Chymase Level Is a Predictive Biomarker of Dengue Hemorrhagic Fever in Pediatric and Adult Patients

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Background. Most patients with dengue experience mild disease, dengue fever (DF), while few develop the life-threatening diseases dengue hemorrhagic fever (DHF) or dengue shock syndrome (DSS). No laboratory tests predict DHF or DSS. We evaluated whether the serum chymase level can predict DHF or DSS in adult and pediatric patients and the influence of preexisting conditions (PECs) on chymase levels.

Methods. Serum chymase levels were measured in patients presenting with undifferentiated fever to hospitals in Colombo District, Sri Lanka. The value of serum the chymase concentration and clinical signs and symptoms as predictors of DHF and/or DSS was evaluated by multivariate analysis. We assessed the influence of age, PECs, and day after fever onset on the robustness of the chymase level as a biomarker for DHF and/or DSS.

Results. An elevated chymase level in acute phase blood samples was highly indicative of later diagnosis of DHF or DSS for pediatric and adult patients with dengue. No recorded PECs prevented an increase in the chymase level during DHF. However, certain PECs (obesity and cardiac or lung-associated diseases) resulted in a concomitant increase in chymase levels among adult patients with DHF.

Conclusions. These results show that patients with acute dengue who present with high levels of serum chymase consistently are at greater risk of DHF. The chymase level is a robust prognostic biomarker of severe dengue for adult and pediatric patients.

Keywords. dengue virus; dengue fever; DHF/DSS; prognosis; biomarker; chymase; mast cells; pediatric dengue.

Dengue is an arboviral infection spread by mosquitoes in the tropical and subtropical regions of the world [1]. It is caused by dengue virus (DENV), which has 4 distinct serotypes. Between 50 million and 400 million DENV infections occur yearly [2]. In DENV-endemic countries, such as the site of this study, Sri Lanka [3], many patients experience multiple DENV infections in a lifetime. An incubation period of approximately 4–7 days occurs before fever onset [4, 5]. Signs such as leukenaemia, thrombocytopoenia, and rash and symptoms such as headache, vomiting, and retro-orbital eye pain are common during acute disease and usually resolve within 3–7 days for those who experience mild disease, diagnosed as dengue fever (DF) [6, 7]. Mild hemorrhagic manifestations such as petechiae and purpura are also common during this time [6, 7]. Resolution of the febrile response usually occurs around days 4–7, when most patients completely recover, but other individuals can suddenly experience complications [6]. Warning signs have been identified to aid in recognizing patients on course to develop dengue hemorrhagic fever (DHF) or dengue shock syndrome (DSS), also known as severe dengue [8]. Warning signs include mucosal bleeding, liver enlargement, abdominal pain, rapid hematocrit increase, persistent vomiting, and clinical fluid accumulation [8]. Many of these signs are subjective and present very late. DHF and DSS are extremely difficult to predict, and there is an urgent need to identify risk factors and biomarkers of severe dengue. Although some studies associated high serum viral titers with DHF and DSS [9, 10], others have reported conflicting information [11–13]. Multiple factors could explain these discrepancies, including faster clearance of viremia during secondary infection [14, 15]. There is an unmet need for diagnostic tests to identify severe DENV during the acute phase of disease, since clinical symptoms often fail to predict it with sufficient warning, and to understand whether those tests are robust in multiple patient populations.
Chymase Level Is a Predictive Biomarker of DHF

Differences in the severity of DENV disease between adult and pediatric cases have been identified [16, 17]. For example, in recent outbreaks in the Philippines, the symptomatic disease burden primarily involved children [18]. In Thailand, signs such as epistaxis, oliguria, and liver enlargement were more prevalent among children than among adults [17]. It is unclear whether there may be differences in the pathophysiological mechanisms in pediatric and adult patients with dengue or whether prior immunity in the adult population can account for the observed differences in severity. One study suggested that children have an inherently higher capillary fragility, possibly explaining why small children present more often with DSS than adults do [19]. In contrast, in some populations, including residents of Singapore, morbidity and mortality are more common in adult and elderly patients, potentially because of the increased incidence of comorbidities [20, 21]. Preexisting conditions (PECs) such as diabetes [22] and obesity [23] have also been associated with severe disease in some studies. The observation in the CYD-TDV (Dengvaxia) trials that a greater percentage of children aged 2–5 years than those >5 years required hospitalization within 3 years of infection emphasizes the importance of evaluating DENV diagnostic tests in multiple age groups [24].

In humans, immune pathology is thought to underlie severe disease [7]. One immune cell type, the mast cell (MC), is critical to the host response to DENV [25–27]. MCs are granulated cells that regulate the host vasculature during inflammation and homeostasis [28]. DENV activation of MCs prompts them to degranulate, which involves the release of their presynthesized inflammatory mediators from intracellular stores [26]. Most granule-associated mediators are proteases, including chymase, which is highly specific to MCs [29]. Chymase is a serine protease and angiotensin-converting enzyme [30]. MC degranulation occurs rapidly. During anaphylaxis, MC-derived mediators are the underlying cause of shock and death and can be detected in the serum as biomarkers [31]. Our prior studies showed that DENV activates MCs, resulting in chymase release, and that MC activation is augmented by DENV-specific antibodies, as are present during secondary infections [25–27, 32]. This led to the observation in a retrospective study of adult DENV-infected patients in Singapore that chymase levels were elevated in acute-phase serum samples from patients with DHF [25].

Here, we questioned whether the MC-derived protease chymase is a robust predictor of DHF and/or DSS in multiple patient populations and we assessed whether PECs attenuate the predictive capacity of chymase. Febrile patients were recruited at study sites in Colombo District, Sri Lanka [3, 33]. Patients’ signs and symptoms were recorded, and their disease severity was documented. Acute-phase serum samples were used to measure the chymase level to assess its potential to predict DHF and/or DSS. Our analysis reveals that the chymase level is a prognostic biomarker for DHF and DSS in pediatric and adult patients, including those with PECs.
15%), and most patients (74%) had secondary infections [3]. On average, patients were enrolled a mean duration (±SD) of 4.43 ± 1.2 days after fever onset (Figure 1B), and there was no difference in the average day of enrollment between patients who received a diagnosis of DF and those who received a diagnosis of DHF (Figure 1C). Of the patients for whom the chymase level was measured, 91% were confirmed to be positive for DENV by either immunoglobulin M, NS1, and/or polymerase chain reaction testing. The blood sample in which the serum chymase level was measured was collected at the time of study enrollment, when 45.2% of patients were admitted with a diagnosis of DF, 7.2% were admitted with a diagnosis of undifferentiated fever or viral fever, 1.1% were admitted with a diagnosis of DHF, and 0.8% were admitted with a diagnosis of urinary tract infection; for 45.4%, the diagnosis was either undetermined or unreported. Only 4% of patients who ultimately had a diagnosis of DHF received this diagnosis at the time of admission, which is consistent with the prevalent observation that the clinical course of DENV disease is difficult to predict during acute disease.

Independent variables associated with a diagnosis of DHF were investigated by multivariate logistic regression analysis (Table 1). Although chymase concentrations were measured in the first blood sample, collected at the time of enrollment, data on the remaining variables were recorded if they were recorded at any time between enrollment and study completion. Of the variables considered, serum chymase concentration at the time of presentation was a highly predictive biomarker for a DHF diagnosis for all patients with a diagnosis of DENV infection and for those with confirmed DENV infection (Table 1). Even after adjustment for the other variables, the odds of having DHF was 1.32 times more likely than the odds of having DF (adjusted odds ratio, 1.32; 95% confidence interval, 1.21–1.44) with every unit increase in serum chymase level (Table 1). As expected, many warning signs associated with subsequent development of severe dengue, such as hepatic and abdominal
Table 1. Serum Chymase Level and Symptoms Associated With Diagnosis of Dengue Fever (DF) Versus Dengue Hemorrhagic Fever (DHF) Among 334 Patients With a Diagnosis of Dengue Virus (DENV) Infection and the Subset of 297 Patients With Confirmed DENV Infection

<table>
<thead>
<tr>
<th>Dengue Sign or Symptom</th>
<th>OR (95% CI)</th>
<th>aOR (95% CI)</th>
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<tbody>
<tr>
<td>Chymase level</td>
<td>1.31³ (1.20–1.42)</td>
<td>1.32³ (1.21–1.44)</td>
</tr>
<tr>
<td>Bleeding per vaginam</td>
<td>3.19 (1.15–8.83)</td>
<td>1.79 (0.48–6.61)</td>
</tr>
<tr>
<td>Abdominal pain</td>
<td>2.31 (1.27–4.20)</td>
<td>1.35 (0.54–3.41)</td>
</tr>
<tr>
<td>Abdominal tenderness</td>
<td>2.29 (1.21–4.33)</td>
<td>2.57 (0.69–9.64)</td>
</tr>
<tr>
<td>Hepatic tenderness</td>
<td>2.08 (1.06–4.08)</td>
<td>0.72 (0.21–2.52)</td>
</tr>
<tr>
<td>Narrow pulse pressure</td>
<td>4.04 (0.95–17.2)</td>
<td>2.78 (0.50–15.4)</td>
</tr>
<tr>
<td>Oliguria</td>
<td>2.72 (1.20–6.20)</td>
<td>2.03 (0.67–6.18)</td>
</tr>
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Abbreviations: aOR, adjusted odds ratio; CI, confidence interval; OR, odds ratio.

³Data are for 223 with DF and 99 with DHF.
³Per unit increase.
³Data are for 174 DF and 73 with DHF.

pains or tenderness, were also significantly associated with DHF. Also, as expected, a narrow pulse pressure was significantly associated with a DHF diagnosis for patients who received a DENV diagnosis (Table 1) but not for the subset with confirmed DENV infection (P = .06; Table 1 and Supplementary Table 2). For patients with confirmed DENV infection, diarrhea was significantly associated with receipt of a DHF diagnosis (Table 1). Hemorrhagic manifestations, aside from bleeding per vaginam, were present in both DF and DHF groups and did not discriminate between them (Table 1 and Supplementary Tables 1 and 2). All recorded variables that were not significantly linked with DHF are included in Supplementary Tables 1 and 2. We found that an early increase in the serum chymase level was a significant predictor for subsequent diagnosis of DHF in DENV-infected patients before the signs were clinically apparent. The significant link between chymase level and a DHF diagnosis was highly correlated for patients with confirmed DENV infection (P ≤ .0001; Supplementary Table 2). Serum chymase levels were unknown to the clinicians at the time of diagnosis. Since approximately 4% of patients with DHF received a DHF diagnosis upon admission, we repeated the analysis without those patients to ensure that chymase levels were truly prognostic. Indeed, the association between chymase level and DHF remained highly significant (P = .028). Thus, the serum chymase level is a prognostic biomarker of DHF.

We next compared the mean serum chymase concentrations of patients grouped on the basis of their final diagnosis: all patients with a diagnosis of DF; a subset of patients with a diagnosis of DF who had signs of bleeding (ie, the DFWB group), which was used to understand whether chymase levels also correlated with mild/moderate bleeding; and patients with a diagnosis of DHF. Patients with a diagnosis of DHF had significantly higher serum chymase levels than all patients with a diagnosis of DF and patients with DFWB. Interestingly, patients in the DFWB group, who had overt symptoms of bleeding yet did not meet the DHF diagnosis criteria, formed a discrete group that differed significantly from the DF and DHF groups (Figure 2A). Similar results were obtained when the analysis was limited to the majority of patients with laboratory-confirmed dengue (Supplementary Figure 1). Grouping by age revealed that children and adolescents with DHF had higher chymase levels than adults with DHF (Figure 2B). The chymase level was also significantly elevated in the serum of patients with DHF, compared with the level in patients with DF, irrespective of the day of enrollment after fever onset (Figure 2C). Thrombocytopenia is a sign of both DF and DHF, although a rapid decline in the platelet count can be a warning sign of severe dengue [8]. A platelet count of <100 000 is one criterion for a DHF diagnosis [8, 35]. However, platelet counts for patients in the DF and DHF groups at the time of study enrollment did not differ (Figure 2D), even though, at later time points, differences in platelet counts became apparent (Supplementary Figure 2). These data show that chymase levels highly correlate with DENV disease severity.

To test how rigorously the chymase concentration performs as a biomarker of severe DENV, we used a ROC analysis. Curves were generated by plotting sensitivity against specificity for chymase level as a predictor of DENV severity. In this test, accuracy
is measured by the area under the ROC curve, where, by convention, an area of 0.9–1 is considered “excellent,” an area of 0.8–0.9 is considered “good,” an area of 0.7–0.8 is considered “fair,” and an area of <0.7 is considered “poor.” The chymase level was, overall, a good predictor of DENV severity (Figure 3A). When evaluated in different age groups, chymase concentration was an excellent predictor of severity in children and adolescent DENV-infected patients (Supplementary Figure 3A–C). Although the chymase level was an excellent predictor of severity in adult patients in Singapore who were recruited ≤72 hours after fever onset (Supplementary Figure 4) [25], in this Sri Lankan cohort, where patients were recruited over a longer time (average, 4.4 days after fever), the chymase level was a good predictor of severity (Supplementary Figure 3D). The chymase level was a weaker predictor of severity for patients aged >45 years, which included elderly adults (Supplementary Figure 3E). However, since most patients enrolled very late after fever onset, we performed the same analysis for patients enrolled <4 days after fever onset. This showed that the chymase concentration is an excellent predictor of severity as an early biomarker (Figure 3B). The ROC analysis results were highly similar for the majority of patients with confirmed DENV infection (Figure 3C), compared with all patients with a diagnosis of DENV infection (Figure 3A). Analysis of all patients recruited days 3–6 after fever onset revealed a sensitivity of 0.96 and a specificity of 0.79 when a chymase level of 1.5 ng/mL was used as a cutoff for DHF (Figure 3D). This cutoff was chosen on the basis of the ROC analysis, to emphasize sensitivity over specificity by choosing the maximal sensitivity value that retained good specificity. This resulted in a positive predictive value of 76%, indicating that 76% of those with chymase levels of >1.5ng/mL should develop DHF, and a negative predictive value of 96%.

Figure 2. Chymase level is a prognostic biomarker of dengue hemorrhagic fever (DHF). A, Patients with DHF (n = 99) had significantly higher levels of serum chymase as compared to patients with DF (n = 223) and the subset of patients with DF who had bleeding (DFWB; n = 15). Patients with DFWB had elevated chymase levels as compared to all patients with DF that were still significantly lower than those in patients with a diagnosis of DHF. B, Chymase levels were significantly higher in patients with DHF in all age groups. Data are for 18 patients with DF and 10 with DHF who were aged 0–5 years; 20 and 18, respectively, aged 6–9 years; 52 and 16, respectively, aged 10–18 years; 130 and 48, respectively, aged 18–45 years; and 27 and 7, respectively, aged >45 years. C, Chymase levels were significantly elevated in patients with DHF, compared with levels among patients with DF, enrolled on days 3–6 after fever onset. Data are for 43 patients with DF and 15 with DHF on day 3; 78 and 36, respectively, on day 4; 63 and 23, respectively, on day 5; and 30 and 14, respectively, on day 6. D, Platelet counts in acute blood samples collected on the day of study enrollment did not differ between DF and DHF groups (analysis was performed using the Student unpaired t test). Error bars represent standard errors of the mean. *P<.05, by 1-way analysis of variance with the Bonferroni post hoc test.
value of 97%, indicating that 97% of those with chymase levels of <1.5 ng/mL should not develop DHF. For patients aged 0–5 years, a sensitivity of 1 is maintained at this cutoff value. For patients aged >45 years, the sensitivity and specificity were 1.0 and 0.92, respectively (Figure 3). These results confirm that an elevated chymase level can identify DHF and DSS with very high sensitivity and good specificity for multiple age groups.

**PECs Influence Serum Chymase Levels**

We assessed whether any PECs could influence chymase levels in DENV-infected patients. PECs were self-reported and included obesity, diabetes, and cardiac, musculoskeletal, cerebrovascular, liver, or lung-associated diseases. The patients’ history of anemia or gastrointestinal disease had been recorded, but none in this cohort had these conditions. The impact of PECs on chymase levels was assessed by multivariate analysis with age in years as a covariate. This revealed that chymase levels during acute dengue are significantly influenced by certain PECs, such as diabetes and cardiovascular diseases (Table 2).

We next questioned whether PECs could interfere with the capacity of the chymase level to predict DHF and DSS. The average chymase levels in the DF and DHF groups between adult patients (age, >18 years) with and those without PECs (no-PEC) were compared (Figure 4). The analysis was limited to adults since nearly all patients with PECs were adults. For all groups of patients with PECs, chymase levels were significantly higher in patients who received a DHF diagnosis as compared to patients who received a DF diagnosis (Figure 4), emphasizing that none of the PECs would prevent the use of chymase level as a biomarker of DHF. For patients with any PEC (Figure 4A) and the subset of patients with lung-associated diseases or obesity (Figure 4B and 4C), patients with DHF had significantly higher levels of chymase than no-PEC DHF patients. For patients with cardiac diseases, diabetes, or musculoskeletal diseases as PECs...
Although the mean levels of chymase were higher, the difference between patients with and those without PECs was not statistically significant. Similarly, there was a nonsignificant trend toward increased chymase levels in patients with DF for certain PECs, such as lung-associated diseases (Figure 4B). Together, these results emphasize that the chymase concentration remains a good indicator of DHF in DENV-infected patients and that it is higher in patients with certain types of PECs, particularly obesity and lung-associated conditions.

**DISCUSSION**

Our results highlight the potential of the chymase level to be used to predict DHF at the time of initial presentation. Importantly, at the time of blood specimen collection, only 4% of patients who would ultimately receive a diagnosis of DHF had received a DHF diagnosis; yet chymase concentrations would have correctly predicted that 96% of those patients would develop DHF. Indeed, for all groups of patients recruited in this cohort, independent of age, chymase levels were correlated with the severity of disease and significantly predicted disease outcomes according to multiple statistical methods. Interestingly, patients with DFWB had significantly higher levels of chymase than patients with DF without bleeding, indicating that the chymase level may assist in discriminating within the broad clinical spectrum of DF. Other early clinical signs, such as mild bleeding, rash, and low platelet count, were not able to prognosticate the subsequent development of DHF. Signs of bleeding or hemorrhaging were also not strong predictors of DHF in this cohort, most likely because they frequently were present in patients in the DFWB group, who did not meet all of the criteria for DHF. This emphasizes the long-standing observation that patients with DF often experience mild-to-moderate hemorrhagic manifestations that have limited predictive capacity for diagnosing DHF. Some warning signs were associated with DHF diagnosis, but these occurred late in the development of disease, during the transition from acute febrile to defervescent stages, and they occurred after an increase in the serum chymase level. One limitation of our study is that more clinical data were available for patients who were admitted to the hospital for a longer time and that inpatient monitoring was not used for the purposes of collecting data for the study. Therefore, some patients could have developed signs or symptoms that were not recorded because they had been discharged. The association in our study between certain warning signs and DHF could reflect their importance to the accepted case definition of severe DENV. Reliance only on these clinical signs for DHF diagnosis is not ideal owing to their late presentation. Since chymase levels were elevated in blood specimens obtained prior to the presentation of most warning signs, they can potentially also identify patients who may go on to develop severe disease and for whom closer clinical monitoring is warranted.

### Table 2. Multivariate Analysis of Preexisting Conditions (PECs) That Significantly Influence Serum Chymase Levels During Dengue Virus Infection

<table>
<thead>
<tr>
<th>PEC</th>
<th>P</th>
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<tbody>
<tr>
<td>Any</td>
<td>.289</td>
</tr>
<tr>
<td>Cardiovascular disease</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Diabetes</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Lung</td>
<td>&lt;.001</td>
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<tr>
<td>Obesity</td>
<td>&lt;.001</td>
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<tr>
<td>Cerebrovascular disease</td>
<td>&lt;.001</td>
</tr>
<tr>
<td>Musculoskeletal disease</td>
<td>.999</td>
</tr>
<tr>
<td>Liver disease</td>
<td>&lt;.001</td>
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</table>

Data were analyzed with chymase concentration as a dependent variable and PECs as independent variables.

(Supplementary Figure 5A–C), although the mean levels of chymase were higher, the difference between patients with and those without PECs was not statistically significant. Similarly, there was a nonsignificant trend toward increased chymase levels in patients with DF for certain PECs, such as lung-associated diseases (Figure 4B). Together, these results emphasize that the chymase concentration remains a good indicator of DHF in DENV-infected patients with PECs and that it is higher in patients with certain types of PECs, particularly obesity and lung-associated conditions.

**Figure 4.** Preexisting conditions (PECs) lead to higher chymase levels in patients with dengue hemorrhagic fever (DHF). Comparison of serum chymase levels in healthy patients (for no-PEC, DF: n = 191, for no-PEC, DHF: n = 82) versus patients with PECs grouped by all PECs (for PEC, DF: n = 33, for PEC, DHF: n = 17, A), patients with lung-associated diseases (with-PEC; n = 6-DF, n = 2-DHF, B), and obesity (with-PEC; n = 5-DF, n = 5-DHF, C). For all groups, with-PEC groups were compared to the without-PEC adult patients (age, >18 years). Data were analyzed by 2-way analysis of variance with the Sidak multiple comparisons test. Post-hoc power analyses are included in Supplementary Table 3. *P < .001.
Our data indicate that the chymase level is a stronger biomarker for children and adolescents than for adults. A lower chymase level in adults with DHF could reflect their increased likelihood of having other comorbidities that could exacerbate disease or influence the diagnosis of severity, potentially independent of vascular complications. Elderly patients frequently have atypical dengue presentations and often experience coinfections [36]. Older adults may also present later with a febrile response after inoculation with virus, which could alter the detection parameters and the study time points since they are anchored here to the onset of fever. Studies have not identified any influence of age on the DENV intrinsic incubation period; however, these studies were performed in young adults [37].

Even though the oldest adults in this study had lower levels of chymase during DHF, a conservative cutoff of 1.5 ng/mL was still able to predict DHF in this population with a sensitivity approaching 1 and a specificity of >0.95. Further studies are needed to determine whether these cutoffs are valid for additional patient cohorts, including those in other dengue-endemic regions.

Previously, we reported that, in a cohort of adult Singaporean patients, chymase levels were significantly correlated with disease severity. Those data had a sensitivity of 1.0, a false-positive rate of 0.83, and an area under the ROC curve of 0.99 for adults recruited up to 72 hours after fever onset (Supplementary Figure 4) [25]. In that study, measurement of the chymase concentration in sequential serum samples showed that levels decreased over time as acute disease resolved [25]. In this cohort, patients were recruited at longer times after fever onset, but we did not observe significant differences between the levels of chymase, based on the day of study enrollment. Longitudinal serum monitoring of individual patients, which was not performed here, might also demonstrate that chymase levels are higher at earlier time points. Going forward, it will be important to assess whether early measurement of the chymase level and intervention can improve the outcomes of DENV-infected patients and promote allocation of healthcare resources to those most likely to develop severe DENV.

Our results show that PECs influence the levels of chymase during DHF. In particular, the link between high chymase levels and DHF in patients with lung-associated PECs is interesting because, although the specific conditions were not specified, some of these patients presumably had asthma, which has a known MC-promoted etiology. In the 1981 Cuban epidemic, asthma was also frequently a PEC of patients with DHF [38]. Higher chymase levels indicate increased activation of MCs in patients with DHF with PECs. Although we previously observed that febrile control patients without dengue (most of whom had respiratory virus infections) did not have elevated chymase levels [25], some other infections activate MCs. One patient with DHF in this study had a chymase concentration that was approximately 100 times the levels of other patients with DHF and received a diagnosis of bacterial coinfection. This raises the point that chymase induction is not necessarily DENV specific and can be detected in other conditions that promote MC activation, including bacterial infections and noninfectious inflammatory conditions, such as asthma or anaphylaxis. Thus, measurement of the serum chymase level alone should not serve as a diagnostic test for DENV and, for DENV infection prognosis, must be interpreted in the context of results of other tests performed to confirm infection or in the context of a strong suspicion of DENV. Furthermore, since PECs influence chymase concentrations in DENV-infected patients, additional studies should determine whether differing cutoffs for patients with PECs would improve the prognostic value of the chymase level.

Our data demonstrate that the chymase concentration is a strong predictor of DHF in febrile patients with acute-phase DENV infection, with the potential to identify those at risk for complications more robustly and at earlier time points than many clinical warning signs. Since chymase is an angiotensin-converting enzyme and can influence vasoconstriction [28], further studies are needed to determine its contributions to the pathophysiology of DENV disease.

Supplementary Data
Supplementary materials are available at The Journal of Infectious Diseases online. Consisting of data provided by the authors to benefit the reader, the posted materials are not copyedited and are the sole responsibility of the authors, so questions or comments should be addressed to the corresponding author.

Notes
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