primary DENV infection.

In addition to revealing a potential role for DENV-reactive IgA in the setting of acute DENV infection, the data presented in this study provide molecular-level resolution and reinforcement to several fundamental features of DENV-elicted humoral immunity. This includes the observations that the majority of the individual antibody clones elicited by a primary DENV infection exhibit relatively poor neutralization activity, and the serotype cross-reactive humoral immune profile associated with secondary DENV infections extends to the level of individual antibody clones. Our findings must be considered in light of several limitations, however. These data were generated by the analysis of samples obtained from a small number of children, most of whom were experiencing DENV-1 infections. Although viral tropism and clinical manifestations of the different DENV serotypes are thought to be similar, some caution must be used in extrapolating these observations to other age groups and to other DENV serotypes. Immunological history and host/pathogen genetics both play significant roles in determining the pathogenesis and cellular/molecular level response to DENV infection [61]. However, tools such as scRNA-seq and other high-throughput/high-resolution modalities can now reliably capture many of these complex variables, hopefully providing better insight and more actionable information in the setting of acute DENV infection.

The induction of virus-specific IgA in response to acute infection is not an uncommon phenomenon, especially for viruses that exhibit significant mucosal tropism [62]. The unique cytokine environment found in mucosal tissues is thought to provide the necessary immunological cues to nascently activated B cells to facilitate the IgA1 antibody class-switch event [63]. However, DENV has no known mucosal involvement either during transmission or replication. DENV-reactive IgA generated in response to acute infection is not without precedent, although much of the previous work on IgA in the context of DENV infection has focused on the feasibility of utilizing urine/saliva restricted DENV-reactive IgA as a non-invasive diagnostic [64–67]. A single paper has compared directly the titers of IgA in primary and secondary DENV infections [67], and no previous study has assessed the prevalence of IgA class-switched plasmablasts present following acute DENV infection. The paucity of previous data on the abundance and specificity of DENV-specific IgA following acute viral infection is partially attributable to the relatively brief window during which significant levels of these DENV-reactive IgA antibodies are present in serum following primary DENV infection (only in early convalescent serum samples in our study). In addition, the majority of previous studies assessing the molecular diversity of DENV-elicted humoral immunity also intentionally limited their analysis to include IgG and IgM antibodies and/or genes [32,35–37].

Although the precise role – if any - that DENV-elicted IgA plays in influencing the immunopathogenesis of DENV is still unclear, it is noteworthy that while we have demonstrated that the antigen-binding domains of DENV-elicted IgA clones can effectively neutralize DENV infectivity, the restricted expression profile of IgA Fc receptors implies that DENV-reactive IgA clones cannot easily facilitate classical antibody-dependent immune enhancement of DENV infection. Therefore, much like IgM, DENV-reactive IgA might contribute neutralizing activity to DENV-reactive serum, while antagonizing IgG1 mediated ADE. As the average neutralization potential of the DENV-reactive IgA clones tested in this study varied widely, the ability of these clones to neutralize DENV in a complex humoral environment may be limited, even if these antibodies were present at an equimolar ratio as their IgG counterparts during the resolution phase of a DENV infection. However, the epitope mapping data presented herein suggests that at least some DENV-reactive IgA clones are capable of binding the same epitopes as their IgG1 counterparts. Therefore, it is possible that DENV-specific IgA may limit intra-host viral propagation during the resolution phase of an acute DENV infection by competing with infection-enhancing DENV-specific IgG and preventing internalization of infectious virions through IgG Fc receptors. This scenario does not necessitate that an IgA opsonized virus is less infectious than a un-opsonized virus in the context of Fc-receptor independent infection, but it does suggest that an IgA opsonized virus might be less infectious than an IgG1 opsonized virus from the standpoint of a cell expressing an IgG1-reactive Fc receptor.

In light of the consistently high SHM burden observed in IgA expressing plasmablasts circulating in response to primary DENV infection, we additionally propose that the majority of these cells are reactivated memory B cells that were initially stimulated by an antigen other than DENV. Memory B cells have both a numerical and functional advantage over naïve B cells upon antigen engagement, and a relatively small population of memory B cells that happen to cross-react with a cognate antigen could certainly be expected to contribute to a primary response to infection. Furthermore, it is perhaps unsurprising that a small number of IgA class-switched memory B cells with some degree of DENV cross reactivity exist in any individual prior to a primary DENV infection, as it has been suggested that more than 70% of the total antibody production in humans is IgA in isotype, albeit mostly at mucosal surfaces [68]. Therefore, it is possible that all primary pathogen infections include a significant IgA component, as the largest pool of pre-existing memory B cells that is subsequently capable of responding/cross-reacting with a “novel” antigen would be IgA in isotype. This interpretation follows the concept of “original antigen sin”, wherein a previous infection has a direct impact on the specificity and efficacy of immunity generated in response to a subsequent infection/challenge, but extends the concept to primary DENV infection [69]. This indelible feature of pre-existing immunity can also be observed in our data from secondary DENV infections, wherein antibody specificity is occasionally biased towards DENV serotypes other than the infecting type. These data suggest that immunological memory generated in response to completely different stimuli may impact future immune responses simply because of putative antigenic similarity.

Declaration of Competing Interest

A.T.W reports grants from Military Infectious Disease Research Program, during the conduct of the study.

Fig. 6. Binding specificity of plasmablast-derived/DENV-reactive monoclonal antibodies. (A) Shotgun mutagenesis epitope mapping data from IgG1-derived mAbs. (B) Shotgun mutagenesis epitope mapping data from IgA1-derived mAbs. Color scheme for indicated residues: green = fusion loop, yellow = domain II hinge region. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
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The authors declare that all data supporting the findings of this study are available within this article and its Supplementary Information files, or from the corresponding author upon reasonable request. Single-cell RNAseq gene expression data have been deposited in the Gene Expression Omnibus database (GSE145307).

The opinions or assertions contained herein are the private views of the authors and are not to be construed as reflecting the official views of the US Army or the US Department of Defense, or the National Institutes of Health. Material has been reviewed by the Walter Reed Army Institute of Research. There is no objection to its presentation and/or publication. The investigators have adhered to the policies for protection of human subjects as prescribed in AR 70–25. 

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Data availability

Supplementary material associated with this article can be found 10.1016/j.ebiom.2020.102733.

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