Serum Metabolome and Lipidome Changes in Adult Patients with Primary Dengue Infection

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Abstract

Background: Dengue virus (DENV) is the most widespread arbovirus with an estimated 100 million infections occurring every year. Endemic in the tropical and subtropical areas of the world, dengue fever/dengue hemorrhagic fever (DF/DHF) is emerging as a major global health concern. The complex array of concurrent host physiologic changes has hampered a complete understanding of underlying molecular mechanisms of dengue pathogenesis.

Methodology/Principle Findings: Systems level characterization of serum metabolome and lipidome of adult DF patients at early febrile, defervescence, and convalescent stages of DENV infection was performed using liquid chromatography- and gas chromatography-mass spectrometry. The tractability of following metabolite and lipid changes in a relatively large sample size (n = 44) across three prominent infection stages allowed the identification of critical physiologic changes that coincided with the different stages. Sixty differential metabolites were identified in our metabolomics analysis and the main metabolite classes were free fatty acids, acylcarnitines, phospholipids, and amino acids. Major perturbed metabolic pathways included fatty acid biosynthesis and β-oxidation, phospholipid catabolism, steroid hormone pathway, etc., suggesting the multifactorial nature of human host responses. Analysis of phospholipids and sphingolipids verified the temporal trends and revealed association with lymphocytes and platelets numbers. These metabolites were significantly perturbed during the early stages, and normalized to control levels at convalescent stage, suggesting their potential utility as prognostic markers.

Conclusions/Significance: DENV infection causes temporally distinct serum metabolome and lipidome changes, and many of the differential metabolites are involved in acute inflammatory responses. Our global analyses revealed early anti-inflammatory responses working in concert to modulate early pro-inflammatory processes, thus preventing the host from development of pathologies by excessive or prolonged inflammation. This study is the first example of how anomic approach can divulge the extensive, concurrent, and dynamic host responses elicited by DENV and offers plausible physiological insights to why DF is self limiting.

Introduction

Dengue virus (DENV) is a member of the Flavivirinae genus of the Flaviviridae family and is classified into four distinct serotypes. It is estimated that 100 million dengue infections occur worldwide every year, ranging from acute dengue fever (DF) to life-threatening dengue haemorrhagic fever/dengue shock syndrome (DHF/DSS) [1,2]. Dengue is emerging as a global health concern with the annual average number of DF/DHF cases reported to World Health Organization increasing dramatically in the past two decades [3].

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Four serotypes are now circulating in Asia, Africa and the Americas [3].

DF is generally self-limiting, but its symptoms can be debilitating and cause considerable incapacitating morbidity, which have a significant health and economic toll in the society. In a small percentage of patients, DF evolves to the more severe forms of DHF and DSS, which are characterized by abnormal hemostasis, vascular leakage and liver damage. There is currently no specific antiviral therapy or vaccine available for DF or DHF/DSS. Furthermore, the underlying molecular mechanisms of DENV pathogenesis are still unclear and why DF resolves in time.
Author Summary

Dengue virus is the most widespread arbovirus and a major public health threat in the tropical and subtropical areas of the world. As yet, little is known about the molecular mechanisms underlying infection, and there is no specific treatment or vaccine that is currently effective against the disease. Metabolomics and lipidomics provide global views of metabolome and lipidome landscapes and implicate metabolic to disease phenotype. We performed serum metabolic and lipidomic profiling on a cohort of dengue patients with three sampling time points at early febrile, defervescence, and convalescent stages via mass spectrometry-based analytical platforms. Compared with healthy subjects, approximately two hundred metabolites showed significant difference in dengue patients, and 60 were identified. This study revealed that in primary dengue infection, the host metabolome is tightly regulated, with active, early anti-inflammatory processes modulating the pre-inflammatory processes, suggesting the self-limiting phenotype of dengue fever. Major perturbed metabolic pathways included fatty acid biosynthesis, fatty acid β-oxidation, phospholipid catabolism, steroid hormone pathway, etc. This represents a first report on the characterization of the serum metabolome and significantly advances our understanding on host and dengue virus interactions. These differential metabolites have the potential as biomarkers for disease monitoring and evaluation of therapeutic interventions.

is largely neglected. Therefore, the pathological differences between the severe DHF/DSS and the mild, self-limiting, DF suggest differential virus-host interactions in the susceptibility to the disease. Both viral and host immune factors seem to be involved, but the role of each is not fully understood [4]. Meanwhile, the lack of an appropriate animal model significantly increases the challenges in the study of DENV pathogenesis.

Metabolomics and lipidomics are rapidly emerging fields of ‘omics’ research that aim to study the global changes of small molecule metabolites and lipids in biological systems in response to biological stimuli or perturbations [5]. Metabolites are the end products of cellular regulatory processes, forming a link between molecular changes and phenotype, and therefore reflect the physiological state of a cell, tissue or organism at a point in time. Lipidomics covers the subset of lipid constituents, which are known to be involved in signalling and infection biology. Cellular homeostasis is interrupted under disease conditions, and the human body would attempt to maintain a basal, internal environment by increasing or decreasing levels of certain endogenous metabolites. Thus, by providing a snapshot of the metabolic status of an organism, metabolomics holds the promise of finding metabolic pathways related to disease processes [6,7]. Metabolomics has been applied to infectious diseases to study host-pathogen interactions [8,9,10]. Similarly, there has been enormous interest in how lipids and their metabolism influence Flaviviridae viral infection, especially from the perspective of microbe lipid usage for virus entry, replication and release [11,12,13]. As we redefine the manner in which we understand the complex and dynamic virus-host interactions of dengue infection in vivo, it necessitates capturing the global changes rather than isolated ones. However, few omics studies exist that describe how the human host response biochemically and physiologically during DENV and other flavivirus infection and how it resolves without intervention [14,15]. Therefore we turned our attention to examine serum metabolites and lipids using a comprehensive approach to globally capture the human responses to DF.

In our previous study, a systematic characterization of serum cytokines, proteome, and markers of macrophage and neutrophil activity was reported from a subset of longitudinally enrolled adult patients with primary dengue infections [16]. In the present study, both untargeted metabolomics and targeted lipidomics on the same serum sample set were conducted with the aim to identify metabolic pathways linked to disease progression and understand the mechanisms of DENV infection. Our results showed that DENV infection caused significant serum metabolome and lipidome-wide changes in DF patients. Sixty differential metabolites were identified in metabolomics analysis and detailed examination of metabolic pathways changes revealed that in primary DENV infections, acute inflammatory responses were quickly met with active pro-resolution counterbalances to restore homeostasis. Our studies also implicated the plausible association of sphingomyelins with CD8+ lymphocyte activation during the early febrile period, and plasmalogen phosphatidylethanolamines and lysophosphatidylethanolamines with platelet numbers during the later phases of defervescence and convalescence. We propose that early widespread host changes in the metabolome and lipidome play an important physiological component in defining the self-limiting outcome of DF.

Materials and Methods

Ethics statement

The early dengue infection and outcome (EDEN) study is a multi-center longitudinal study, which prospectively recruits and follows-up adult dengue patients in Singapore, through early febrile, defervescence as well as convalescence stages of the disease [17]. Enrolment of all eligible individuals was based on written informed consent and the protocols were approved by the National Healthcare group (DSRB B/05/013). All samples were anonymized.

Clinical samples

The details of patient recruitment, sample collection and the study protocols of EDED study have been described earlier [18]. In brief, adult patients (>21 years) presenting with acute onset fever (≥38.0°C for less than 72 hours) without rhinitis or other clinical alternatives were included in the study (Visit 1; febrile). Initial dengue diagnosis was made by real time RT-PCR and followed by serology and subsequent serotyping by virus isolation and immunofluorescence using serotype specific monoclonal antibodies (ATCC: HB46-49). Venous blood samples were also collected at fever day 4 to 7 (Visit 2; defervescence) and weeks 3 to 4 (Visit 3; convalescence), aliquoted and frozen at −80°C. ‘Fever day’ here refers to number of days post onset of fever. Classification of DF was made based on the guidelines provided by the World Health Organisation [19]. In brief, acute febrile patients positive for dengue with one or more of the following symptoms: headache, retro orbital pain, myalgia, rash, leukopenia, hemorrhage were classified as DF. Of the 133 dengue patients that were finally enrolled in this study (September 2005–October 2006), 44 DF patients tested negative for dengue IgG antibodies in the acute sera, using a commercially obtained ELISA (PanBio, Brisbane, Australia). These patients were deemed to have primary DENV infection, all of which were included in this study. A detailed hematological and virological analysis was also performed. Additionally we used serum samples from 50 asymptomatic age-matched healthy subjects participated during a hospital staff annual examination as controls (Table 1). This study was...
Table 1. Demographics of healthy subjects and dengue patients enrolled in this study.

| Patient Groups | Serotype | Age    | Gender | Blood sampling time* |
|----------------|----------|--------|--------|----------------------|
| DF (n = 44)    | D1 (54.5%) | 39.0±13.0 | 27 (male) | 46±20 hours (Visit 1) |
|                | D2 (2.3%)  | 17 (female) | 73±31 hours (Visit 2) |
|                | D3 (43.2%) | 16±5 days (Visit 3) |
| Controls (n = 50) | NA       | 40.0±18.0 | 14 (male) | NA |
|                |          |        |        | 36 (female) |

*Average time from fever to phlebotomy (Visit 1), between Visits 1 & 2 (Visit 2) and between Visits 2 & 3 (Visit 3).

Serum sample preparation

For metabolomics analysis, a volume of 50 μL from each serum sample was thawed at 4°C. Serum proteins were precipitated with 200 μL ice-cold methanol, which contained 10 μg/mL 9-fluorenylmethoxycarbonyl-glycine as an internal standard. After vortexing, the mixture was centrifuged at 16,000 rpm for 10 minutes at 4°C and the supernatant was divided into two parts, one for liquid chromatography/mass spectrometry (LC-MS) analysis and the other for gas chromatography/mass spectrometry (GC-MS) analysis. All samples were kept at 4°C and analyzed within 48 h. In order to prevent batch effect, the assays were conducted in random manner. Quality control (QC) samples were prepared by mixing equal amounts of serum samples from 10 healthy subjects and 10 DF patients at all the three time points.

For lipidomics analysis, due to limited sample available, sera from three patients at each visit were randomly pooled (14 pools total). Lipids were extracted from serum using a modified Bligh and Dyer method [20]. Briefly, 900 μL of chloroform-methanol, 1:2 (v/v) was added to 100 μL serum. After 20 min vortexing and incubation at 4°C, 300 μL of chloroform and 300 μL of ddH2O were added to the mixture and centrifuged at 9000 rpm, 4°C for 2 min. Lipids were then recovered from the lower organic phase after centrifugation. Subsequently, 500 μL of chloroform was added, vortexed at 4°C for 20 min. After centrifugation, lipids were recovered from the organic phase and combined with the previous fraction. The lipid extracts were vacuum-dried, stored at −80°C and analyzed within a week.

Metabolomics analysis by LC-MS

The supernatant fraction from sample preparation step was analyzed using Agilent 1290 ultrahigh pressure liquid chromatography system (Waldbronn, Germany) equipped with a 6520 Q-TOF mass detector managed by a MassHunter workstation. The column used for the separation was an Agilent rapid resolution HT Zorbax SB-C18 (2.1×50 mm, 1.8 μm; Agilent Technologies, Santa Clara, CA, USA). The oven temperature was set at 50°C. The gradient elution involved a mobile phase consisting of [A] 0.1% formic acid in water and [B] 0.1% formic acid in methanol. The initial condition was set at 5% B. A 7 min linear gradient to 70% B was applied, followed by a 12 min gradient to 100% B which was held for 3 min, then returned to starting conditions over 0.1 min. Flow rate was set at 0.4 ml/min, and 5 μL of samples was injected. The electrospray ionization mass spectra were acquired in both positive and negative ion mode. Mass data were collected between m/z 100 and 1000 at a rate of two scans per second. The ion spray voltage was set at 4,000 V, and the heated capillary temperature was maintained at 350°C. The drying gas and nebulizer nitrogen gas flow rates were 12.0 L/min and 50 psi, respectively. Two reference masses were continuously infused to the system to allow constant mass correction during the run: m/z 121.0509 (C3H5N+O) and m/z 922.0098 (C11H10O5N2P3F25+). The stability of the LC-MS method was examined and evaluated by a subset of peaks covering a range of masses, intensities, and retention times across the QC samples (Table S1). Variations of retention time and m/z values of the peaks were less than 0.2 min and 10 mDa, respectively, and the relative standard deviations of peak areas were below 20%, indicating good reproducibility and stability of the chromatographic separation and mass accuracy of mass measurement during the whole sequence.

Metabolomics analysis by GC-MS

100 μL supernatant fraction from sample preparation step was dried under nitrogen and derivatised with 150 μL methoxamine (50 μg/mL in pyridine, 37°C 2 h) followed by 150 μL N,N-dimethyl-trimethylsilyl-trifluoroacetamide (37°C 16 h). After centrifugation (4°C, 6000 rpm×1 min), the supernatant fraction was injected into GC-MS. The derivatised sample (1.0 μL) was introduced by splitless injection with an Agilent 7890 Series autosampler into an Agilent 6890 GC system equipped with a fused-silica capillary column HP-5MSi (30 m×0.25 mm i.d., 0.25 μm film thickness) as reported previously [21]. The inlet temperature was set at 250°C. Helium was used as the carrier gas at a constant flow rate of 1.0 ml/min. The column effluent was introduced into the ion source of an Agilent Mass selective detector. The transfer line temperature was set at 280°C and the ion source temperature at 230°C. The mass spectrometer was operated in electron impact mode (70 eV). Data acquisition was performed in full scan mode from m/z 50 to 550 with a scan time of 0.5 sec. The compounds were identified by comparison of mass spectra and retention time with those of reference standards, and those available in libraries (NIST 0.5). A summary of the workflow utilized in metabolomics study is shown in Figure S1.

Lipidomics analysis by LC-MS/MS

To facilitate interrogation of the serum lipidome in a high-throughput, targeted and accurate fashion, we employed LC-MS/MS in Multiple Reaction Monitoring (MRM) mode to survey more than 200 individual glycerophospholipids and sphingolipids simultaneously at linear dynamic ranges of 0.1 to 100 ng/mL. Prior to ESI-LC-MS/MS, each lipid extract was reconstituted in 400 μL of chloroform-methanol (1:1 v/v). LC-MS/MS via Triple Quadrupole 6460 with Jet StreamTM (Agilent Technologies) was used for the quantification of glycerophospholipids and sphingolipids (total >200 molecular species). C-18 reversed phase LC (Zorbax Eclipse 2.1×50 mm I.D., 1.8 μm; Agilent Technologies) was used to separate 5 μL reconstituted lipids at 400 μL/min before entering the mass spectrometer. The column thermostat and autosampler temperatures were maintained at 40°C and 6°C, respectively. The optimized mobile phase consisted of 5 mM ammonium acetate in water (solvent A) and 5 mM ammonium acetate in methanol (solvent B). The gradient of the samples was: 0–2 min, 60–100% B; 2–7 min, maintained at 100% B; 7–9 min, 100–60% B and 9–11 min, 60% for column re-equilibration before the next injection. Positive ionization mode [M+H]+ MRM
was based on product ion $m/z$ 264.4 [sphingosine-H$_2$O]$^+$ and 266.4 [sphinganine-H$_2$O]$^+$ for ceramide (Cer), monohexosykeramide (MHexCer), lactosykeramide (LacCer) and ceramide-1-phosphate (C1P), and $m/z$ 184.1 [phospholipine]$^+$ for sphingomyelin (SM). Negative ionization [M-H]$^-$ post hoc Multiple Comparison one-way ANOVA was used for paired analysis with the Bonferroni limits of quantification and relative standard deviation of the quantification. The linear dynamic ranges, limits of detection, sera, spiked into extracted samples and used in absolute Polar Lipids, Alabaster, Al) were verified of their low abundance in PA 14:0, PC 14:0, PE 14:0, PG 14:0, PS 14:0 and PI 14:0; Avanti specific parent and product ion pairs was employed for various lipids are shown in Table S2. Known concentrations (20–100 ng/mL) of non-naturally occurring internal standards, namely Cer d18:1/17:0, GlcCer d18:1/8:0, LacCer d18:1/8:0, SM d18:1/12:0, C1P d18:1/17:0, PA 14:0, PC 14:0, PE 14:0, PG 14:0, PS 14:0 and PI 14:0; Avanti Polar Lipids, Alabaster, Al) were verified of their low abundance in sera, spiked into extracted samples and used in absolute quantification. The linear dynamic ranges, limits of detection, limits of quantification and relative standard deviation of the various lipids are shown in Table S3. The effect of acyl length on signal intensity [23,24] was compensated by adjusting the collision energy and fragmentor voltage (Table S2). A summary of the workflow utilized in the lipidomics study is shown in Figure S2.

Data analysis

For metabolomics analysis, raw spectrometric data were converted to mzData (LC-MS/MS) and NetCDF (GC-MS) formats via Masshunter (Agilent, US) and input to open-source software MZmine 2.0 for peak finding, peak alignment and peak normalization across all samples [25]. The MZmine 2.0 report table was exported into SIMCA-P software (version 11.0; Umetrics AB, Umeå, Sweden) for multivariate data analysis. Data sets were mean-centered and pareto-scaled to adjust the importance of high and low abundance metabolites to an equal level before carrying out statistical analyses. Unsupervised multivariate analysis principle component analysis (PCA) was first performed on LC-MS data as an unbiased statistical method to observe intrinsic trends within healthy controls and DF patients at different visits. To achieve the maximum separation among the groups, supervised multivariate analysis orthogonal partial least-squares discriminant analysis (OPLS-DA) was sequentially applied. Potential differential metabolites were selected according to the Variable Importance in the Projection (VIP) values and the variables with VIP$>1$ were considered to be influential for the separation of samples in OPLS-DA analysis. In addition, the Kruskal-Wallis test was performed to determine if the differential metabolites obtained from OPLS-DA modelling were statistically significant ($p<0.05$) among groups at the univariate analysis level.

For lipidomics analysis, undetectable lipids were replaced with 1/10 of the lowest non-zero value measured. In cases where 70% of data are missing from patients, lipids were excluded. Missing data represents 1.8% of the data. In total 207 lipids remained for further analysis. Comparisons between groups were achieved using non-parametric analysis namely two-tailed Mann-Whitney tests and Kruskal-Wallis tests with Dunn’s post hoc analysis. Repeated one-way ANOVA was used for paired analysis with the Bonferroni Multiple Comparison post hoc Test.

Compound identification and pathways analysis

The structure identification of the differential metabolites was based on a previously described strategy [26]. The identification process is illustrated here with hexanoylcarnitine, one of the ten differential acylcarnitines. First, the element composition C$_{13}$H$_{25}$NO$_4$ of the $m/z$ 260.18 ion was calculated based on the exact mass, the nitrogen rule and the isotope pattern by Masshunter software from Agilent (Figure S3A). Then, the elemental composition and exact mass were used for open source database searching, including LIPIDMAPS [http://www.lipidmaps.org/], HMDB [http://www.hmdb.ca/], METLIN [http://metlin.scripps.edu/] and MassBank [http://www.massbank.jp/]. Next, MS/MS experiments were performed to obtain structural information via the interpretation of the fragmentation pattern of the metabolite (Figure S3B). In the case of acylcarnitines, $m/z$ 85.0 is the characteristic fragment ion in the MS/MS spectrum [27]. The MS/MS spectra of possible metabolite candidates in the databases were also searched and compared. As a result, the metabolite was identified as hexanoylcarnitine, which was finally confirmed by comparison with the standard (Figure S3C). Both Metabo Analyst [28] and Ingenuity Pathway Analysis software (IPA, www.ingenuity.com) were used to identify relevant pathways.

Results

Global metabolome changes with dengue fever

In order to obtain a high level snapshot of metabolic flux at the different stages of dengue infection represented in our clinical samples, we examined variances in our data both by principal component analysis (PCA) as well as OPLS-DA. The PCA score plot (Figure 1A) indicates that the metabolome changes in DF patients are reversible. The most predominant metabolome changes in DF patients, as compared with healthy controls, happened at early febrile stage (Visit 1), followed by the defervesence stage (Visit 2) and finally returned to the control levels at the convalescence stage (Visit 3). This indicates that the most intense responses of the DF patients to DENV infection happen within the first 72 hours upon infection, and homeostatic recovery appears to begin around the fourth day from the onset of fever. This, more or less, coincides with the course of viraemia, which has been observed to peak during early febrile phase and are generally undetectable during the defervesence stage. The duration of DF illness in the EDEN study is reported to be 7–14 days [18], and a majority of patients recover within three to four weeks as seen at Visit 3, which could explain the metabolome similarities between patients at Visit 3 and healthy controls in the PCA score plot. In addition, PCA did not reveal distinct metabolome profiles between DENV serotype 1 and DENV serotype 3. This suggests converging host responses between the two serotypes which resulted in DF. While host molecular changes may be present between infections with different DENV serotypes, a larger patient set will be required to tease out the significant subtle differences.

OPLS-DA, which is a more rigorous classification tool, revealed clearer segregation of the healthy controls and DF patients during the three visits (Figure 1B). The quality of the OPLS-DA model was evaluated by $R^2$ (Y) and $Q^2$ parameters, which indicated the fitness and prediction ability, respectively. By using four principle components, the OPLS-DA model showed a $R^2$ value of 0.95 and $Q^2$ value of 0.91, indicative of optimal performance of the model. The OPLS-DA was also applied to GC-MS data, which showed a $R^2$ value of 0.65 and $Q^2$ value of 0.60.

Identification of significantly altered metabolites and pathways

Approximately 200 differential metabolites identified via both LC-MS/MS and GC-MS analyses were selected based on the
Figure 1. PCA (A) and OPLS-DA (B) score plots of dengue fever based on metabolomics data. The principle component analysis (PCA) and orthogonal partial least-squares discriminant analysis (OPLS-DA) models were constructed using LC-MS/MS metabolomics data from healthy subjects and dengue patients at three Visits. doi:10.1371/journal.pntd.0002373.g001

criteria- VIP>1 in OPLS-DA analysis and p<0.05 in Kruskal-Wallis test (Table S4, S5). Sixty of these metabolites were structurally identified, with 36 through LC-MS/MS analysis, and 9 by GC-MS analysis, 4 of which were common to both these methods. These metabolites belonged to classes such as free fatty acid, acylcarnitine, phosphatidylcholine (PC), lysophosphatidylcholine (LPC), LysoPC, amino acid and derivative, sphingolipid, monoacylglycerol (MG), diacylglycerol (DG), purine, bile acid, bile pigment, steroid hormone, glucose and phenol sulfate (Table S4). Consistent with the PCA score plot (Figure 1A), the quantitative heatmap (Figure 2) of these differential metabolites revealed that the metabolome changes were reversible in DF patients. They were significantly perturbed during the early stages, and normalized to control levels at convalescent stage.

These differential metabolites show two contrasting time-course changes, an elevated and a decreased trend, at Visit 1 and Visit 2. Most of the metabolites have either the highest or the lowest levels at Visit 1, which is the acute stage of DENV infection, and gradually return to the control levels at Visit 3, indicating that they might serve as prognostic markers of the disease.

It is interesting that the metabolites within each metabolic class generally showed a similar change trend. Specifically, seven FFAs, including four polyunsaturated fatty acids (PUFAs): arachidonic acid (AA; 20:4n-6; Figure 3A), linoleic acid (18:2n-6), docosahexaenoic acid (DHA; 22:6n-3) and α-linolenic acid (18:3n-3), showed an elevated trend at Visit 1 and Visit 2. The PUFA s are released from membrane phospholipids by various phospholipase enzymes, predominantly phospholipase A2, which is elevated in dengue patients [29]. Similarly, ten acylcarnitines, two sphingolipids (glucosylceramide and SM), two MG, one DG, one steroid hormone (cortisol), two purine nucleosides (hypoxanthine and inosine), one bile pigment (biliverdin), two conjugated bile acids (glycoursodeoxycholic acid and glycocholic acid), and glucose showed an increasing trend at Visit 1 and Visit 2 (Figure 3B–3F). Conversely, a total of twenty PCs, LPCs and lysoPEs showed a decreased trend (Figure 3G). Ten amino acids and derivatives, including tryptophan (Figure 3H), glutamic acid etc, also displayed a downward trend, with the exception of phenylalanine (Figure 3I), pyroglutamic acid, L-leucyl-L-proline, which showed an upward trend. The changes of amino acids observed are consistent with a previous report that described an overall decrease in plasma amino acid levels in dengue patients, with the exception of increased phenylalanine in the acute phase of DF patients [30].

We used two well-known pathway analysis tools- MetaboAnalyzer and IPA, to determine if the metabolites identified here revealed underlying biochemical pathways. Both these tools provided similar results, revealing that disturbed metabolic pathways included fatty acid biosynthesis, fatty acid β-oxidation, phospholipid catabolism, arachidonic acid metabolism, sphingolipid metabolism, tryptophan metabolism, steroid hormone biosynthesis, purine metabolism, heme degradation pathway, bile acid biosynthesis, etc (Figures S4). The Networks function in IPA indicated that the most significant changes happened with lipid metabolism and energy production pathways, which is summarized in Figure 4.

Global serum lipid changes with dengue fever

OPLS-DA revealed distinct temporal serum lipidome profiles with DENV infection (Figure 3A). To understand the lipids and blood chemistry parameters responsible for the differentiation of the three visits in the OPLS score plots, the VIP plot was generated (Figure 3B). Lipidome changes, SMs, PCs and lysoPEs influenced the model most significantly. We noted strong negative correlation between total SM with lymphocyte percentage (Pearson r² = 0.631, 95% C.I. = -0.6476 to -0.647, p<0.0001; Figures 6A, S3A). The downward trend of SM levels across the visits (repeated measures 1-way ANOVA, p<0.0001) was dominated by SM d18:1/16:0 (m/z 703.7→184.1) which represented ~1/3 of total SM and exhibited a 5-fold decrease from Visit 1 to 2 (Figure 6B). In cytotoxic T lymphocytes, acid sphingomyelinase, the enzyme that hydrolyzes SM to Cer, is a critical factor in the secretory granule-mediated cell death of virus infected cells [31]. Concordant with SM hydrolysis to ceramides [32], we observed increased ceramide levels parcellating the attenuation of SM (Figure S5B). Unlike the more biologically inert SMs, Cers are potent apoptosis-inducers, and their glycosylated forms, MHexCers or phosphorylated forms, C1Ps are bioactive sphingolipids [33]. The initial low Cer levels were not due to its glycosylation or phosphorylation which remained low (Figures S5C, S5D), but plausibly compartmentized as the less bioactive SM during the early onset of DF. This is evidenced by the relatively high SMs levels in the febrile phase compared to the later defervescence and convalescence phases (Figure 6C). The development of DENV-specific CD4+ and CD8+ T-lymphocyte responses [34], and the activation and contraction of antiviral CD4+ T lymphocytes in the defervescence and convalescence phases of dengue fever has previously been documented [35]. Therefore, the decrease in SM and increase in Cer correlated well with the action of T lymphocyte sphingomyelinase on SM and Cer levels in generating cytotoxic granules [31].

In our study cohort, MHexCers significantly increased from Visit 1 to Visit 3 (repeated measures 1-way ANOVA, p<0.01; Figure 5C), verifying our untargeted metabolomics results. Conversely, the higher glycosphingolipid, LacCer did not show significant differences across the visits (Figure S5E).

In our study we found many lysoPEs and pPEs closely correlated to platelet numbers (Pearson r² = 0.263–0.4, p<0.0001; Figures 6C, S6A). By contrast, PE s did not correlate with platelet numbers (Pearson r² = 0.404, 95% C.I. = -0.4760 to 0.1096, p = 0.217). In addition, their temporal trends differed from pPEs and lysoPEs (Figures S5B–D). These pPEs are polyunsaturated and long-chained lipids (34p:2, 36p:4, 38p:4; 38p:5, 38p:6 and 40p:6). Phosphatic acid glycerophospholipids are characterized by vinyl ether bonds at sn-1 position of the glycerol backbone, and two predominant glycerophospholipids are PEs and PCs. We also found that LPCs and lysoPEs strongly correlated to white blood cell count (Pearson r² = 0.425, p<0.0001). pPEs with no or single double bonds and of chain lengths more than 40 carbons correlated poorly, suggesting apparent alkyl chain length specificity to white blood cell count. Furthermore, univariate 1-way ANOVA showed that lysoPEs are significantly increased between Visits 2 and 3 (p<0.0001), and to a lesser degree between Visits 1 and 2 (p<0.05; Figure S3B). Taken together, our combined lipidome and blood chemistry analyses revealed that SMs are associated with early host response whereas lysoPEs are more associated with late recovery responses.
Figure 2. Heat map of identified differential metabolites from both LC-MS/MS and GC-MS analyses. Each row shows ion intensity for a specific metabolite after mean centering and unit variance scaling of the data. Each column shows the serum metabolic profiles of healthy subjects and dengue patients at three visits. * differential metabolites from GC-MS analysis.
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**Discussion**

DF is a self-limiting febrile illness and in most DF cases, there is complete recovery with little to no manifestations of chronic negative clinical consequences, clearly indicating human-adapted protective mechanisms against DENV in DF patients. The protective mechanisms are complex and unfortunately, poorly studied, in part due to the focus on heterotypic infection and DHF/DSS. The study of the recovery of DF from a physiologic perspective is therefore important in advancing our understanding how humans mount a defense and

### Figure 3. Typical change trends based on differential metabolite classes.

- **A.** Elevated change trend of arachidonic acid (free fatty acid).
- **B.** Elevated change trend of hexanoylcarnitine (acylcarnitine).
- **C.** Elevated change trend of glucosylceramide (sphingolipid).
- **D.** Elevated change trend of cortisol (steroid hormone).
- **E.** Elevated change trend of inosine (purine).
- **F.** Elevated change trend of biliverdin (bile pigment).
- **G.** Decreased change trend of PC 38:8 (phospholipid).
- **H.** Decreased change trend of tryptophan (amino acid).
- **I.** Elevated change trend of phenylalanine.

Median-horizontal line in the box. The bottom and the top of the box are the 25th and the 75th percentiles, and the black band near the middle of the box is the median peak area of the metabolite. *p* < 0.05, **p** < 0.01, ***p*** < 0.001, ns, non-significant by test Kruskal-Wallis test. The statistical comparison was with control levels.

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**Discussion**

DF is a self-limiting febrile illness and in most DF cases, there is complete recovery with little to no manifestations of chronic negative clinical consequences, clearly indicating human-adapted protective mechanisms against DENV in DF patients. The protective mechanisms are complex and unfortunately, poorly studied, in part due to the focus on heterotypic infection and DHF/DSS. The study of the recovery of DF from a physiologic perspective is therefore important in advancing our understanding how humans mount a defense and
recover from DENV infection and potentially other flaviviruses-induced illnesses. Our current integrated metabolomics and lipidomics study provides an extensive map of physiologic changes in adult DF patients, through which we have identified altered metabolic pathways linked to DENV infection. The relatively large population size (44 patients and 50 healthy controls) with repeated sampling at multiple time-points of infection, including early febrile, defervescence and convalescence stages, captured dynamic changes in metabolites and the discovery of reliable differential metabolites that were closely associated with dengue pathophysiology. By using both LC-MS/MS and GC-MS, we found that approximately 200 serum metabolites were significantly changed, of which 60 were identified. Lipids, which form a biochemically important subclass of metabolites, were also independently verified and studied. These metabolites constituted a mix of pro- as well as anti-inflammatory mediators, suggesting the rapid host resolution to limit excessive inflammation and curtail further exacerbation of DENV infection. Furthermore, these metabolites showed reversible trend changes, indicating that they might be able to potentially serve as the basis to develop prognostic markers of the disease.

Infection of humans with DENV causes early pronounced immunological reactions that shift towards pro-inflammatory responses [36]. While acute inflammatory response is necessary in initiating pathogen killing, activation of anti-inflammatory and pro-resolution processes is important to prevent excessive pathological inflammatory damage to the host. Omega-6 PUFAs, which

![Diagram of altered lipid and energy metabolic pathways in dengue patients.](image)

**Figure 4. Major altered lipid and energy metabolic pathways in dengue patients.** The four points on each line represent the levels of each metabolite at control, Visit 1, Visit 2 and Visit 3, respectively. Differential metabolites colored in red or green represent elevated or decreased change trend at acute stages (within 1 week) after dengue virus infection compared with healthy control subjects. These changes are reversible and there is no significant difference for most of the differential metabolites between healthy controls and dengue patients at convalescence stage (week 3–4). Black-colored metabolites either indicate similar level in healthy controls and dengue fever patients or are undetectable in this study. PC, phosphatidylcholine; PUFA, polyunsaturated fatty acid; LPC, lysophosphatidylcholine; MG, monoacylglyceride; DG, diacylglyceride; TG, triacylglyceride; SM, sphingomyelin.

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are released from cell-membrane phospholipids by phospholipid hydrolysis, including AA and AA-derived mediators such as prostaglandins, thromboxanes, hydroxyeicosatetraenoic acids and leukotrienes (collectively known as eicosanoids) are potent mediators of inflammation and are involved in modulating both the intensity and duration of inflammatory responses [37,38]. Contrary to the effects of AA, omega-3 PUFAs, e.g. DHA are known for their anti-inflammatory activity [39]. Our results showed an elevated trend of pro- and anti-inflammatory PUFAs during the febrile stage of DENV infection, including both AA and DHA, as are total hydroxyeicosatetraenoic acids [40]. Such effects of active resolution, where AA-associated inflammation incited early during DENV infection is concomitantly resolved by DHA-derived anti-inflammatory mediators, and the suppressed production of pro-inflammatory cytokines [39], is believed to embrace the current model of inflammation whereby the return to homeostasis is an active process, as opposed to a passive one [41]. We also found increased levels of cortisol, a major, cytokine-activated glucocorticoid of potent anti-inflammatory and immunosuppressive properties in the febrile stage of DF. Viral infections are capable of inducing an increase of cortisol in serum during DENV infection, it had been reported that cortisol level at early febrile stage was significantly higher than its level at convalescent stage [42], consistent with our results. The functional role of cortisol in restraining inflammatory and immune responses by inhibiting the production of cytokines and other pro-inflammatory mediators [43] could possibly down-regulate immune responses and protect the DF patients against cytokine-mediated pathologies. Consistent

Figure 5. Serum lipids and blood chemistry describes distinct time points in primary dengue infection. A. Orthogonal partial least-squares discriminant analysis (OPLS-DA) models of median-centred and standard deviation scaled glycerophospholipids and sphingolipids and blood chemistry parameters collected from Visit 1 (1 to 3 days after the onset of fever; febrile phase), Visit 2 (4 to 7 days post fever onset; defervescence phase) and Visit 3 (21 to 27 days post fever onset; convalescence phase). B. OPLS-DA VIP plot showing the association of serum glycerophospholipids and sphingolipids with the blood chemistry parameters. Only parameters of VIP values >1.5 are shown. LYMPH x10^3/μL, lymphocyte numbers; LYMPH %, percentage of lymphocytes. doi:10.1371/journal.pntd.0002373.g005
with elevated oxidative stress in adult DF patients [44] and elevated inosine concentrations in the extracellular space at times of inflammation [45], we found an increased serum inosine in the febrile phase. Inosine, potentially formed through nitrosative deamination of adenosine, has been shown to exert wide-ranging anti-inflammatory effects both in vivo and in vitro, such as inhibition of pro-inflammatory cytokine/chemokine production and the enhancement of anti-inflammatory cytokine IL-10 production [46]. Taken together, our metabolomics results reveal a dynamic metabolite flux in dengue where an acute inflammation is limited by swift, dampening responses to restore homeostasis.

In parallel, host immunity has interested dengue investigators in exploring the mechanisms in the immune system underlying DF pathogenesis. In primary infections, neutralizing antibody responses develop after DENV infection and provide persistent protection against the same serotype even in repeated infections; even though this may lead to the enhancement of disease severity upon challenges of a following heterotypic DENV infection. On the basis of the aforementioned observations, we postulated that the relationship between SM, Cer concentrations and immune cells describe multiple early, molecular roles of sphingolipids in mediating the host interactions during DF. Central among the sphingolipid enzymes is acid sphingomyelinase, which has been implicated in numerous immune responses including, T lymphocyte activation and positive infection by ebola viruses [53], entry of measles viruses into dendritic cells [54] and exocytosis of cytolytic granules by T cells [31]. The binding of extracellular ceramide to the leukocyte mono-immunoglobulin-like receptor 3 has been recently reported in repressing mast cell activation [55] and serum ceramides could potentially influence mast cell surveillance in DENV infection [56]. Likewise, we also observed the correlation of LPEs and pPEs with white blood cells and may potentially be related to iNKT activation. Plasmalogen lysoPEs (pLPE) are self-antigenic and are required for iNKT stimulation under both physiological conditions and when challenged with bacteria [48]. We expect that, our findings that sphingolipids are associated with lymphocytes during DF may further unravel the significance of T-cell responses and warrants further investigations. The lack of any significant change in phosphatidic acids and its lyso form was intriguing given its

Figure 6. Correlation of serum lipids with specific serological parameters. A. Linear correlation of lymphocyte numbers with total sphingomyelin (SM). B. Quantitative mass spectrometry analysis of individual SM species. C. Scatter plot showing the temporal trend of total SM during dengue fever. *** P<0.0001. D. Linear correlation of platelet numbers with total lysoPE. doi:10.1371/journal.pntd.0002373.g006
commonly assigned roles in platelet activation. However, it could be because unlike DHF, DF does not entail severe thrombocytopenia and our blood chemistry assays suggest the same.

In the wide spectrum of responses occurring simultaneously with pathogenic invasions, some metabolic changes may appear to be general and others more specific. In influenza 2009 pandemic H1N1 infection, extensive changes in serum eicosanoid (e.g. prostaglandin-F12, prostaglandin-G, LTA4) and linolenic acid levels were observed [57]. While linolenic acid is a FFA common to both H1N1 infection and dengue infection, there was no observable difference in eicosanoids in DF. Another plausible distinctive metabolome change in DF compared to H1N1 is the xanthine derivatives. In 2009 pandemic H1N1 infection, serum methyloxanthine levels were lower during infection relative to after treatment [57]. This result can be interpreted in two ways: theophylline, a methyloxanthine has been shown to elicit immunomodulator, anti-inflammatory, and bronchoprotective properties [58] and such properties may help attenuate the bronchi damaging effects of 2009 pandemic H1N1 virus as part of the host response [59] or constitute part of the unspecified treatment. Regardless of which is the reason, methyloxanthine points towards the lung-injury specific consequence of 2009 pandemic H1N1 infection. On the other hand, in DF, circulating inosine and hypoxanthine levels increased in the febrile phase where systemic inflammation may be concerned [60,61]. While studies of increased inosine and hypoxanthine levels were demonstrated in models of chronic systemic inflammation, it is conceivable that similar events could be observed in infectious diseases where acute bouts of inflammation occur. During inflammation, increased adenosine-to-inosine editing in cytokine mRNA may interfere with the degradation of cytokine transcripts and support the early pro-inflammatory and resolving responses necessary in the early phase of dengue. This may be reflected in a mixture of pro- and anti-inflammatory cytokines such as significant IL-4, IL-8 and IFN increases in the febrile phase of DF [16].

On the other hand, certain metabolic changes may be more general across different pathogenic infections. For example, we found increased phenylalanine levels in the early stages of DF. Tetrahydrobiopterin (BH4) is a co-factor for phenylalanine (4)-hydroxylase (PHA), an enzyme required for converting phenylalanine to tyrosine, and also NO synthase (NOS) [62]. Through immune and inflammatory responses-mediated up-regulation of NOS, NO is elevated during dengue infection [63,64]. Therefore, NOS competes with PHA for BH4 and this causes an accumulation of unconverted phenylalanine in the blood. This is aggravated in the elevated oxidative stress status in DF patients [44] where superoxide and peroxynitrite increases BH4 oxidation, thereby depleting the available BH4 pool. Yet, an increased level of phenylalanine has been commonly observed in other infectious diseases [65] and given NO’s diverse functions in immunity and inflammation responses it is not surprising that such changes could occur in both bacterial and viral infections [66,67]. Another general effect is energy metabolism. In this study we identified ten acylcarnitines, which are essential intermediates of fatty acid β-oxidation (FAO). FAO in liver is suppressed during infection and inflammation, and increased FFA substrates are then moved away from oxidation and directed toward re-esterification into TG [68]. The elevated levels of acylcarnitines suggest disturbed FAO in DF patients, which is in agreement with the observation that the TCA cycle was affected in an endothelial cell line infected with DENV [14]. Proteomics and lipidomics work by Diamond et al., showed a shift in energy metabolism during HCV infection in Huh7.5 cells, plausibly as a compensatory measure for the energy-demanding biosynthesis [15]. Similarly, a metabolomics study on severe pneumonia identified energy metabolism as a perturbed pathway [10]. Our relatively large number of human samples not only corroborated the in vitro findings but also showed that energy metabolism is likely to be a general perturbed pathway in acute infections. In the absence of large datasets of host-pathogen ‘functional omic’ datasets, we are limited in our ability to make comparisons and evaluate pathways specific to either bacterial or viral infections in humans. However, with increasing need to comprehensively capture the complexity of host-pathogen interactions, and aided by advancements in mass spectrometry and computational biology [69], we expect more datasets made available to the community for meaningful comparisons.

The limitations of our study can be discussed in the context of the host. Firstly, since the current study is restricted to adult primary infections the findings still need to be expanded and evaluated in pediatric patients as well as patients with secondary infections. Secondly, DENV progressively infects a multitude of cells and organs, including dendritic cells, monocytes, macrophages, endothelial cells and liver [70–72]. When we approached the host response from a systemic perspective, changes of the chemical composition in the serum could reflect tissue lesions, organ dysfunctions, pathological states and also compensatory responses. Thus pinpointing the source or cause of these changes can be challenging. Nevertheless, our results strongly raise the implication of distinctive metabolome changes with specific physiological responses at different phases of dengue infection. It will require increasing volumes of investigative work to tease apart the sources of these metabolites, including lipids. More importantly, as the body of metabolic work on in vivo viral infections grows, the potential to derive distinctive and accurate biomarkers alluding to host responses of various viral infections increases.

In summary, our findings provide a first detailed description of the metabolome changes in patients with acute dengue infection and offer a global molecular view that leads to the overall homeostatic physiological outcome of DF. In addition to providing host-pathogen insights into the underlying mechanism in symptom manifestation, metabolites identified in this study might be used both to monitor disease progression as well as evaluate the efficacy of therapeutic interventions. Furthermore, certain pathways might serve as useful therapeutic targets to alleviate severity of dengue.

Such work would be our next endeavour.

Supporting Information

Figure S1 Workflow of serum metabolomics study based on LC-MS/MS and GC-MS analysis. (TIF)

Figure S2 Schematic workflow of mass spectrometry-based lipidomics on serum samples collected from EDEN study. (TIF)

Figure S3 Identification of hexanoylcarnitine as a differential metabolite in dengue fever. A. the extracted ion chromatogram (EIC) and matched formula of the ion m/z 260.18. B. MS/MS spectrum and proposed fragmentation pathways of the ion m/z 260.18. C. MS/MS spectrum of a commercial standard hexanoylcarnitine. (TIF)

Figure S4 Pathway analysis results by MetaboAnalyst. (TIF)

Figure S5 Temporal trends of A. lymphocyte numbers and B. Cer, C. GlcCer, D. C1P and E. LacCer levels. *p<0.05; **p<0.01. (TIF)
were determined according to the International Conference on Harmonization (ICH) guidelines.

Table S4 Identified differential metabolites from LC-MS/MS analysis.

(DOCX)

Table S5 Unidentified differential metabolites from LC-MS/MS analysis.

(DOCX)

Author Contributions

Conceived and designed the experiments: LC YHL YK EEO SRT CNO. Performed the experiments: LC YHL. Analyzed the data: LC YHL YK FX KL. Contributed reagents/materials/analysis tools: LC YHL YK FX KL EEO SRT CNO. Wrote the paper: LC YHL YK EEO SRT CNO.

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