Innate immune cytokine profiling and biomarker identification for outcome in dengue patients

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Abstract: Biomarkers of progression to severe dengue are urgently required for effective patient management. Innate immune cells have been implicated in the enhancement of infection and "cytokine storm" associated with dengue severity. Using intracellular cytokine staining and flow cytometry, we observed significantly higher proportions of innate immune cells secreting inflammatory cytokines dominated by IFN- γ and TNF- α at admission associated with good prognosis. Secondary dengue predisposed to severe outcomes. In patients with severe dengue and those with liver impairment, early activation as well as efficient down-regulation of innate

responses were compromised. IFN-γ⁺CD56⁺CD3⁺ NKT cells and IL-6⁺ granulocytes served as novel biomarkers of progression to severity (composite AUC=0.85-0.9). Strong correlations among multiple cytokine-secreting innate cell subsets pointed to coordinated activation of the entire innate immune system by DENV.

Short Title: Innate cytokine biomarkers of dengue severity

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One Sentence Summary: Activation and efficient attenuation of innate immunity are both compromised in severe dengue.

Main Text: Dengue virus (DENV) afflicts ~130 million people annually with 70% contribution from Asia (1). Majority of patients present with mild dengue fever (DF) while ~5-20% develop severe dengue (SD)- characterized by plasma leakage leading to shock and/or organ impairment that predominantly manifest during defervescence (2, 3). The need to identify severe cases early, to facilitate interventions that reduce mortality has spawned numerous investigations to find biomarkers of severity. The phenomenon of antibody dependent enhancement (ADE) of viral infectivity has been implicated in dengue pathology (4, 5) with monocytes that both support virus growth and express all three classes of Fc-receptors (FcγR, FcεR and FcαR) being major contributors (6, 7). Deficits in T cell signaling and cytokine secretion have been reported in dengue patients from Sri Lanka, India and Thailand (8, 9). Combined with spontaneous high production of multiple cytokines from unstimulated PBMCs (9), this implicated innate cells as likely sources of the 'cytokine storm' in SD (10), supported by in vitro assays of infected monocytes and natural killer (NK) cells (11, 12). Conflicting patterns for serum levels of TNF-α, IP-10 and IFN-γ as a function of severity were reported (13-16), perhaps due to variation in sampling time following infection. The failure to exploit flow cytometry made it impossible to identify the source of cytokines measured by ELISA. Against this backdrop, we carried out a blinded study (fig. S1) to

query the secretion of cytokines by human innate immune cells in response to DENV using intracellular cytokine staining and flow cytometry (fig. S2 & S3).

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Dengue infection, in contrast to fevers from other etiologies, caused a surge in secretion of inflammatory cytokines from a variety of innate cells (Fig. 1, A-F). TNF-α was by far the most abundant cytokine secreted by multiple cell subsets from a vast majority of patients while an impressive 95.5% of patients secreted IFN-γ from CD56+CD3+ NKT cells (table S1). We also observed significantly greater numbers of dual-functional IL- 10^{+} TNF- α^{+} granulocytes and IFN- γ^{+} TNF- α^{+} CD56⁺CD3⁺ NKT cells, in dengue patients than in febrile controls (fig. S4A, B). Polyfunctional CD56⁺CD3⁺ NKT cells among others, could indeed be visualized by t-Distributed Stochastic Neighbor Embedding analysis (fig. S5). Thus, in addition to monocytes which support dengue replication (11), all other innate cell subsets were also activated by DENV. As reported previously (17), DENV infection significantly increased the numbers of CD14⁺ and CD14⁺CD16⁺ monocytes and significantly reduced CD56+CD3+ NKT cell numbers compared to controls (table S2; fig. S6A, B). Some cell subsets progressively expanded with increase in disease severity (table S3). CD14⁺CD16⁺ intermediate monocytes or CD19⁺ B cells were the highest per cell secretors of all cytokines (fig. S7, A-E; table S1). The median florescence intensity (MFI) was comparable across severity except for TNF-α⁺CD56⁺CD3⁺ NKT cells (fig. S7F, G).

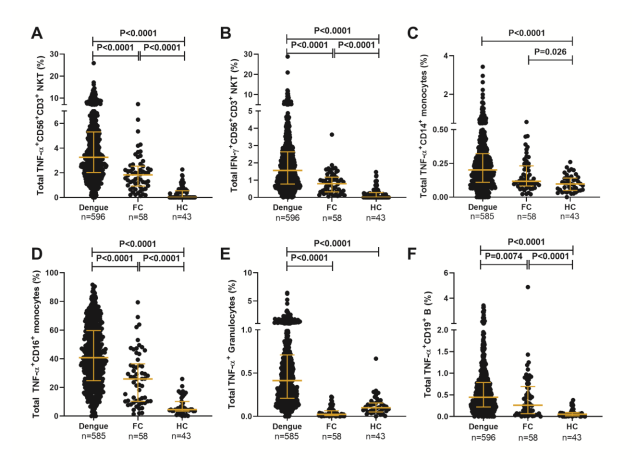


Fig. 1. DENV activates cytokine secretion from innate immune cells. Frequency of total (**A**) TNF- α ⁺CD56⁺CD3⁺- and (**B**) IFN- γ ⁺CD56⁺CD3⁺-NKT cells, (**C**) TNF- α ⁺CD14⁺ monocytes, (**D**) TNF- α ⁺CD16⁺ monocytes, (**E**) TNF- α ⁺ granulocytes and (**F**) TNF- α ⁺CD19⁺ B cells from dengue patients compared to febrile controls (FC) and healthy controls (HC). Each dot represents one patient; P value displayed with median and IQR.

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When assessed as a function of disease severity, a significantly greater proportion of TNF- α^{+} granulocytes were evident in DF relative to DFWS/SD (Fig. 2A), a trend also evident for dual secretion of TNF-α and IL-6 by this cell subset and IL-10 secretion by CD56⁺CD3⁺ NKT cells (Fig. 2B, C). When we used bleed-scores or liver enzyme levels as a surrogate of severity, those with no bleeding or normal liver enzyme levels carried a significantly greater proportion of innate cells secreting inflammatory cytokines compared to those with varying degrees of hemorrhage or abnormal liver enzyme levels (Fig. 2, D-F). As expected, the odds of severity were greater in patients with high liver enzymes (fig. S8A, B) and secondary dengue (fig. S8C, D). The latter predisposition was accompanied by a significantly lower proportion of TNF-α secreting monocytes, granulocytes and CD19⁺ B cells (fig. S8, E-H) reinforcing the reported (4, 5) deleterious role of pre-existing immunity on dengue outcomes. In addition, age and gender also influenced cytokine secretion from monocyte subsets (fig. S8I, J). Despite variations within the cohort based on gender, age and primary/secondary dengue, the higher percentages of innate cell subsets secreting inflammatory cytokines, impressively correlated with better prognosis as suggested by earlier blood transcriptome studies (18). In contrast, higher frequency of monofunctional IP-10⁺CD19⁺ B cells was associated with severity (fig. S9A, B). Total TNF- α^+ percentages within all subsets displayed strong positive correlation with one another (r=0.63-0.77), while IFN- γ^+ and IFN- γ^+ TNF- α^+ within CD56+CD16+ NK or CD56+CD3+ NKT cells positively correlated with each other (table S4), demonstrating synchronized activation of all innate cells by DENV. In our large cohort, non-structural (NS) protein 1 levels, a measure of viral load did not vary with disease duration (fig. S10A). Patients with highest NS1 levels displayed highest innate cell cytokine secretion (fig. S10, B-D) and also avoided severe outcomes (fig.

S10E), suggesting a requirement for high viral antigen levels to achieve efficient innate cell activation.

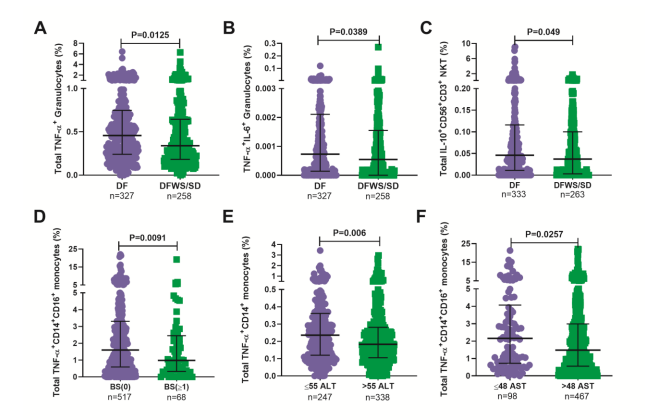


Fig. 2. Higher percentages of innate immune cells secreting inflammatory cytokines are associated with better prognosis. Frequency of (**A**) total TNF-α⁺ granulocytes, (**B**) TNF-α⁺IL-6⁺ granulocytes, and (**C**) total IL-10⁺CD56⁺CD3⁺ NKT cells compared between DF and DFWS/SD. (**D**) Frequency of total TNF-α⁺CD14⁺CD16⁺ monocytes compared between bleed-scores (BS). Frequency of total (**E**) TNF-α⁺CD14⁺ monocytes and (**F**) TNF-α⁺CD14⁺CD16⁺ monocytes compared between normal and elevated ALT (>55 IU/L; E) and AST (>48 IU/L; F). P values with median and IQR reported.

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In order to query the link if any, between kinetics of innate immune activation by DENV and disease severity, we compared innate cell cytokine secretion between different measures of severity at early (days 1-3), intermediate (days 4-6) and late (days 7-15) times of hospital presentation. Patients admitted 1-3 days post symptom onset (dpso) had impressively higher TNFα (Fig. 3A) and IFN-γ (Fig. 3B, S11A) -secreting innate cells in those with normal compared to above-normal liver enzyme levels. Those admitted 4-6 dpso had a significantly greater proportion of IFN-γ⁺CD56⁺CD3⁺ NKT cells as well as TNF-α-secreting granulocytes, CD19⁺ B cells and CD56⁺CD3⁺ NKT cells in DF relative to DFWS/SD (Fig. 3, C-F). In contrast, patients with high liver enzymes failed to down-regulate secretion of TNF-α, IL-6 and IP-10 from different innate cell subsets during the late stage (7-15 dpso) that was prominent in patients with normal levels (Fig. 3G, H & S11, B-D). Severe dengue was also characterized by a late surge of IP-10 (fig. S11, E-F). Dysregulated high levels of IFN- γ^+ CD56 $^+$ CD3 $^+$ NKT cells and TNF- α^+ CD19 $^+$ B cells were also evident in DFWS/SD relative to DF patients during this late time window (fig. S11, G-H). Thus, robust early secretion of TNF-α and IFN-γ by innate cells during acute disease phase combined with efficient attenuation of all innate cytokines during later stages of disease averted severity.

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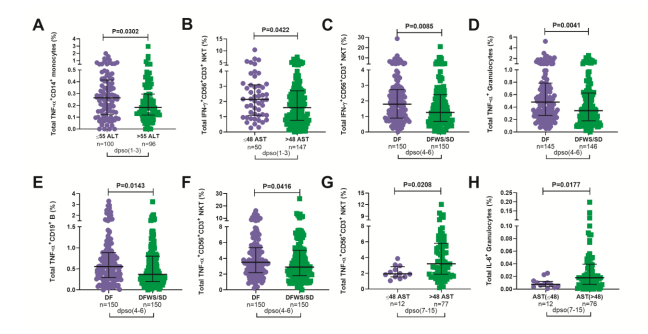


Fig. 3. Correlation between disease severity and kinetics of innate immune cell activation. Frequency of total (A) TNF- α ⁺CD14⁺ monocytes and (B) IFN- γ ⁺CD56⁺CD3⁺ NKT cells during 1-

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3 dpso compared between normal and high levels of ALT (A; IU/L) and AST (B; IU/L). Frequency of total (**C**) IFN- γ ⁺CD56⁺CD3⁺ NKT cells, (**D**) TNF- α ⁺ granulocytes, (**E**) TNF- α ⁺CD19⁺ B cells and (**F**) TNF- α ⁺CD56⁺CD3⁺ NKT subsets during 4-6 dpso compared between DF and DFWS/SD. Frequency of total (**G**) TNF- α ⁺CD56⁺CD3⁺ NKT cells and (**H**) IL-6⁺ granulocytes during 7-15 dpso compared between normal and high AST levels.

To identify potential biomarkers of progression to severity, we compared those who worsened after recruitment (as evidenced by a shift from DF/DFWS to DFWS/SD or death), with patients who readily recovered from DF and DFWS/SD. Both recovered DF and DFWS/SD patients carried a significantly greater proportion of IFN-γ⁺TNF-α⁺CD56⁺CD3⁺-, IFN- γ^+ CD56 $^+$ CD3 $^+$ - NKT cells and monofunctional IL-6 $^+$ granulocytes (Fig. 4, A-C) relative to worsened patients. To assess the biomarker performance of these cells, receiver operating characteristic (ROC) curve analysis was performed. IFN-γ⁺TNF-α⁺CD56⁺CD3⁺-, IFN- γ^+ CD56 $^+$ CD3 $^+$ NKT and monofunctional IL-6 $^+$ granulocytes provided AUC of 0.77, 0.76 and 0.75 respectively with 90% sensitivity and 60 to 66% specificity when DF was compared with the worsened group (fig. S12, A-C). Combining IL-6⁺ granulocytes and IFN- γ ⁺CD56⁺CD3⁺ NKT cells using binary logistic regression resulted in composite AUC of 0.85 (Fig. 4D; table S5) and revealed that every one percentage rise in IFN- γ ⁺CD56⁺CD3⁺ NKT resulted in 3.12 fold lower odds of worsening (95% CI=1.1-9.2, P=0.035). In patients with elevated AST, this composite biomarker predicted the progression to severity with higher accuracy (AUC=0.9) and displayed 100% sensitivity with 81.9% specificity (table S5).

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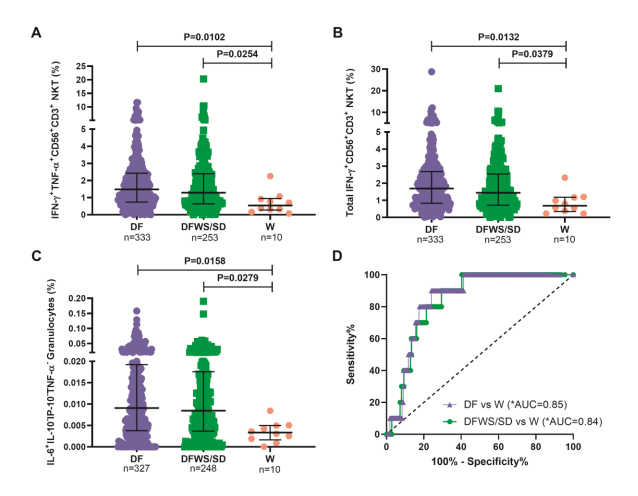


Fig. 4. Innate cytokine secreting cells predicted outcome in dengue patients. Frequency of (**A**) IFN- γ^+ TNF- α^+ CD56 $^+$ CD3 $^+$, (**B**) total IFN- γ^+ CD56 $^+$ CD3 $^+$ NKT cells and (**C**) IL-6 $^+$ IL-10 $^-$ IP-10 $^-$ TNF- α^- granulocytes compared between DF, DFWS/SD and worsened (W) patients. P value with median and IQR reported. (**D**) Composite ROC curve for total IFN- γ^+ CD56 $^+$ CD3 $^+$ NKT cells and IL-6 $^+$ IL-10 $^-$ IP-10 $^-$ TNF- α^- granulocytes comparing DF (purple) or DFWS/SD (green) with worsened patients. (*AUC), composite AUC.

This study, the first to query the cellular source of innate inflammatory cytokines in a large,

blinded dengue cohort, conclusively demonstrated the beneficial role of early innate cell activation

triggered by high viral antigen levels, in ensuring recovery from dengue. DENV activated all

innate immune subsets resulting in mono and polyfunctional cytokine production that correlated

with good outcome. The innate immune cytokine signature for each pathogen may be unique and

rewarding to investigate.

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The observed persistence of innate cell-derived cytokines during late phase of disease pointed to

dysregulated innate responses in SD (20). Severe COVID-19 patients also displayed sustained

plasma levels of IP-10 and IL-6 (21); increased plasma IP-10 also correlated with liver impairment

in HIV/HBV patients (22). In light of the reported requirement of IP-10 for B cell activation (23),

our finding of abnormal high IP-10 levels during late stages of disease is a likely contributor to the

B cell mediated pathology attributed to severe dengue (24). A potential role for persistently

elevated innate responses in provoking the reported inappropriate TCR signaling and T cell

apoptosis in SD (8, 9) warrants further investigation. Thus, failure to achieve both robust activation

and prompt attenuation of innate immune cells was a hallmark of severe dengue.

Our composite biomarker performed well despite limited number of patients with transitions to

greater severity and inclusion of patients with varying disease duration. Potential enhancement of

its performance by including additional hitherto unidentified cytokine secreting cells will enhance

its utility in a clinical setting. Ease of processing and ready availability of flow cytometers in

diagnostic laboratories assures feasibility of host blood-based biomarker deployment whereas

biomarkers reliant on expensive instruments and high-end technical skills, may have limited utility

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in resource constrained geographies (25, 26).

CD56⁺CD16⁺ NK cells secreting TNF-α showed strong positive correlation with multiple subsets pointing to its central role in the coordinated innate immune activation by DENV. Our findings are in line with the 20 gene transcript signature that included anti-viral IFN-γ signaling pathway genes, being under-expressed in NK and NKT cells of SD patients (26). Weaker activation of early innate effector mechanisms was also reported in patients who later developed severe manifestations (27). The reported impaired innate immune activation in severe SARS-CoV-2 patients (28), suggests that severe disease in multiple viral infections may be a shared consequence of defective modulation of kinetics of early host innate activation as well as its subsequent attenuation.

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supplementary materials. De-identified flow cytometry files (.fcs) that support the results reported in this article, and analyzed flow cytometry data (.xlsx) will be made available on request.

Supplementary Materials:

Materials and Methods
Supplementary Text
Figures S1-S13
Tables S1-S11
References (29, 30)

STARD Checklist

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Supplementary Materials for

Innate immune cytokine profiling and biomarker identification for outcome in dengue patients

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This PDF file includes:

Materials and Methods Supplementary Text Figs. S1 to S13 Tables S1 to S11

Other Supplementary Materials for this manuscript include the following:

STARD Checklist

Materials and Methods

Ethics statement

This study was carried out in accordance with the Declaration of Helsinki. Institutional ethics

committee approval to conduct the study was obtained from the four participating hospitals;

Bangalore Medical College and Research Institute (BMCRI; BMCRI/PS/25/2018-19),

Kempegowda Institute of Medical Sciences (KIMS; KIMS/IEC/A1-2018), St. John's Medical

College (SJMC; IEC/1/473/2019), M S Ramaiah Medical College (RMCH; MSRMC/EC/19) and

Indian Institute of Science (IISc; 10-14032018), Bengaluru, India.

Study subjects and clinical data collection

778 participants were enrolled for this blinded study between June 24 and November 29, 2019

which represents the annual dengue season following onset of monsoon rains at the four hospitals

listed above. Written informed consent was obtained from all participants before sample collection

and analysis. Blood samples were coded and labeled as follows: DEN/hospital abbreviation/19/###

before being sent to IISc for flow cytometry analysis. Consecutive suspected adult dengue patients

(≥ 18 years) who tested positive using a dengue specific NS1/IgM rapid dengue day 1 test kit (J

Mitra and Co., India) were recruited. Those who tested negative were recruited as febrile controls

(FC); volunteers with no illness for the past 3 months were enrolled as healthy controls (HC; fig.

S1).

Sample size analysis for a desired power of 80%, type I error tolerance of 0.05, and a hypothesized

effect size of 0.75, required at least 29 dengue patients who would transition post admission, to a

worse condition as defined by the World Health Organization (WHO) categorization of dengue

severity as follows: dengue fever (DF) was defined by headache, body ache, rash, nausea, or mild

bleeding; dengue fever with warning signs (DFWS) included symptoms like persistent vomiting,

mucosal bleeding, pleural effusion, ascites, and hepatomegaly. Severe dengue (SD) included symptoms such as plasma leakage, ≥1000 IU/L of alanine aminotransferase (ALT) / aspartate aminotransferase (AST), severe bleeding which leads to shock, and/or organ impairment (3). However, when the dengue cases ceased to appear in the hospitals by end November 2019, we had obtained only 10 patients who transitioned to a worse category, primarily owing to the clinical interventions following admission. The study was also designed to collect longitudinal samples at days 3 and 7 post-admission; however, only 154 patients provided a second sample and a single patient donated three consecutive samples. Demographic characteristics (i.e., gender and age), clinical features (i.e., days post symptom onset, nausea, head ache, body ache, abdominal pain, rashes, splenomegaly, hepatomegaly and bleeding manifestations) and routine hematological laboratory findings (i.e., complete blood cell count, serum albumin, liver enzymes, platelet count and hematocrit) were recorded. Patients were assigned bleed-scores (BS) as follows: no bleeding, 0; petechiae, 1; epistaxis/gingival bleeding/menorrhagia, 2; gastrointestinal bleeding, 3; intracranial/intrapulmonary bleeding, 4. Plasma leakage in pleural and/or peritoneal cavities was confirmed using X-ray/ultrasound scans. HIV patients were not recruited. Data from samples of patients with co-infections (typhoid, sepsis, malaria, urinary tract infection, Hepatitis B) or those who were discharged against medical advice (DAMA) and samples with experimental errors (clotted blood samples, QC failure of flow cytometer, sample processing errors) were excluded from analysis.

Serology

Dengue virus (DENV) infection was confirmed using a commercial IgM, IgG and NS1 enzymelinked immunosorbent assay (ELISA; Panbio, Australia) and results were interpreted according to manufacturer's instructions. The kits were used to distinguish primary (IgM to IgG ratio >1.2)

from secondary (IgM to IgG ratio <1.2) infection. Primary dengue status was also assigned to those

who tested positive for DENV specific NS1 (index value >1.1) but were negative for IgM and IgG.

Ex vivo intracellular cytokine staining of innate immune cells in whole blood

Blood samples collected in sodium citrate vacutainer tubes (BD Biosciences) were immediately

processed, no later than 4 hours from collection. RBCs from 500µl blood were lysed using 4ml of

1X ammonium chloride buffer (166mM ammonium chloride, 9.9mM potassium bicarbonate and

0.126mM EDTA). The centrifuged cells were washed with 1X phosphate buffered saline (PBS)

and stained with Fixable Viability Stain 450 [BD, Cat#562247] for 10 minutes at room

temperature, to exclude dead cells. This was followed by staining for appropriate surface markers

(table S6) for 30 minutes at 4°C. The surface marker TLR2 was superior to HLA-DR owing to its

stable expression during infection in contrast to the latter which is reported to be down-regulated

in all manner of inflammatory conditions, and was therefore used to identify monocytes (30). Cells

were fixed with 2% paraformaldehyde [Sigma Aldrich, Cat#P6148], washed and permeabilized

with 0.1% saponin. The permeabilized cells were stained with intracellular antibodies (monocyte

panel - IL-6, IP-10, IL-10, TNF-α; NK panel - IP-10, IL-10, TNF-α, IFN-γ) for 30 minutes at 4°C

(table S6). Cells were washed, resuspended in 1X PBS and data were acquired on a Beckman

Coulter DxFlex flow cytometer.

Analysis of flow cytometry data

Gating strategy for NK, monocyte and granulocyte subsets is shown in fig. S2. Cytokine-secreting

cells are represented as percentage of parent population. Control samples were stained with surface

antibodies and lacked all four intracellular markers keeping in mind the paucity of blood volumes

(fluorescence minus four; FMF). Fluorescence minus one (FMO) controls were compared with

FMF controls in five random individuals to confirm absence of non-specific binding of the

intracellular cytokine-specific antibodies. FMF controls were used to set the positive gates for each

cytokine (fig. S3). Positive cytokine production was based on the criteria given for each cytokine

from each cell subset in tables S7 and S8. Data were analyzed using FlowJo software (version

10.6.1). Polyfunctional cytokine secretion was assessed by Boolean gating. The analyzed data

from FlowJo was submitted to the clinical statistician for unblinding of patient characteristics prior

to statistical analysis.

Optimized t-Distributed Stochastic Neighbor Embedding (t-SNE; (31)) analysis to visualize the

clusters within the NK/NKT cells was performed. All 32 patient samples with SD were included

along with 32 each from DF and DFWS (WHO 2009 categorization) which were selected at

random using the RAND function in Excel 2016. A subset of 10,000 events were selected from

CD19⁻ cells for each sample using DownSample plugin, followed by concatenation of all events.

A total of 960,000 events and 7 markers (CD56, CD16, CD3, IP-10, IL-10, TNF- α , and IFN- γ)

were used to generate the t-SNE map. We used the KNN algorithm (random projection forest –

ANNOY) and Barnes-Hut gradient algorithm implementation with the recommended parameters

(iterations -1000; learning rate (eta) -67200) at perplexity =50 in t-SNE plugin built within

FlowJo.

Statistical analysis

All analyses were done using IBM SPSS statistics 23.0 and GraphPad prism version 8.

Significance between two or multiple groups was tested using Mann–Whitney U test (two-tailed)

and non-parametric Kruskal-Wallis test with a Bonferroni correction for multiple comparisons,

respectively. In patient cohort characteristics, normally distributed data were tested using one-way

ANOVA. Chi square test of independence and Fisher's exact test were used to evaluate the

association of clinical parameters with WHO categorization of patients based on severity.

Differences between proportions of primary and secondary infection across WHO categories were assessed using the Z test for proportions. Confidence intervals for odds ratio were determined using Baptista-Pike method. Spearman's correlation (two-tailed) analysis was performed to assess the positive or negative correlation between various cytokine secreting cell subsets. Receiver operating characteristic (ROC) curve analysis was performed to assess accuracy of proposed biomarker and 95% confidence intervals were calculated using Wilson/Brown method. Multivariate binary logistic regression was performed to compare DF or DFWS/SD with worsened groups as the dependent variables. Independent variables for multivariate analysis were selected if they were significantly different in univariate analysis (two-tailed Mann–Whitney U test for nonparametric continuous data and Chi square test for categorical variables). Required assumptions such as dichotomous mutually exclusive dependent variable, two or more independent variables, linear relationship between each independent variable and odds ratio, absence of multicollinearity were all met. Even though three independent variables were significantly different two of them directly correlated with each other (IFN-γ+TNF-α+CD56+CD3+ NKT cells and total IFN- γ^+ CD56 $^+$ CD3 $^+$ NKT cells). Hence we used a combination of monofunctional IL-6 $^+$ granulocytes with either total IFN-γ⁺CD56⁺CD3⁺ NKT cells or IFN-γ⁺TNF-α⁺CD56⁺CD3⁺ NKT cells to generate logistic regression models. The latter was not significant and was not used. Monofunctional IL-6⁺ granulocytes with total IFN-γ⁺CD56⁺CD3⁺ NKT cells regression model was a good fit confirmed by the Hosmer and Lemeshow goodness of fit test. The estimated probabilities obtained from logistic regression model were used to plot composite ROC curves.

Supplementary Text

Cohort Characteristics

Of the 596 subjects with laboratory confirmed dengue who were included in the final data analyses, the mean age was 30.4±10.79 (mean±SD, range 17- 69). 72% of enrolled patients were male and 28% were female (table S9). They were admitted to the hospital at a median of 4 days (range 1-15) post symptom onset. Dengue specific IgM and IgG ELISA distinguished 336 (56.4%) primary patients from 256 (42.9%) secondary dengue patients. 281 patients tested positive for dengue specific NS1 ELISA. In our cohort, 333 (55.8%) patients were classified as DF, 227 (38.7%) as DFWS and 32 (5.4%) as SD. The clinical laboratory parameters that correlated with the diagnoses are listed in tables S10 and S11. The longitudinal samples collected were not used for data analysis/interpretation since the innate immune activation status in these samples reflected the effect of medical intervention rather than disease progression. We therefore used the day of presentation at hospital to query the alterations in immune status as a function of kinetics of disease progression. Post recruitment and hospitalization, 7 patients worsened sufficiently to transition from DF to DFWS and 1 from DFWS to SD while 2 died. Interestingly, 9 among 10 worsened patients were men.

Cytokine signature of innate immune cells against DENV

The most abundant cytokines were IFN-γ from CD56⁺ NK cell subsets/CD19⁺ B cells and TNF-α from all innate cell subsets. CD16⁺ non-classical monocytes, CD56⁺CD3⁺ NKT cells, and CD14⁺CD16⁺ intermediate monocytes were the highest secretors of TNF-α. IFN-γ⁺TNF-α⁺ dual secreting cells were dominant in CD56⁺CD3⁺ NKT, CD19⁺ B cells, CD56⁺CD16⁺ and CD56⁺CD16⁻ NK cell subsets (fig. S12, A-D). CD56⁺CD3⁺ NKT cells also had abundant dual-functional IL-10⁺TNF-α⁺ cells followed by IP-10⁺TNF-α⁺ cells (fig. S12A). CD56⁺CD16⁺ NK cells carried polyfunctional profile dominated by IP-10 in combination with IFN-γ or TNF-α (fig.

S12B). CD19⁺ B cells also showed the presence of IL-10⁺TNF- α ⁺ dual functional cells followed by IP-10⁺TNF- α ⁺ and IP-10⁺IFN- γ ⁺. The triple positive cells from this subset included IFN- γ and TNF- α in combination with IL-10 or IP-10 (fig. S12C). Granulocytes and monocyte subsets had abundant IL-10⁺TNF- α ⁺ cells (fig. S12, E-H). Granulocytes were the predominant secretor of multiple cytokines (IL-6, IL-10, IP-10 and TNF- α ; table S1) in addition to multiple combinations of polyfunctional cells (fig S12H). IL-6 and IL-10 were the least abundant cytokines secreted by all queried innate cell types against DENV.

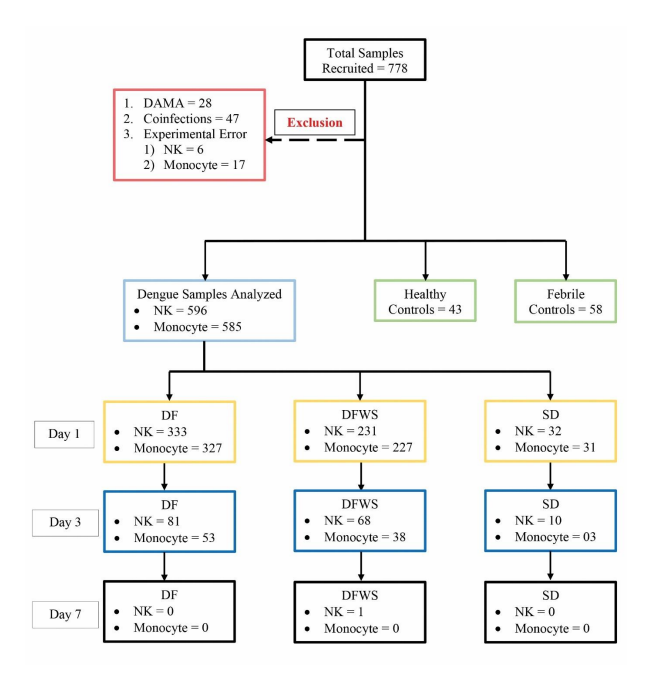


Fig. S1. Schematic of patient recruitment. A total of 778 patients were initially recruited; patients who were discharged against medical advice (DAMA), those with co-infections (i.e. typhoid, sepsis, malaria, urinary tract infection, hepatitis B infection) and samples with experimental errors, were excluded as shown. Abbreviations: DF – dengue fever; DFWS – dengue fever with warning signs; SD –severe dengue, NK – natural killer cell.

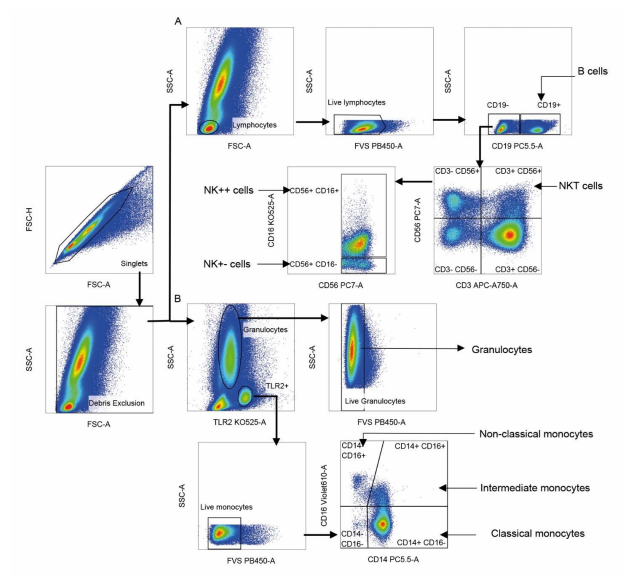


Fig. S2. Gating strategy for innate immune cell subsets. Singlet cells were selected based on FSC-A and FSC-H scatter and debris was excluded based on FSC-A and SSC-A. (A) Lymphocytes were gated using FSC-A and SSC-A scatter; dead cells and B cells staining for live/dead dye and CD19, respectively were sequentially excluded. Natural Killer T (NKT) cells were identified as CD56⁺ CD3⁺ (CD56 vs. CD3). Two CD3-negative natural killer (NK) cells subsets were identified as CD56⁺ CD16⁺ (NK++) cells and CD56⁺ CD16⁻ (NK+-) cells displayed on CD56 vs. CD16. Cells positive for CD19 lineage marker were identified as B cells. (B) Live monocytes were identified as TLR2⁺ and negative for live/dead dye. Three monocyte subsets were distinguished as

CD14⁺CD16⁻ classical monocytes (CM), CD14⁺CD16⁺ intermediate monocytes (IM) and CD14⁻ CD16⁺ non-classical monocytes (NCM) based on CD16 vs. CD14. Live granulocytes were distinguished based on SSC-A scatter vs TLR-2 followed by those negative for live/dead dye.

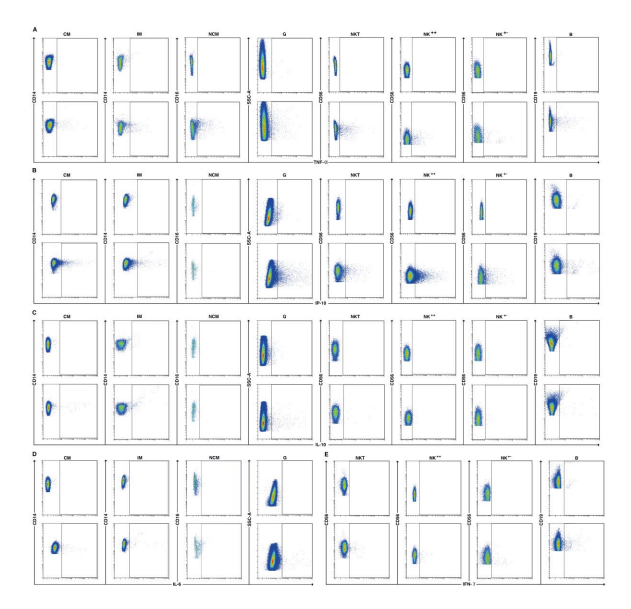


Fig. S3. Fluorescence minus four controls for flow cytometry. Pseudo color flow cytometry plots for a representative patient comparing the fluorescence minus four (FMF; top row) control with completely stained sample (bottom) for secretion of (A) TNF-α, (B) IP-10, (C) IL-10, (D) IL-6 and (E) IFN-γ from the indicated cell subsets. Abbreviations: CM – CD14⁺CD16⁻ classical monocytes; IM – CD14⁺CD16⁺ intermediate monocytes; NCM – CD14⁻CD16⁺ non-classical monocytes; G – Granulocytes; NKT – CD56⁺CD3⁺ cells; NK⁺⁺ - CD56⁺CD16⁺ cells; NK⁺⁻ - CD56⁺CD16⁻ cells; B – CD19⁺ cells.

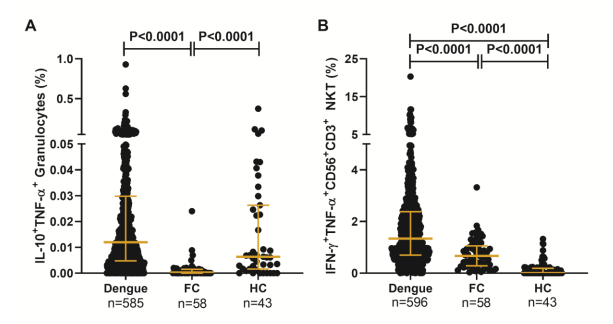


Fig. S4. DENV activates dual-functional cytokine secreting innate immune cells. Frequency of (**A**) IL-10⁺TNF- α ⁺ granulocytes and (**B**) IFN- γ ⁺TNF- α ⁺CD56⁺CD3⁺ NKT cells compared between dengue patients, febrile controls (FC) and healthy controls (HC). Each dot represents a patient sample. P value determined using Kruskal-Wallis test, followed by Bonferroni correction for multiple comparisons with median and IQR reported.

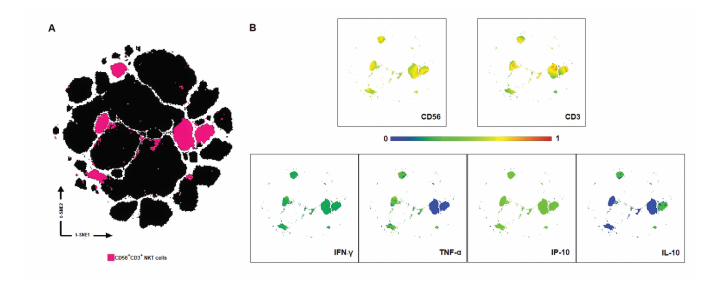


Fig. S5. Visualization of CD56⁺CD3⁺ NKT cells in t-SNE dimensional reduced space. (A) Two-dimensional representation identified five major CD56⁺CD3⁺ NKT cell clusters (pink) visualized by t-SNE map. (B) Differential expression of CD56, CD3, IFN-γ, TNF-α, IP-10 and IL-10 in NKT cell clusters visualized by two-dimensional multicolored t-SNE maps.

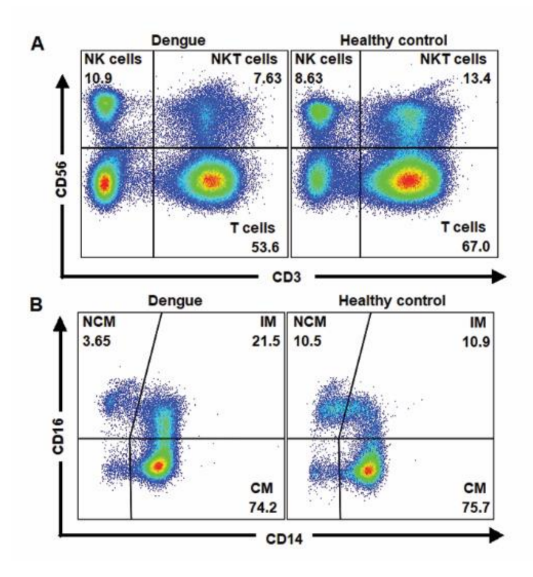


Fig. S6. Modulation of innate immune cell subsets by dengue. Pseudo color flow cytometry plots for a representative patient and healthy control show (**A**) reduction in percentages of CD56⁺CD3⁺ NKT cells and (**B**) expansion of percentage of CD14⁺CD16⁺ intermediate monocytes in dengue patients relative to healthy control. Abbreviations: NK - CD56⁺ cells; NKT – CD56⁺CD3⁺ cells; T – CD3⁺ T cells; NCM – CD14⁻CD16⁺ non-classical monocytes; IM – CD14⁺CD16⁺ intermediate monocytes; CM – CD14⁺CD16⁻ classical monocytes.

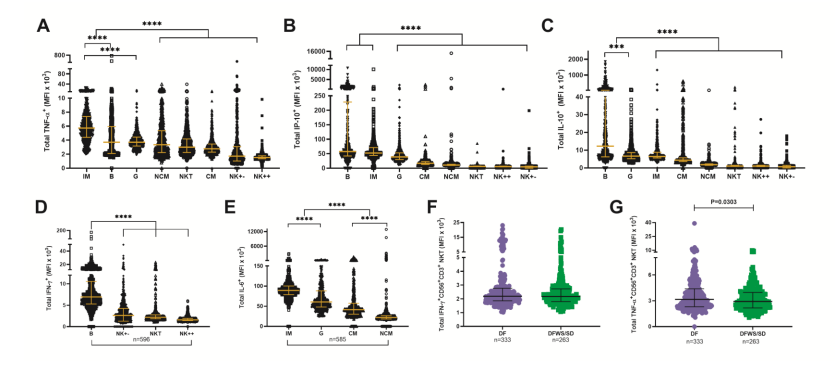


Fig. S7. Per cell secretion of cytokines from each cell subset. MFI values for each cytokine were compared between subsets. (A) TNF- α (B) IP-10 (C) IL-10 (D) IFN- γ (E) IL-6 compared among cell subsets. P values were determined using Kruskal-Wallis test, followed by Bonferroni correction for multiple comparison between groups with median and IQR reported. **** P<0.0001. MFI of (F) total IFN- γ ⁺ and (G) total TNF- α ⁺ cells within CD56⁺CD3⁺ NKT cells compared between DF and DFWS/SD. Mann-Whitney *U* test was performed and median with IQR are reported. Abbreviations: MFI – median florescence intensity; CM – CD14⁺CD16⁻ classical monocytes; IM – CD14⁺CD16⁺ intermediate monocytes; NCM – CD14⁻CD16⁺ non-classical monocytes; G – Granulocytes; NKT – CD56⁺CD3⁺ cells; NK⁺⁺ - CD56⁺CD16⁺ cells; NK⁺⁻ - CD56⁺CD16⁻ cells; B – CD19⁺ B cells.

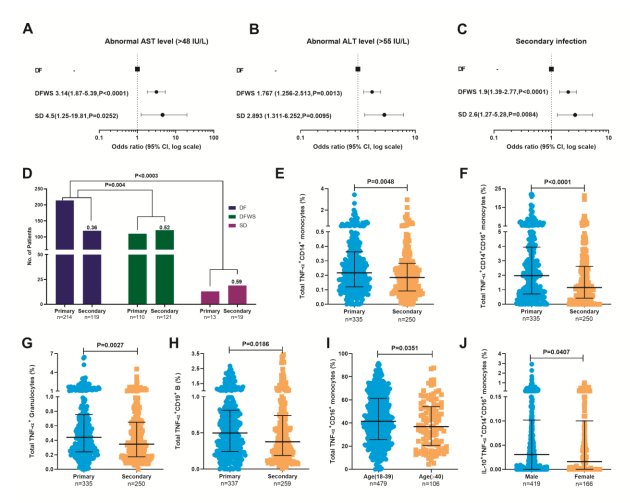


Fig. S8. Factors influencing innate immune cytokine secretion in dengue patients. Forest plot showing association of dengue severity with (A) AST levels, (B) ALT levels and (C) serostatus. Data are odds ratio (95% CI, P value). (D) Z-score test to compare the proportion of primary/secondary dengue in DF, DFWS and SD. Significantly higher frequency of total (E) TNF- α +CD14+ monocytes, (F) TNF- α +CD14+CD16+ intermediate monocytes, (G) TNF- α + granulocytes and (H) TNF- α +CD19+ B cells in primary compared to secondary patients. (I) Significantly higher frequency of total TNF- α +CD16+ non-classical monocytes in young (18-39 years) compared to old (\geq 40 years) patients. (J) Significantly higher frequency of IL-10+TNF- α + CD14+CD16+ intermediate monocytes in men compared to women. P values were determined using Mann-Whitney U test and median with IQR are reported.

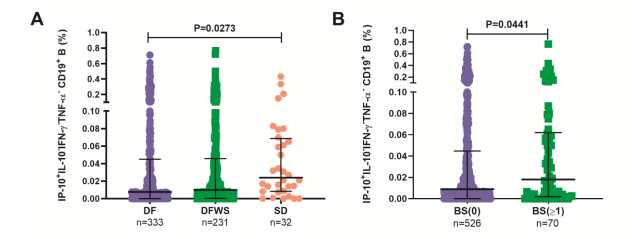


Fig. S9. IP-10 secreting CD19⁺ B cells directly correlate with severity. Frequency of monofunctional IP-10⁺IL-10⁻IFN- γ -TNF- α -CD19⁺ B cells were (**A**) significantly higher in SD compared to DF and (**B**) significantly higher in patients with different degrees of hemorrhage (BS≥1) compared to patients with no bleeding (BS=0). P values were determined using Kruskal-Wallis test, followed by Bonferroni correction for multiple comparisons and using Mann-Whitney *U* test between two groups with median and IQR reported.

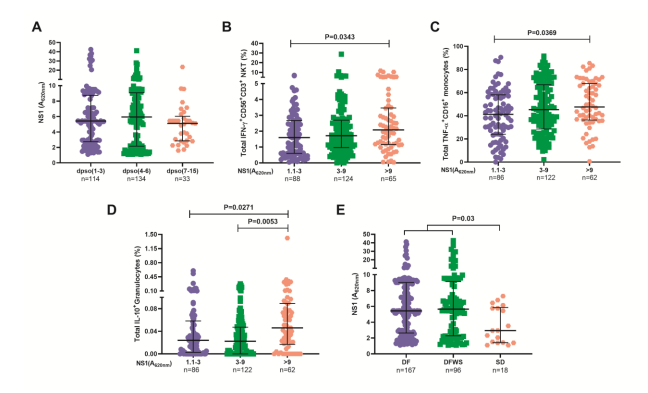


Fig. S10. DENV NS1 protein levels correlate directly with innate immune activation. (**A**) All patients who had detectable dengue specific NS1 levels in serum were compared based on days post symptom onset (dpso). Frequency of total (**B**) IFN- γ +CD56+CD3+ NKT cells, (**C**) TNF- α +CD16+ non-classical monocytes and (**D**) IL-10+ granulocytes compared between patients with low NS1 (A_{620nm} 1.1 to 3), intermediate NS1 (A_{620nm} 3 to 9) and high NS1 (A_{620nm} > 9) values. (**E**) Levels of dengue NS1 compared between DF, DFWS and SD patients. P values were determined using Kruskal-Wallis test, followed by Bonferroni correction for multiple comparison between groups with median and IQR reported.

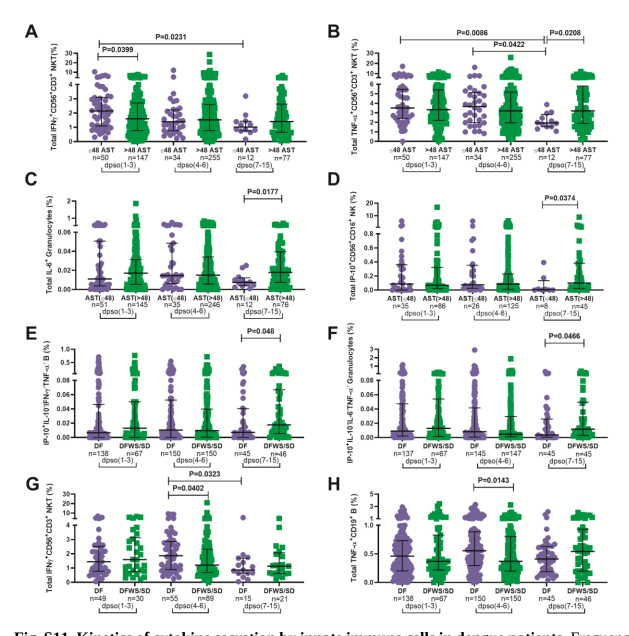


Fig. S11. Kinetics of cytokine secretion by innate immune cells in dengue patients. Frequency of total (**A**) IFN-γ+CD56+CD3+ NKT cells, (**B**) TNF-α+CD56+CD3+ NKT cells, (**C**) IL-6+ granulocytes in total cohort and (**D**) IP-10+CD56+CD16+ NK cells in primary dengue cohort compared between normal (≤48 IU/L) and abnormal (>48 IU/L) levels of AST. Frequency of (**E**) monofunctional IP-10+IL-10-IFN-γ-TNF-α-CD19+ B cells in total cohort, (**F**) monofunctional IP-10+IL-10-IL-6-TNF-α- granulocytes in total cohort, (**G**) total IFN-γ+CD56+CD3+ NKT cells in secondary dengue and (**H**) total TNF-α+CD19+ B cells in total cohort compared between DF and

DFWS/SD severity groups as well as between time window of hospital presentation during 1-3, 4-6 or 7-15 days post symptom onset (dpso). P values were determined using Mann-Whitney *U* test between normal and abnormal levels of AST or DF and DFWS/SD groups for any single time interval and Kruskal-Wallis test, followed by Bonferroni correction for multiple comparison of normal or abnormal AST / DF or DFWS/SD patients between the three time intervals. Medians with IQR are reported.

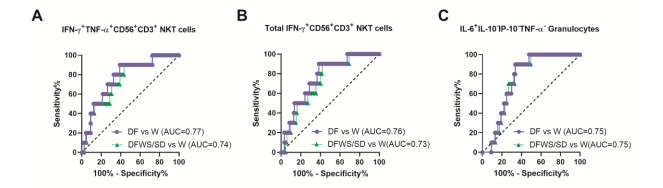


Fig. S12. Biomarker performance of cytokine-secreting innate immune cell subsets. Receiver operating characteristic curve of (**A**) IFN- γ ⁺TNF- α ⁺CD56⁺CD3⁺ NKT cells, (**B**) total IFN- γ ⁺CD56⁺CD3⁺ NKT cells and (**C**) monofunctional IL-6⁺IL-10⁻IP-10⁻TNF- α ⁻ granulocytes in discriminating worsened (W) patients from recovered DF (purple) or DFWS/SD (green) patients. Area under the curves (AUC) were determined by ROC curve analyses.

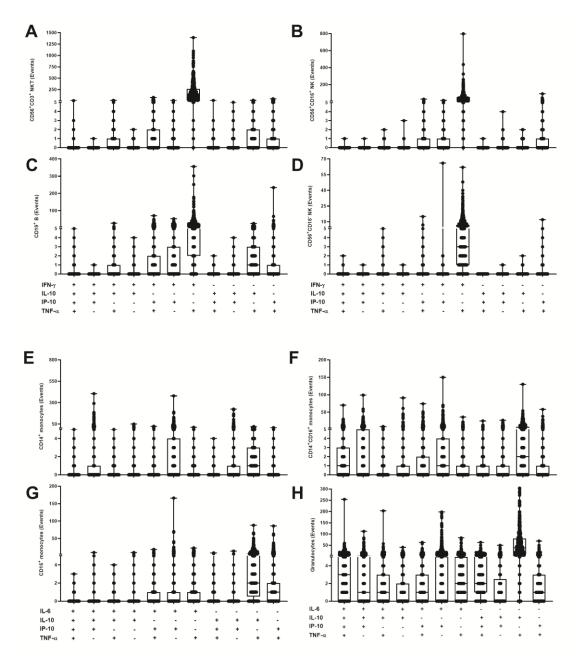


Fig. S13. Polyfunctional profiles of innate immune cells. Polyfunctional cytokine signature of **(A)** CD56⁺CD3⁺ NKT cells **(B)** CD56⁺CD16⁺ NK cells **(C)** CD19⁺ B cells **(D)** CD56⁺CD16⁻ NK cells **(E)** CD14⁺ classical monocytes **(F)** CD14⁺CD16⁺ intermediate monocytes **(G)** CD16⁺ nonclassical monocytes and **(H)** Granulocytes. Cytokine-secreting cell events over the control are plotted as box and whisker plot. Each dot represents a patient.

	IFN-γ	,	IP-10		IL-10		TNF-a	
Cell Subsets	% Producers [Median (IQR)]	MFI (AU x 10 ³)	% Producers [Median (IQR)]	MFI (AU x 10 ³)	% Producers [Median (IQR)]	MFI (AU x 10 ³)	% Producers [Median (IQR)]	$MFI \\ (AU \times 10^3)$
NKT	95.5	2.2	44.3	6.1	33.9	1.1	98.2	3.0
	[1.5 (0.77-2.6)]	(1.8-2.7)	[0.06 (0.004-0.3)]	(5.4-7.4)	[0.04 (0.007-0.1)]	(0.8-1.5)	[3.25 (2.0 -5.3)]	(2.3-4.2)
NK++	81.9	1.6	48.3	5.5	13.6	1.0	95.1	1.5
	[0.29 (0.16-0.56)]	(1.3-1.9)	[0.065(0.1-0.28)]	(4.6-6.9)	[0.009 (0-0.03)]	(0.8-1.5)	[1.007 (0.57-1.61)]	(1.3-1.8)
NK+-	13.8	2.6	21.0	4.8	1.5	0.7	44.1	1.7
	[0.13 (0-0.33)]	(1.3-4.2)	[0.29 (0-0.29)]	(0-6.9)	[0 (0-0.03)]	(0-1.4)	[0.52 (0.2-0.9)]	(1.0-3.1)
B cell	50.8	6.9	38.3	57.3	27.7	12.2	88.6	3.7
	[0.07 (0.02-0.16)]	(5.3-10.6)	[0.02 (0.002-0.049)]	(42.1-228.6)	[0.017 (0.003-0.049)]	(6.2-41.2)	[0.44 (0.22-0.78)]	(2.1-5.9)
	IL-6		IP-10		IL-10		TNF-α	
CM	28.2	41.6	47.0	16.9	35.6	4.4	87.9	2.8
	[0.01 (0-0.14)]	(30.3-57.4)	[0.027 (0-0.13)]	(12.9-22.9)	[0.01 (0-0.05)]	(3.6-5.7)	[0.2 (0.11-0.32)]	(2.3-3.5)
IM	37.8	89.2	41.2	52.2	42.4	6.5	88.7	5.7
	[0.31 (0-0.2)]	(78.8-100.3)	[0.05 (0-0.25)]	(44.1-71.2)	[0.02 (0-0.29)]	(5.6-8.8)	[1.49 (0.5-3.1)]	(4.4-7.4)
NCM	11.5	22.1	15.6	10.9	19.8	1.8	98.8	3.3
	[0.54 (0-0.28)]	(18.4-26.5)	[0.1 (0-0.42)]	(9.6-13.5)	[0.09 (0-0.44)]	(1.5-2.3)	[40.6 (24.6-59)]	(2.2-5.3)
Granulocytes	76.8 [0.015 (0.005- 0.03)]	60.5 (48.1-88.2)	74.4 [0.019 (0.005-0.06)]	38.7 (30.0-52.8)	73.3 [0.025 (0.003-0.05)]	6.8 (5.2-8.6)	98.8 [0.414 (0.20-0.71)]	3.7 (3.6-4.4)

Table S1. Percentage producers and MFI of cytokines from innate cell subsets. % producers describe the percentage of patients in our cohort with detectable cytokine events from each cell subset; median cytokine secreting cells as a percent of parent and IQR are represented within

brackets. The tabulated MFI (AU x 10^3) are median (IQR). Abbreviations: MFI - Median Fluorescence Intensity; AU - Arbitrary Unit; IQR - Inter Quartile Range; NKT - CD56 $^+$ CD3 $^+$ cells; NK $^{++}$ - CD56 $^+$ CD16 $^+$ cells; NK $^{+-}$ - CD56 $^+$ CD16 $^-$ cells; B cells - CD19 $^+$ cells; CM - CD14 $^+$ classical monocytes; IM - CD14 $^+$ CD16 $^+$ intermediate monocytes; NCM - CD16 $^+$ non-classical monocytes.

	Dengu	e (n=596)	FC ((n=58)	НС	(n=43)	P v	alue
Cell Subsets	%	Absolute count (/µl)	%	Absolute count (/µl)	%	Absolute count (/µl)	%	Absolute count (/µl)
Lymphocytes	31.9 (21.9–39.7)	1423 (923–2422)	29.3 (20.8–41.1)	1758 (849–2940)	34.7 (22.8–40.4)	1568 (737–2867)	ns	ns
NK ⁺⁺ cells ^a	6.4 (4.2–9.0)	92 (47–166)	5.7 (3.8–8.8)	106 (56–179)	6.8 (4.3–10.4)	105 (54–150)	ns	ns
NK+- cellsa	1.1 (0.8–1.6)	18 (9–31)	0.9 (0.7–1.3)	17 (8–29)	0.7 (0.4–0.9)	11 (4–19)	< 0.0001	0.0009
NKT cells ^a	4.9 (3.7–7.3)	77 (45–132)	6.9 (4.7–9.3)	126 (67–208)	6.7 (4.8–8.2)	120 (33–186)	< 0.0001	0.0007
B cells ^a	10.6 (7.6–14.5)	153 (81–305)	8.6 (5.3–11.8)	131 (62–241)	8.7 (6.3–13.6)	123 (58–280)	0.0004	ns
T cells ^a	59.4 (52.8–64.9)	815 (448–1373)	57 (51.2–65.8)	1013 (451–1503)	54.3 (49–60.5)	815 (382–1567)	0.0250	ns
	Dengu	e (n=585)	FC ((n=58)	НС	(n=43)		
Monocytes	6.4 (4.7–8.5)	301 (173–464)	5.2 (2.9–6.5)	252 (149–357)	4.3 (3.4–5.4)	179 (113–296)	<0.0001	0.0002
CM^b	68.6 (60.6–76.2)	228 (122–347)	64.9 (53.3–73.9)	178 (104–288)	70.7 (64.9–74.2)	140 (89–238)	0.0229	0.0018
${f IM^b}$	17.3 (12.6–24.7)	53 (31–92)	17.1 (13.6–22.2)	44 (27–91)	9.5 (7.7–11.9)	20 (12–35)	< 0.0001	< 0.0001
NCM^b	3.2 (2.1–4.9)	10 (6–16)	4.8 (2.6–8.3)	11 (8–17)	6.3 (5.0–7.9)	13 (8–21)	< 0.0001	ns
Granulocytes	43.0 (32.6–56.1)	1921 (1275–2932)	52.5 (34.3–63.7)	2551 (1664–4212)	48.1 (39.8–58.0)	1874 (1240–2545)	ns	0.0022

Table S2. Immune cell subsets in dengue and control groups. The presented values are median (IQR). Kruskal-Wallis test followed by Bonferroni correction for multiple comparisons was performed to compare the difference of absolute count /μl blood and % of cell subsets between volunteer groups and the P values are indicated. Number of subjects in each group is given in parentheses. % of cell subsets - (a) represents

percentage of lymphocytes and (b) represents percentage of monocytes. Abbreviations: FC – febrile controls; HC – healthy controls; ns - not significant; IQR – Inter Quartile Range; NKT – CD56+CD3+ cells; NK++ - CD56+CD16+ cells; NK+- - CD56+CD16- cells; B cells – CD19+ cells; T cells – CD3+ cells; CM – CD14+CD16- classical monocytes; IM – CD14+CD16+ intermediate monocytes; NCM – CD14-CD16+ non-classical monocytes.

	DF (n=333)	DFWS	S (n=231)	SD	(n=32)	P v	alue
Cell Subsets	%	Absolute count (/µl)	%	Absolute count (/µl)	%	Absolute count (/µl)	%	Absolute count (/µl)
Lymphocytes	32.3 (22.2–39.3)	1402 (823–2220)	31.1 (21.8–39.7)	1502 (952–2685)	32.2 (19.5–38.6)	1677 (995–2835)	ns	ns
NK ⁺⁺ cells ^a	6.5 (4.3–9.2)	87 (47–165)	6.2 (4.0–8.6)	94 (50–172)	5.2 (2.8–9.6)	100 (32–180)	ns	ns
NK+- cellsa	1.1 (0.8–1.6)	16 (9–29)	1.2 (0.8–1.7)	19 (10–33)	1.3 (0.9–1.6)	20 (14–33)	ns	ns
NKT cells ^a	4.9 (3.8–7.5)	75 (42–132)	4.8 (3.7–7.1)	78 (48–138)	3.9 (3.2–6.0)	80 (36–109)	ns	ns
B cells ^a	9.9 (7.2–14)	142 (72–262)	10.9 (7.7–15.1)	166 (339–87)	12.6 (9.5–17.9)	224 (107–405)	0.0177	0.0144
T cells ^a	60.2 (53.9–65.7)	769 (380–1264)	59.2 (52.4–64.1)	876 (472–1427)	53.2 (44.9–59.6)	803 (555–1380)	<0.0001	ns
	DF (n=327)	DFWS	S (n=227)	SD	(n=31)		
Monocytes	6.2 (4.4–8.5)	283 (162–441)	6.5 (4.8–8.4)	318 (175–496)	6.5 (5.1–7.6)	372 (255–529)	ns	0.0251
$\mathbf{C}\mathbf{M}^{\mathbf{b}}$	68.8 (60.4–76.1)	207 (113–326)	68 (60.6–76.5)	241 (123–377)	68.7 (62.5–76.2)	268 (190–399)	ns	0.0339
IM^b	17.8 (12.6–24.5)	49 (28–88)	16.7 (12.6–24.8)	56 (33–93)	17.4 (10.3–24.1)	57 (40–91)	ns	ns
NCM^b	2.1 (3.3–4.9)	10 (5–15)	3.2 (1.9–4.9)	10 (6–17)	2.9 (2.1–3.8)	13 (7–17)	ns	ns
Granulocytes	43 (32.7–56.7)	1844 (1215–2706)	43.5 (31.9–55.7)	2019 (1290–3128)	43.1 (35.2–51.9)	2574 (1762–3160)	ns	0.0184

Table S3. Cell subsets as a function of dengue severity. The presented values are median (IQR). Kruskal-Wallis test followed by Bonferroni correction for multiple comparisons was performed to compare the difference of absolute count/µl blood and % of cell subsets between dengue severity groups and the P values are indicated. Number of subjects in each group is given in parentheses. % of cell subsets - (a) represents percentage

of lymphocytes and (b) represents percentage of monocytes. Abbreviations: DF – dengue fever; DFWS – dengue fever with warning signs; SD – severe dengue; ns - not significant; IQR – Inter Quartile Range; NKT – CD56⁺CD3⁺ cells; NK⁺⁺ - CD56⁺CD16⁺ cells; NK⁺⁻ - CD56⁺CD16⁻ cells; B cells – CD19⁺ cells; T cells – CD3⁺ cells; CM – CD14⁺CD16⁻ classical monocytes; IM – CD14⁺CD16⁺ intermediate monocytes; NCM – CD14⁻ CD16⁺ non-classical monocytes.

	n	Spearman r (95% CI)	P value
TNF- α^+ NK ⁺⁺ & TNF- α^+ NKT cells	596	0.77 (0.73-0.80)	< 0.0001
$TNF-\alpha^+ NK^{++}$ & $TNF-\alpha^+ B$ cells	596	0.64 (0.58-0.68)	< 0.0001
$TNF-\alpha^+ NK^{++} \& TNF-\alpha^+$ granulocytes	582	0.64 (0.58-0.68)	< 0.0001
TNF- α^+ CM & TNF- α^+ granulocytes	585	0.71 (0.67-0.75)	< 0.0001
TNF- α^+ IM & TNF- α^+ granulocytes	585	0.63 (0.59-0.68)	< 0.0001
TNF- α^+ NK ⁺⁺ & IFN- γ^+ TNF- α^+ NKT cells	596	0.73 (0.69-0.77)	< 0.0001
IFN- γ^+ TNF- α^+ NKT & IFN- γ^+ TNF- α^+ NK ⁺⁺ cells	596	0.76 (0.72-0.79)	< 0.0001
IFN- γ^+ TNF- α^+ NKT & IFN- γ^+ NK ⁺⁺ cells	596	0.61 (0.56-0.66)	< 0.0001
IFN- γ^+ TNF- α^+ NK ⁺⁺ & IFN- γ^+ NKT cells	596	0.72 (0.67-0.75)	< 0.0001
IFN- γ^+ NKT & IFN- γ^+ NK ⁺⁺ cells	596	0.71 (0.66-0.74)	< 0.0001
IFN-γ ⁺ NKT & TNF-α ⁺ NK ⁺⁺ cells	596	0.69 (0.64-0.73)	<0.0001

Table S4. Correlation analysis of cytokines between subsets. Spearman's correlation coefficient and P values for cytokines from various subsets are reported. Abbreviations: n – number of samples; NKT – CD56⁺CD3⁺ cells; NK⁺⁺ - CD56⁺CD16⁺ cells; B cells – CD19⁺ cells; CM – CD14⁺CD16⁻ classical monocytes; IM – CD14⁺CD16⁺ intermediate monocytes.

Subset and cytokine	Patient groups	AUC (95% CI)	Cut-off	Sensitivity (%)	Specificity (%)	Likelihood Ratio
IFN-γ+CD56+CD3+	Total	0.76 (0.64-0.88)	<1.202 ^a	90	61.56	2.34
NKT cells	Abnormal liver enzymes	0.83 (0.74-0.92)	<1.216 a	100	60.32	2.52
IFN-γ ⁺ TNF-α ⁺	Total	0.77 (0.64-0.90)	<1.072 a	90	60.96	2.31
CD56 ⁺ CD3 ⁺ NKT cells	Abnormal liver enzymes	0.83 (0.74-0.93)	<1.072 a	100	59.52	2.47
IL-6+ Granulocytes	Total	0.75 (0.67-0.83)	<0.005 a	90	66.36	2.67
II. 6+ Cuanula autos	Total	0.85 (0.78-0.92)	>0.042 b	90	75.93	3.73
IL-6 ⁺ Granulocytes, IFN-γ ⁺ CD56 ⁺ CD3 ⁺ NKT cells	Abnormal AST	0.90 (0.85-0.95)	>0.056 b	100	81.9	5.53
	Abnormal liver enzymes	0.88 (0.83-0.94)	>0.51 ^b	100	75.7	4.12

Table S5. Biomarker performance of IFN-γ+CD56+CD3+ NKT cells, IFN-γ+TNF-α+CD56+CD3+ NKT cells and monofunctional IL-6+granulocytes. The tabulated values are AUC, cut-off used, sensitivity (%), specificity (%) and likelihood ratio from ROC analysis in total and homogeneous patient groups. ROC analysis was carried out between recovered DF patients and those who worsened during the study. Abbreviations: AUC – area under the curve; CI – confidence interval; ROC – receiver operating characteristic; DF – dengue fever. Cut off represents (a) % cytokine secreting cells or (b) probability obtained from regression analysis.

Target (Clone)	Species	Fluorochrome/Conjugate	Company	Catalog no.
	NK cell pa	anel – Surface Markers		
CD19 (HIB19)	Mouse IgG1, κ	PE-Cy5	BioLegend	302210
CD56 (B159)	Mouse IgG1, κ	PE-Cy7	BD Bioscience	557747
CD3 (SK7)	Mouse BALB/cIgG1, κ	APC-H7	BD Bioscience	560176
CD16 (3G8)	Mouse CDF1 IgG1, κ	BV510	BD Bioscience	563830
	Monocyte l	Panel – Surface Markers		
CD14 (M5E2)	Mouse IgG2a, κ	PerCP- Cy5.5	BD Bioscience	550787
TLR2 (11G7)	Mouse IgG1, κ	BV510	BD Bioscience	742767
CD16 (3G8)	Mouse CDF1 IgG1, κ	BV605	BD Bioscience	563172
	Intracellu	ılar Cytokine Markers		
IP-10 (6D4/D6/G2)	Mouse IgG2a, κ	PE	BD Bioscience	555049
IL-10 (JES3-19F1)	Rat IgG2a, κ	APC	BD Bioscience	554707
IFN-γ (B27)	Mouse IgG1, κ	BV605	BD Bioscience	562974
TNF-α (MAb11)	Mouse IgG1, κ	BV750	BD Bioscience	566359
IL-6 (MQ2-6A32)	Rat IgG2a, κ	FITC	BD Bioscience	557696

Table S6. Antibodies used for flow cytometry.

Cell subset	II	ΤΝ-γ	IL-	10	IP	P-10	T	NF-α
Events	Minimum	over control						
NKT	15	twice	10	10	10	10	15	twice
NK++	15	15	10	10	10	10	15	twice
NK+-	15	twice	10	10	10	10	15	10
В	15	10	10	10	10	10	15	15
	I	L-6	IL-	10	IP	P-10	T	NF-α
CM	10	10	10	10	10	10	15	15
IM	10	10	10	10	10	10	15	15
NCM	10	10	10	10	10	10	15	twice
Granulocytes	15	15	15	15	15	15	15	twice

Table S7. **Criteria used to define positive responses in test samples.** The minimum number of events and the number of events over the control used to define a positive response in test samples for all cytokines from innate cell subsets are reported. Abbreviations: NKT – CD56⁺CD3⁺ cells; NK⁺⁺ - CD56⁺CD16⁺ cells; NK⁺⁻ - CD56⁺CD16⁻ cells; B cells – CD19⁺ cells; CM – CD14⁺ classical monocytes; IM – CD14⁺CD16⁺ intermediate monocytes; NCM – CD16⁺ non-classical monocytes.

Call Cubacta	Erronta	TN	lF-α	IL	-6	IP-	10	IL.	-10
Cell Subsets	Events	Control	Test	Control	Test	Control	Test	Control	Test
	Median	4	70	5	9	5	19	10	17
\mathbf{CM}	IQR	2-8	33-128	2-17	4-28	1-20	4-78	3-35	7-44
	Range	0-122	0-1557	0-402	0-1066	0-727	0-2388	0-759	0-1146
	Median	37	191	15	21	18	24	31	37
IM	IQR	16-89	86-341	8-29	9-45	8-33	12-47	17-65	16-79
	Range	1-2044	3-3312	1-843	0-709	0-1191	0-507	1-441	0-711
	Median	2	564	2	3	1	4	4	6
NCM	IQR	1-4	324-927	0-5	1-8	0-4	1-9	1-10	2-15
	Range	0-385	0-2686	0-107	0-205	0-66	0-229	0-107	0-157
	Median	38	1272	21	76	23	91	92	180
Granulocytes	IQR	23-66	637-2136	9-47	40-138	11-49	40-223	56-147	110-268
	Range	4-1565	38-31503	0-709	1-969	0-883	2-3534	9-1210	10-1079
		TN	IF-α	IF	V -γ	IP-	10	IL.	-10
	Median	6	364	4	160	2	12	4	11
NKT	IQR	3-13	186-665	1-8	81-310	0-7	4-35	1-9	4-23
	Range	0-188	0-2578	0-129	1-2786	0-376	0-995	0-65	0-1032
	Median	1	131	3	44	2	13	2	4
NK++	IQR	0-2	63-253	1-8	22-86	0-5	4-47	0-6	1-10
	Range	0-22	1-1977	0-85	0-1147	0-305	0-3825	0-46	0-59
	Median	0	12	0	4	0	2	0	1
NK+-	IQR	0-1	5-25	0-0	1-10	0-1	0-9	0-1	0-2
	Range	0-31	0-226	0-135	0-1188	0-335	0-383	0-18	0-49
	Median	17	118	7	26	2	10	4	9
В	IQR	8-37	53-251	2-18	11-54	1-7	3-32	2-9	5-17
	Range	0-398	0-1764	0-108	0-430	0-318	0-304	0-61	0-92

Table S8. **Total number of cytokine-secreting innate immune cell events from control and test samples.** The median, IQR and range are reported. Abbreviations: IQR – Inter Quartile Range; CM – CD14⁺ classical monocytes; IM – CD14⁺CD16⁺ intermediate monocytes; NCM – CD16⁺ non-classical monocytes NKT – CD56⁺CD3⁺ cells; NK⁺⁺ - CD56⁺CD16⁺ cells; NK⁺⁻ - CD56⁺CD16⁻ cells; B cells – CD19⁺ cells.

]	Dengue Pa	atients Rec	ruited				
	KI	MS	BMe	CRI	RM	СH	SJN	МС	To	tal
	N	%	N	%	N	%	N	%	N	%
Male	147	80	65	65	80	60	136	76	428	72
Female	36	20	35	35	53	40	44	24	168	28
Total	183		100		133		180		596	
				Total Der	ngue Admis	ssions				
	KI	MS	BM	CRI	RM	СН	SJN	МС	To	tal
	N	%	N	%	N	%	N	%	N	%
Male	2383	68	935	54	642	60	1280	59	5238	62
Female	1139	32	798	46	429	40	898	41	3265	38
Total	3522		1733		1071		2178		8503	

Table S9. Gender distribution among dengue patients. Numbers of total dengue patients admitted and those recruited for our study (between June 24, 2019 and November 29, 2019) in each hospital are listed according to gender. Abbreviations: N – Number of patients; % - percentage of male/female patients; KIMS - Kempegowda Institute of Medical Sciences; BMCRI - Bangalore Medical College and Research Institute; SJMC - St. John's Medical College; RMCH - M S Ramaiah Medical College and Hospital.

	DF	DFWS	SD		P value	
Variable	(n=333)	(n=231)	(n=32)	DF-DFWS	DF- SD	DFWS-SD
Age; median	26.8	28	29		0.098 ^a	
(IQR)	(21.8, 35.04)	(22, 36.3)	(26.1, 43)		0.070	
dpso;	4	4	4		0.0008^{a}	
median(IQR)	(3,5)	(4,5)	(3,5)		0.0000	
Male : Female	238:95	164:76	26:6	0.902	0.238	0.225
Nausea; n (%)	127 (38.1)	127 (55)	18 (56.3)	< 0.0005	0.046	0.89
Head ache; n (%)	155 (46.5)	132 (57.1)	16 (50)	0.013	0.708	0.446
Abdomen Pain; n (%)	0 (0)	109 (18.3)	16 (50)	< 0.0005	<0.0005*	0.76
Body Pain; n (%)	171 (51.4)	146 (63.2)	14 (43.8)	0.005	0.411	0.035
Rashes; n (%)	22(6.6)	25(10.8)	2(6.3)	0.075	1*	0.549*
Splenomegaly; n (%)	3(0.9)	8(3.5)	2(6.3)	0.03*	0.063*	0.349*
Hepatomegaly; n (%)	0	24(10.5)	4(12.5)	< 0.0005	<0.0005 ^a	0.536*
Petechiae; n (%)	1(0.3)	13(5.6)	0	< 0.0005	1*	0.378*
Ascites; n (%)	0	39(16.9)	7(21.9)	< 0.0005	<0.0005*	0.486
Primary : Secondary	214:118	109:120	13:18	< 0.0005	0.008	0.457
Hemorrhage; n (%)	2 (0.6)	47 (20)	21 (65)	<0.0001*	<0.0001*	<0.0001*

Table S10. Demographic characteristics and clinical symptoms of dengue patients. Age and days of fever at presentation are represented as median (IQR). Statistical differences in proportion of gender, clinical symptoms and primary/secondary infection between dengue patient groups based on severity was determined by either Chi-Square or Fisher's exact test. P values are tabulated; ^a - Kruskal-Wallis test; * - Fisher's exact test. Abbreviations: dpso – days post symptom onset; IQR – Inter Quartile Range; n – number of patients; DF – dengue fever; DFWS – dengue fever with warning signs; SD – severe dengue.

Variable	DF (n=333)	DFWS (n=231)	SD (n=32)	P value	Statistical test
Hemoglobin at enrollment; mean (SD)	14.43 (1.95)	14.48 (2.1)	15.14 (2.25)	0.10	ANOVA
Lowest Platelet count (x10 ³); median (IQR)	50 (27,85)	29 (14-56)	16 (10,32)	<0.0001	Kruskal-Wallis test
Highest Hematocrit %; mean (SD)	42.9 (5.23)	43.84 (5.5)	44.49 (6.4)	0.09	ANOVA
Albumin g/dL; median (IQR)	3.8 (3.5,4.1)	3.6 (3.3,4)	3.4 (3.2,3.7)	<0.0001	Kruskal-Wallis test
Aspartate transaminase IU/L; median (IQR)	86 (50,143)	114 (75,189)	123.5 (91,880)	<0.0001	Kruskal-Wallis test
Alanine transaminase IU/L; median (IQR)	56 (29.5,95.5)	70 (45, 118)	98 (51, 265)	<0.0001	Kruskal-Wallis test

Table S11. Clinical laboratory parameters of dengue patients. Hemoglobin and highest hematocrit are represented as mean (standard deviation); Lowest platelet count, albumin, aspartate transaminase and alanine transaminase as median (IQR). Based on normality test, ANOVA test was used for parametric data and Kruskal-Wallis test for non-parametric data to find significant difference across dengue severity based on clinical parameters. Abbreviations: IQR – Inter Quartile Range; DF – dengue fever; DFWS – dengue fever with warning signs; SD – severe dengue.

Section & Topic	No	Item	Reported on pag
TITLE OR ABSTRACT			
	1	Identification as a study of diagnostic accuracy using at least one measure of accuracy (such as sensitivity, specificity, predictive values, or AUC)	Main page 1,2
ABSTRACT			
	2	Structured summary of study design, methods, results, and conclusions	Main page 1,2
		(for specific guidance, see STARD for Abstracts)	
INTRODUCTION			
	3	Scientific and clinical background, including the intended use and clinical role of the index test	Main page 2,3
	4	Study objectives and hypotheses	Main page 2,3
METHODS			р-8/-
Study design	5	Whether data collection was planned before the index test and reference standard	Supplementary
cady acsign	•	were performed (prospective study) or after (retrospective study)	page 2,3
Participants	6	Eligibility criteria	Supplementary
- articipants	U	Liigibility Citieria	page 2,3
	7	On what basis potentially eligible participants were identified	Supplementary
		(such as symptoms, results from previous tests, inclusion in registry)	page 2,3
	8	Where and when potentially eligible participants were identified (setting, location and dates)	Supplementary
			page 2
	9	Whether participants formed a consecutive, random or convenience series	Supplementary page 2
Test methods	10a	Index test, in sufficient detail to allow replication	Supplementary page 4,5
	10b	Reference standard, in sufficient detail to allow replication	Supplementary page 2
	11	Rationale for choosing the reference standard (if alternatives exist)	n/a
	12a	Definition of and rationale for test positivity cut-offs or result categories	n/a
		of the index test, distinguishing pre-specified from exploratory	
	12b	Definition of and rationale for test positivity cut-offs or result categories	n/a
		of the reference standard, distinguishing pre-specified from exploratory	
	13a	Whether clinical information and reference standard results were available	Supplementary
		to the performers/readers of the index test	pages 2, 5
	13b	Whether clinical information and index test results were available	Supplementary
		to the assessors of the reference standard	page 5
Analysis	14	Methods for estimating or comparing measures of diagnostic accuracy	Supplementary
			page 5,6
	15	How indeterminate index test or reference standard results were handled	Supplementary page 3
	16	How missing data on the index test and reference standard were handled	Supplementary page 3
	17	Any analyses of variability in diagnostic accuracy, distinguishing pre-specified from exploratory	n/a
	18	Intended sample size and how it was determined	Supplementary
RESULTS			page 2
Participants	19	Flow of participants, using a diagram	Supplementary page 9
	20	Baseline demographic and clinical characteristics of participants	Supplementary
	21a	Distribution of severity of disease in those with the target condition	page 37,38 Supplementary
	21b	Distribution of alternative diagnoses in those without the target condition	page 37,38 n/a
	210	Time interval and any clinical interventions between index test and reference standard	n/a
Test results	23	Cross tabulation of the index test results (or their distribution)	Main page 5,6
rest resuits	23	by the results of the reference standard	iviaiii page 3,0



	24	Estimates of diagnostic accuracy and their precision (such as 95% confidence intervals)	Supplementary page 31
	25	Any adverse events from performing the index test or the reference standard	n/a
DISCUSSION			
	26	Study limitations, including sources of potential bias, statistical uncertainty, and generalisability	Main page 11,12
	27	Implications for practice, including the intended use and clinical role of the index test	Main page 11,12
OTHER INFORMATION			
	28	Registration number and name of registry	n/a
	29	Where the full study protocol can be accessed	n/a
	30	Sources of funding and other support; role of funders	Main page 15



STARD 2015

AIM

STARD stands for "Standards for Reporting Diagnostic accuracy studies". This list of items was developed to contribute to the completeness and transparency of reporting of diagnostic accuracy studies. Authors can use the list to write informative study reports. Editors and peer-reviewers can use it to evaluate whether the information has been included in manuscripts submitted for publication.

EXPLANATION

A diagnostic accuracy study evaluates the ability of one or more medical tests to correctly classify study participants as having a target condition. This can be a disease, a disease stage, response or benefit from therapy, or an event or condition in the future. A medical test can be an imaging procedure, a laboratory test, elements from history and physical examination, a combination of these, or any other method for collecting information about the current health status of a patient.

The test whose accuracy is evaluated is called **index test.** A study can evaluate the accuracy of one or more index tests. Evaluating the ability of a medical test to correctly classify patients is typically done by comparing the distribution of the index test results with those of the **reference standard**. The reference standard is the best available method for establishing the presence or absence of the target condition. An accuracy study can rely on one or more reference standards.

If test results are categorized as either positive or negative, the cross tabulation of the index test results against those of the reference standard can be used to estimate the **sensitivity** of the index test (the proportion of participants *with* the target condition who have a positive index test), and its **specificity** (the proportion *without* the target condition who have a negative index test). From this cross tabulation (sometimes referred to as the contingency or "2x2" table), several other accuracy statistics can be estimated, such as the positive and negative **predictive values** of the test. Confidence intervals around estimates of accuracy can then be calculated to quantify the statistical **precision** of the measurements.

If the index test results can take more than two values, categorization of test results as positive or negative requires a **test positivity cut-off**. When multiple such cut-offs can be defined, authors can report a receiver operating characteristic (ROC) curve which graphically represents the combination of sensitivity and specificity for each possible test positivity cut-off. The **area under the ROC curve** informs in a single numerical value about the overall diagnostic accuracy of the index test.

The **intended use** of a medical test can be diagnosis, screening, staging, monitoring, surveillance, prediction or prognosis. The **clinical role** of a test explains its position relative to existing tests in the clinical pathway. A replacement test, for example, replaces an existing test. A triage test is used before an existing test; an add-on test is used after an existing test.

Besides diagnostic accuracy, several other outcomes and statistics may be relevant in the evaluation of medical tests. Medical tests can also be used to classify patients for purposes other than diagnosis, such as staging or prognosis. The STARD list was not explicitly developed for these other outcomes, statistics, and study types, although most STARD items would still apply.

DEVELOPMENT

This STARD list was released in 2015. The 30 items were identified by an international expert group of methodologists, researchers, and editors. The guiding principle in the development of STARD was to select items that, when reported, would help readers to judge the potential for bias in the study, to appraise the applicability of the study findings and the validity of conclusions and recommendations. The list represents an update of the first version, which was published in 2003.

More information can be found on http://www.equator-network.org/reporting-guidelines/stard.

