Predictive Symptoms and Signs of Severe Dengue Disease for Patients with Dengue Fever: A Meta-Analysis

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The aim of the meta-analysis was to provide more solid evidence for the reliability of the new classification. A systematic literature search was performed using PubMed, Armed Forces Pest Management Board Literature Retrieval System, and Google Scholar up to August 2012. A pooled odds ratio (OR) was calculated using either a random-effect or a fixed-effect model. A total of 16 papers were identified. Among the 11 factors studied, five symptoms demonstrated an increased risk for SDD, including bleeding [OR: 13.617; 95% confidence interval (CI): 3.281, 56.508], vomiting/nausea (OR: 1.692; 95% CI: 1.256, 2.280), abdominal pain (OR: 2.278; 95% CI: 1.631, 3.182), skin rashes (OR: 2.031; 95% CI: 1.269, 3.250), and hepatomegaly (OR: 4.751; 95% CI: 1.769, 12.570). Among the four bleeding-related symptoms including hematemesis, melena, gum bleeding, and epistaxis, only hematemesis (OR: 6.174; 95% CI: 2.66, 14.334; \( P < 0.001 \)) and melena (OR: 10.351; 95% CI: 3.065, 34.956; \( P < 0.001 \)) were significantly associated with SDD. No significant associations with SDD were found for gender, lethargy, retroorbital pain, diarrhea, or tourniquet test, whereas headache appeared protective (OR: 0.555; 95% CI: 0.455, 0.676). The meta-analysis suggests that bleeding (hematemesis/melena), vomiting/nausea, abdominal pain, skin rashes, and hepatomegaly may predict the development of SDD in patients with DF, while headache may predict otherwise.

1. Introduction

Dengue is an infectious disease caused by dengue virus (DENV). It is endemic in many tropical and subtropical areas. Patients infected with DENV have a wide spectrum of clinical manifestation, ranging from silent infections with no symptoms to a mild flu-like syndrome, dengue fever (DF), or severe dengue disease (SDD), including dengue haemorrhagic fever (DHF) and dengue shock syndrome (DSS) [1–3]. Recently, DF has become one of the most challenging public health problems in affected regions, as the DF incidence increases rapidly worldwide [4]. There are approximately 2.5 billion people at risk for DF worldwide. Fifty million people would acquire DENV annually, and half a million among them would develop dengue hemorrhagic fever, including 22,000 deaths [5].

Several methods have been used for the diagnoses of DF. However, there lacks an accurate means to predict the severity of disease at early stages of the infection. Since patients with mild or classical DF can develop SDD later [2], it is important to look for symptoms/signs to facilitate the early prediction of the progression into SDD. The establishment of predictive symptoms/signs is essential for preventing unnecessary hospitalization, reducing disease burden, and controlling potential SDD. Based on the dengue guidelines (2009), the warning symptoms/signs for SDD include abdominal pain or tenderness, persistent vomiting, mucosal bleed, lethargy, and restlessness.

Published studies about symptoms/signs that are associated with SDD have been inconclusive. For instance, Khan et al. found that male DF patients were more likely to progress into DHF (OR: 2.3, 95% CI: 1.1–4.5, \( P \) value 0.021) [6], while
there was no association with SDD [7, 8]. The frequencies of symptoms/signs of vomiting/nausea, abdominal pain, skin rashes, and bleeding were also found to be correlated with SDD [6–11]. However, the published studies were not able to conclude that these symptoms/signs are associated with SDD. In addition, although some findings such as viral factors, varying host immune conditions, host immune reactions, and laboratory tests can predict SDD [12], the clinical manifestations might always offer the earliest markers in predicting SDD. For example, patients with nonsevere dengue could be clustered into two groups: one with warning signs, such as abdominal pain, mucosal bleeding, and liver enlargement, and the other without those signs [2], as most of the warning signs were associated with an indication for ICU admission and were severe, even for the relationship of death [13]. Because of these inconsistent reports, more accurate methods to predict SDD are needed. We conducted the meta-analysis to identify which clinical symptoms/signs are associated with SDD and to help find better methods to predict the development of SDD in patients with DF.

2. Materials and Methods

2.1. Literature Searches. Our study was performed according to the recommendations of the PRISMA Statement [14], which is available in supporting information (see Table S1 available online at http://dx.doi.org/10.1155/2014/359308). Computerized searches were conducted on NCBI PubMed, Armed Forces Pest Management Board Literature Retrieval System, and Google Scholar. As few studies before 2000 met the criteria of WHO guidelines (1997), the search time window was set between January 1, 2000, and August 1, 2012, with no language limit. Because severe dengue disease (SDD) is classified as DHF and DSS, we used the following key words for searching: dengue fever, DF, dengue haemorrhagic fever, DHF, dengue shock syndrome, DSS, and clinical diagnosis. We also manually searched the reference lists of the retrieved articles to identify more qualified studies.

2.2. Inclusion and Exclusion Criteria. Studies were eligible for inclusion if they met the following criteria: (1) retrospective, prospective, or cross-sectional studies providing the details of symptoms/signs as well as any information regarding gender, vomiting/nausea, abdominal pain, skin rashes, bleeding, headache, lethargy, retroorbital pain, diarrhea, hepatomegaly, or tourniquet test; (2) the symptoms/signs of DF and SDD were distinguished; (3) cases with DF in the study were confirmed by laboratory tests; cases with SDD were defined by one or more of the following: plasma leakage that may lead to shock (dengue shock) and/or fluid accumulation, with or without respiratory distress, and/or severe bleeding, and/or severe organ impairment. When two or more publications reported the same study, we chose the most recent one. Reports providing inadequate information were excluded.

2.3. Quality Assessment. The quality of the selected studies was assessed independently by two authors using the Newcastle-Ottawa Scale (NOS) [15]. The NOS uses different tools for case-control and cohort studies and consists of 3 parameters of quality: selection, comparability, and exposure/outcome assessment. The NOS assigns a maximum of 4 points for selection, 2 for comparability, and 3 for exposure or outcome. We assigned NOS scores of 1–3, 4–6, and 7–9 for low, intermediate, and high-quality studies, respectively. Discrepancies were settled by consensus after joint reevaluation of the original studies.

2.4. Data Extraction. For each eligible manuscript, the following information was extracted: (1) first author’s name and year of publication; (2) study design (prospective, retrospective, or cross-sectional); (3) study populations (children, adults, or both); (4) distinctive numbers of patients with specific symptoms in DF and SDD groups.

2.5. Statistical Analysis. The prevalence rates of specific symptoms/signs in DF and SDD groups were compared by calculating an odds ratio (OR) with a 95% confidence interval (CI) using either a fixed-effect model or a random-effect model. Predictive factors of interest included gender, vomiting/nausea, abdominal pain, skin rashes, bleeding (hematemesis, melena, gum bleeding, and epistaxis), headache, lethargy, retroorbital pain, diarrhea, hepatomegaly, and tourniquet test.

Heterogeneity between studies was assessed using both the Chi-square test with a $P \leq 0.10$ and the inconsistency index ($I^2$) with a cut-off of 50% [16]. To explore the potential sources of heterogeneity among studies, subgroup analyses and metaregression were performed on the strata of study design, study population, and publication year.

Potential publication bias was comprehensively assessed by Begg’s funnel plot and Egger’s rank correlation test of asymmetry. Publication bias was determined present when the $P$ value $\leq 0.10$ by Egger’s or Begg’s test. All statistical analyses were performed using STATA version 11.0 (STATA Corporation, College Station, TX, USA).

3. Results

3.1. Study Characteristics and Quality. The search strategy identified 446 citations. Sixteen articles published between 2000 and 2012 were ultimately included in this meta-analysis based on the inclusion and exclusion criteria (Figure 1). The final collection consists of 10 prospective [9–11, 17–23], four retrospective [7, 8, 24, 25], and two cross-sectional studies [26, 27]. As listed in Table 1, eight of the 16 studies reported on a population study of children, two on adults and six on both. The factor of gender was included in five studies. The clinical symptoms/signs of vomiting/nausea were included in 13 studies, abdominal pain in 13, skin rashes in 10, bleeding in 13, headache in 13, lethargy in 6, retroorbital pain in 9, diarrhea in 7, hepatomegaly in 8, and tourniquet test in 4 studies. Four common kinds of bleeding symptoms were present in these studies, including hematemesis in five, melena in four, gum bleeding in seven, and epistaxis in five. Based on the NOS scores, 12 studies (75%) were of high quality and the other four (25%) were acceptable.
| Author (publication year) | Study design | Population | Gender | Vomiting/nausea | Bleeding | Headache | Abdominal pain | Retroorbital pain | Rashes | Diarrhea | Hepatomegaly | Lethargy | Tourniquet test |
|--------------------------|--------------|------------|--------|----------------|----------|----------|----------------|-----------------|--------|----------|-------------|----------|----------------|
| Ahmed et al. (2001) [17]| Prospective  | Children   | DF     | SD (M/F)       | DF (Y/N) | DF (Y/N) | DF (Y/N)       | DF (Y/N)        | DF (Y/N) | DF (Y/N) | DF (Y/N)   | DF (Y/N) | DF (Y/N)       |
| Narayanan et al. (2002) [21]| Prospective | Children   | 22/21  | 9/7            | 34/9     | 15/1     | 23/20         | 16/0            | 13/30   | 4/12     | 10/33       | 4/12     | 4/12           |
| Endy et al. (2002) [19]| Prospective  | Children   | —      | 4/48          | DF (Y/N) | DF (Y/N) | DF (Y/N)       | DF (Y/N)        | DF (Y/N) | DF (Y/N) | DF (Y/N)   | DF (Y/N) | DF (Y/N)       |
| Phong et al. (2004) [22]| Prospective  | Children   | 170/134| 254/85        | 56/256   | 36/283   | 140/167       | 88/230          | 156/156 | 234/84   | DF (Y/N)   | DF (Y/N) | DF (Y/N)       |
| Carlos et al. (2005) [18]| Prospective  | Children   | 143/96 | 72/48         | DF (Y/N) | DF (Y/N) | DF (Y/N)       | DF (Y/N)        | DF (Y/N) | DF (Y/N) | DF (Y/N)   | DF (Y/N) | DF (Y/N)       |
| Shah et al. (2006) [9] | Prospective  | Children   | —      | 11/0          | DF (Y/N) | DF (Y/N) | DF (Y/N)       | DF (Y/N)        | DF (Y/N) | DF (Y/N) | DF (Y/N)   | DF (Y/N) | DF (Y/N)       |
| Malavog et al. (2006) [23]| Prospective | Adult      | —      | 18/15         | DF (Y/N) | DF (Y/N) | DF (Y/N)       | DF (Y/N)        | DF (Y/N) | DF (Y/N) | DF (Y/N)   | DF (Y/N) | DF (Y/N)       |
| Lee et al. (2006) [24]| Retrospective| Both       | 177/235| 119/113       | DF (Y/N) | DF (Y/N) | DF (Y/N)       | DF (Y/N)        | DF (Y/N) | DF (Y/N) | DF (Y/N)   | DF (Y/N) | DF (Y/N)       |
| Raz et al. (2009) [27]| Cross-sectional| Both      | —      | 13/35         | DF (Y/N) | DF (Y/N) | DF (Y/N)       | DF (Y/N)        | DF (Y/N) | DF (Y/N) | DF (Y/N)   | DF (Y/N) | DF (Y/N)       |
| Lee et al. (2009) [25]| Retrospective| Adult      | 1183/672| 53/29         | DF (Y/N) | DF (Y/N) | DF (Y/N)       | DF (Y/N)        | DF (Y/N) | DF (Y/N) | DF (Y/N)   | DF (Y/N) | DF (Y/N)       |
| Priyadarshini et al. (2010) [10]| Prospective | Both      | —      | 9/60          | DF (Y/N) | DF (Y/N) | DF (Y/N)       | DF (Y/N)        | DF (Y/N) | DF (Y/N) | DF (Y/N)   | DF (Y/N) | DF (Y/N)       |
| Khan et al. (2010) [8] | Retrospective| Both       | 99/62  | 30/10         | DF (Y/N) | DF (Y/N) | DF (Y/N)       | DF (Y/N)        | DF (Y/N) | DF (Y/N) | DF (Y/N)   | DF (Y/N) | DF (Y/N)       |
| Giraldo et al. (2011) [7]| Retrospective| Children   | 77/74  | 16/14         | DF (Y/N) | DF (Y/N) | DF (Y/N)       | DF (Y/N)        | DF (Y/N) | DF (Y/N) | DF (Y/N)   | DF (Y/N) | DF (Y/N)       |
| Falconer et al. (2012) [20]| Prospective | Both       | —      | 13/15         | DF (Y/N) | DF (Y/N) | DF (Y/N)       | DF (Y/N)        | DF (Y/N) | DF (Y/N) | DF (Y/N)   | DF (Y/N) | DF (Y/N)       |
| Kardi et al. (2012) [26]| Cross-sectional| Both      | —      | 13/83         | DF (Y/N) | DF (Y/N) | DF (Y/N)       | DF (Y/N)        | DF (Y/N) | DF (Y/N) | DF (Y/N)   | DF (Y/N) | DF (Y/N)       |
| Sirvichayakul et al. (2012) [11]| Prospective | Children   | —      | 95/28         | DF (Y/N) | DF (Y/N) | DF (Y/N)       | DF (Y/N)        | DF (Y/N) | DF (Y/N) | DF (Y/N)   | DF (Y/N) | DF (Y/N)       |

Note: "M": male; "F": female; "Y": yes, the group has the symptom; "N": no, the group has no symptom. "—": no statistics.
Table 2: Results of meta-analysis for the clinical manifestations between DF and SDD.

| Clinical manifestation | Number of studies | Odds ratio (95% CI) | Test for OR | Test of heterogeneity | Publication bias |
|------------------------|-------------------|---------------------|-------------|------------------------|-----------------|
|                         |                   |                     | Test of OR | Test of heterogeneity | Publication bias |
| Gender                 | 6                 | 1.230 (0.999, 1.513) | 0.051       | 0                      | 0.686           | 0.991 | 0.707 |
| Vomiting/nausea*       | 13                | 1.692 (1.256, 2.280) | 0.001       | 39.7                   | 0.069           | 0.455 | 0.428 |
| Abdominal pain*        | 13                | 2.278 (1.631, 3.182) | <0.001      | 55.5                   | 0.008           | 0.343 | 0.669 |
| Skin rashes*           | 10                | 2.031 (1.269, 3.250) | 0.003       | 69.7                   | <0.001          | 0.581 | 0.592 |
| Bleeding*              | 12                | 13.617 (3.281, 56.508) | 0.001       | 94.0                   | <0.001          | 0.033 | 0.373 |
| Hematemeses*           | 5                 | 6.174 (2.66, 14.334)  | <0.001      | 0                      | 0.476           | 0.137 | 0.211 |
| Melena*                | 4                 | 10.351 (3.065, 34.956) | <0.001      | 0                      | 0.955           | 0.36  | 0.734 |
| Gum bleeding           | 7                 | 2.518 (0.463, 13.685) | 0.285       | 90.5                   | <0.001          | 0.058 | 0.548 |
| Epistaxis              | 5                 | 2.319 (0.599, 8.976)  | 0.562       | 86.9                   | <0.001          | 0.981 | 0.462 |
| Headache*              | 13                | 0.555 (0.455, 0.676)  | <0.001      | 0                      | 0.594           | 0.177 | 0.583 |
| Lethargy               | 6                 | 1.552 (0.714, 3.370)  | 0.267       | 49.6                   | 0.078           | 0.864 | 1     |
| Retroorbital pain      | 9                 | 1.096 (0.531, 2.261)  | 0.804       | 45.2                   | 0.067           | 0.810 | 0.602 |
| Diarrhea               | 7                 | 1.149 (0.555, 2.380)  | 0.708       | 64.1                   | 0.010           | 0.185 | 0.230 |
| Hepatomegaly*          | 8                 | 4.751 (1.769, 12.570) | 0.002       | 72.5                   | 0.001           | 0.014 | 0.063 |
| Tourniquet test        | 4                 | 2.194 (0.395, 12.206) | 0.369       | 82.4                   | 0.001           | 0.715 | 1     |

Note: "Italics": using the fixed-effect model, the other symptoms/signs: using the random-effect model; * significantly different between DF and SDD.

3.2. Potential Predictive Indicators of SDD. In this meta-analysis, the fix-effect model was analyzed in gender, hematemesis, melena, and headache, while the random-effect model was used in vomiting/nausea, abdominal pain, skin rashes, bleeding (gum bleeding and epistaxis), lethargy, retroorbital pain, diarrhea, hepatomegaly, and tourniquet. According to the meta-analysis results, there were no significant differences between the DF and SDD groups in association with the following factors ($P > 0.05$, Table 2): gender of patients, clinical symptoms/signs of lethargy, retroorbital, diarrhea, and tourniquet test. There were significant differences in association between the two groups with the following factors: symptoms/signs of vomiting/nausea, abdominal pain, skin rashes, bleeding, and hepatomegaly ($P < 0.05$, Figures 2(a)–2(e)). In particular, bleeding and hepatomegaly were highly correlated with the progression of DF into SDD, with the ORs at 13.617 (95% CI: 3.281, 56.508) and 4.751 (95% CI: 1.769, 12.570). The results indicate that these two factors strongly predict greater risk of the development of SDD. The other factors such as vomiting/nausea, abdominal pain, and skin rashes were also associated with SDD, while the strength of association was not as strong. In the bleeding symptoms, the ORs for predicting SDD of hematemesis and melena were 6.174 (95% CI: 2.66, 14.334; $P < 0.001$) and 10.351 (95% CI: 3.065, 34.956; $P < 0.001$), respectively, demonstrating significant differences between the DF and SDD groups, while the frequencies of the other two kinds of bleeding, gum bleeding and epistaxis, were not significantly different between the two groups. The details were shown in Table 2, Figures 2 and 3. Interestingly, headache was not associated with the low risk of SDD (OR: 0.555; 95% CI: 0.455, 0.676; Figure 2(f)).

3.3. Heterogeneity Analysis. Metaregression analysis was conducted to examine which factors could have brought heterogeneity across the studies, and 11 clinical symptoms/signs were analyzed, including vomiting/nausea, abdominal pain, skin rashes, bleeding (epistaxis and gum bleeding), lethargy, retroorbital pain, diarrhea, hepatomegaly, and tourniquet test. It turned out that two factors, study design and population, contributed to the heterogeneity in the studies of gum bleeding ($P < 0.10$) and epistaxis ($P < 0.10$). Based on the subgroup analyses, the epistaxis ratio was not significantly different between DF and SDD groups in children ($P = 0.562$), while the difference was significant in adults ($P < 0.001$, OR: 14.139, 95% CI: 6.622, 30.187). Similar results applied to the subgroup analysis of gum bleeding based on the retrospective and prospective studies (data not shown).
| Study ID | OR (95% CI) | Weight (%) |
|----------|-------------|------------|
| Ahmed et al. (2001) | 0.67 (0.16, 2.76) | 3.75 |
| Narayanan et al. (2002) | 3.97 (0.46, 34.20) | 1.77 |
| Endy et al. (2002) | 5.66 (1.92, 16.73) | 5.74 |
| Phuong et al. (2004) | 2.07 (1.48, 2.90) | 18.28 |
| Shah et al. (2006) | 1.52 (0.07, 33.71) | 0.89 |
| Malavige et al. (2006) | 1.77 (0.76, 4.10) | 8.24 |
| Riaz et al. (2009) | 0.83 (0.47, 1.48) | 12.58 |
| Lee et al. (2009) | 1.13 (0.71, 1.80) | 15.10 |
| Priyadarshini et al. (2010) | 1.52 (0.73, 3.16) | 9.77 |
| Khan et al. (2010) | 1.52 (0.07, 33.71) | 9.77 |
| Giraldo et al. (2011) | 4.62 (0.46, 46.67) | 1.55 |
| Falconar et al. (2012) | 4.72 (1.06, 20.91) | 3.43 |
| Sirivichayakul et al. (2012) | 1.69 (1.26, 2.28) | 100.00 |

Note: weights are from random-effects analysis

(a) Vomiting/nausea

| Study ID | OR (95% CI) | Weight (%) |
|----------|-------------|------------|
| Narayanan et al. (2002) | 1.10 (0.29, 4.18) | 4.62 |
| Endy et al. (2002) | 2.56 (0.87, 7.53) | 6.22 |
| Phuong et al. (2004) | 2.79 (2.00, 3.89) | 14.99 |
| Carlos et al. (2005) | 1.83 (1.15, 2.89) | 13.23 |
| Shah et al. (2006) | 9.67 (1.21, 77.12) | 2.26 |
| Malavige et al. (2006) | 2.30 (0.61, 8.60) | 4.69 |
| Riaz et al. (2009) | 1.26 (0.77, 2.04) | 12.87 |
| Lee et al. (2009) | 1.31 (0.73, 2.36) | 11.37 |
| Priyadarshini et al. (2010) | 4.48 (2.36, 8.50) | 10.70 |
| Khan et al. (2010) | 1.33 (0.15, 12.24) | 2.01 |
| Giraldo et al. (2011) | 2.57 (1.08, 6.14) | 8.03 |
| Falconar et al. (2012) | 627.00 (11.20, 35092.88) | 0.66 |
| Sirivichayakul et al. (2012) | 3.01 (1.30, 6.98) | 8.34 |
| Overall ($I^2 = 55.5\%, \ P = 0.008$) | 2.28 (1.63, 3.18) | 100.00 |

Note: weights are from random-effects analysis

(b) Abdominal pain

| Study ID | OR (95% CI) | Weight (%) |
|----------|-------------|------------|
| Ahmed et al. (2001) | 8.36 (2.20, 31.78) | 7.09 |
| Narayanan et al. (2002) | 0.65 (0.07, 6.30) | 3.40 |
| Endy et al. (2002) | 1.16 (0.13, 10.18) | 3.65 |
| Riaz et al. (2009) | 1.07 (0.64, 1.77) | 13.85 |
| Lee et al. (2009) | 1.62 (1.03, 2.54) | 14.36 |
| Priyadarshini et al. (2010) | 2.98 (1.57, 5.65) | 12.67 |
| Khan et al. (2010) | 9.16 (4.04, 20.78) | 11.01 |
| Giraldo et al. (2011) | 1.60 (0.73, 3.51) | 11.31 |
| Sirivichayakul et al. (2012) | 1.49 (0.69, 3.18) | 11.53 |
| Karoli et al. (2012) | 1.08 (0.48, 2.41) | 11.13 |
| Overall ($I^2 = 69.7\%, \ P = 0.000$) | 2.03 (1.27, 3.25) | 100.00 |

Note: weights are from random-effects analysis

(c) Skin rashes

Figure 2: Continued.
| Study ID            | OR (95% CI)     | Weight (%) |
|---------------------|-----------------|------------|
| Ahmed et al. (2001) | 59.18 (3.28, 1067.11) | 7.03       |
| Narayanan et al. (2002) | 28.79 (1.62, 510.35)   | 7.06       |
| Endy et al. (2002)  | 3.61 (0.31, 41.87)   | 7.66       |
| Phuong et al. (2004) | 0.58 (0.37, 0.91)    | 9.82       |
| Carlos et al. (2005) | 1.15 (0.41, 3.19)    | 9.43       |
| Shah et al. (2006)  | 73.28 (4.14, 1295.76) | 7.06       |
| Malavige et al. (2006) | 5.45 (1.90, 15.64)  | 9.40       |
| Riaz et al. (2009)  | 120.89 (46.84, 312.02) | 9.50       |
| Khan et al. (2010)  | 68.11 (23.34, 198.72) | 9.39       |
| Girald et al. (2011) | 1.72 (0.78, 3.79)    | 9.62       |
| Falconar et al. (2012) | 23.22 (1.20, 451.11) | 6.93       |
| Karoli et al. (2012) | 525.74 (30.51, 9059.21) | 7.10       |
| Overall ($I^2 = 94.0\%, P = 0.000$) | 13.62 (3.28, 56.51) | 100.00     |

Note: weights are from random-effects analysis

Figure 2: Forrest plots of the relationship between DF and the risk of SDD. (a)–(e) Pooled ORs of SDD are greater than one in 5 symptoms and signs with vomiting/nausea, abdominal pain, skin rashes, bleeding, and hepatomegaly; (f) pooled OR of SDD is smaller than one in headache.
Study ID | OR (95% CI) | Weight (%)  
--- | --- | ---  
Ahmed et al. (2001) | 3.07 (0.98, 9.60) | 67.24  
Carlos et al. (2005) | 4.17 (0.37, 46.50) | 11.80  
Shah et al. (2006) | 44.87 (2.56, 787.30) | 5.49  
Malavige et al. (2006) | 6.27 (0.34, 114.55) | 11.60  
Sirivichayakul et al. (2012) | 11.06 (0.44, 277.72) | 3.87  
Overall ($I^2 = 0.0\%, P = 0.476$) | 6.17 (2.66, 14.33) | 100.00  

(a) Hematemesis

Study ID | OR (95% CI) | Weight (%)  
--- | --- | ---  
Ahmed et al. (2001) | 13.41 (2.75, 65.37) | 41.81  
Shah et al. (2006) | 5.16 (0.29, 92.10) | 27.47  
Malavige et al. (2006) | 10.74 (0.61, 188.94) | 22.70  
Sirivichayakul et al. (2012) | 11.06 (0.44, 277.72) | 8.03  
Overall ($I^2 = 0.0\%, P = 0.955$) | 10.35 (3.07, 34.96) | 100.00  

(b) Melena

Study ID | OR (95% CI) | Weight (%)  
--- | --- | ---  
Ahmed et al. (2001) | 0.99 (0.26, 3.75) | 15.63  
Carlos et al. (2005) | 1.13 (0.41, 3.13) | 16.29  
Shah et al. (2006) | 3.37 (0.19, 61.13) | 11.48  
Malavige et al. (2006) | 2.78 (0.32, 24.08) | 13.51  
Lee et al. (2009) | 30.16 (18.41, 49.39) | 17.06  
Falconar et al. (2012) | 2.08 (0.17, 25.31) | 12.58  
Sirivichayakul et al. (2012) | 0.72 (0.08, 6.34) | 13.45  
Overall ($I^2 = 90.5\%, P = 0.000$) | 2.52 (0.46, 13.68) | 100.00  
Note: weights are from random-effects analysis

(c) Gum bleeding

Study ID | OR (95% CI) | Weight (%)  
--- | --- | ---  
Ahmed et al. (2001) | 1.38 (0.32, 5.85) | 19.65  
Carlos et al. (2005) | 1.04 (0.55, 1.93) | 24.05  
Shah et al. (2006) | 1.50 (0.08, 28.89) | 11.46  
Lee et al. (2009) | 15.96 (7.00, 36.41) | 23.17  
Sirivichayakul et al. (2012) | 1.46 (0.48, 4.42) | 21.66  
Overall ($I^2 = 86.9\%, P = 0.000$) | 2.32 (0.60, 8.98) | 100.00  
Note: weights are from random-effects analysis

(d) Epistaxis

**Figure 3:** Forrest plots of four kinds of bleeding. (a)-(b) Pooled ORs of SDD are greater than one in hematemesis and melena; (c)-(d) pooled ORs of SDD are not significantly different from one in epistaxis and gum bleeding.
3.4. Publication Bias. Funnel plots showed no publication bias in the studies covering vomiting/nausea, abdominal pain, skin rashes, bleeding, or retroorbital pain (Figure S1 and Table 2). The $P$ values of Egger’s and Begg’s tests also suggested that publication bias had little impact on the results. There were three signs, bleeding, gum bleeding, and hepatomegaly showing publication bias (Egger’s test: $P = 0.041, 0.058, 0.014$).

4. Discussion

The present study is the meta-analysis to comprehensively evaluate the correlation of clinical symptoms/signs with the development of SDD in patients with DF. The results showed that a total of five symptoms/signs significantly predict dengue patients progressing into SDD: vomiting/nausea, abdominal pain, skin rashes, bleeding, and hepatomegaly. The other five factors were not associated with the disease progression, including tourniquet versus nontourniquet, female versus male patients, lethargy, retroorbital pain, and diarrhea. We found that patients with bleeding after DENV infection had approximately a 14-fold increased risk for progression into SDD (including DHF and DSS). When compared with the frequencies of leukopenia and thrombocytopenia, haemorrhagic manifestations, such as gum bleeding, epistaxis, and gastrointestinal bleeding, are less frequent, but not rare [28]. Our analysis included four kinds of bleeding: hematemesis, melena, gum bleeding, and epistaxis. Previous studies showed that the frequencies of hematemesis, melena, gum bleeding, and epistaxis were higher in SDD patients than in DF patients, but none of them were related to the risk of development of SDD in patients with these symptoms [1, 17, 18, 25]. A recent study also showed that the gastrointestinal bleeding was associated with DSS, although it is not a strong association (OR $= 1.84$) [29]. According to our meta-analysis, the two kinds of gastrointestinal bleeding that strongly predicted SDD were hematemesis (OR: 6.174; 95% CI: 2.66, 14.334; $P < 0.001$) and melena (OR: 10.351; 95% CI: 3.065, 34.956; $P < 0.001$), while the other two kinds of bleeding were not significant risk factors. The other four clinical symptoms and signs proved significant for predicting the progression into SDD are vomiting/nausea (OR: 1.692; 95% CI: 1.256, 2.280), abdominal pain (OR: 2.278; 95% CI: 1.631, 3.182), skin rashes (OR: 2.031; 95% CI: 1.269, 3.250), and hepatomegaly (OR: 4.751; 95% CI: 1.769, 12.570). Although the vomiting/nausea, abdominal pain, and skin rashes showed a weak association with SDD compared with DF patients, these warnings must be taken seriously as recent studies demonstrated that these symptoms were associated with the mortality caused by dengue [30, 31]. We found that patients with hepatomegaly after DENV infection had approximately a 5-fold increased risk of progression into SDD; however, the CI was with a wide range. The possible reason is that the rate of hepatomegaly was significantly higher in adults than the elderly [32].

The most accepted hypothesis for progression of DF is that subneutralizing levels of DENV-specific antibodies exacerbate the disease by means of an antibody-dependent enhancement of infection (ADE) [33], which induces a complicated immunopathogenesis in the host. The extent of vascular permeability is enhanced as a result of ADE [34] and patients with SDD as well as alterations of endothelial cells have been shown to experience thrombocytopenia and coagulation disorders [35]. These significant symptoms/signs, especially the bleeding (hematemesis/melena) and hepatomegaly, are manifested in patients with SDD as a result of the aforementioned alterations. In the in vivo model for ADE-induced SDD, gastrointestinal bleeding and viral RNA increased in the liver were observed [36]. Additionally, based on skin biopsies, IgM, beta 1 C-globulin, dengue antigen, and fibrinogen deposits were found to be present within or about blood vessel walls of dermal papillae or in the blood vessels [37], implying that skin rashes that appeared in DHF were caused by an immunopathologic process. So in patients with SDD, the host immune system plays a central role in triggering symptoms like bleeding, hepatomegaly, and skin rashes, which could be used to triage patients in need of intensive care.

The unassociated factors/manifestations were gender, lethargy, retroorbital pain, diarrhea, and positivity of a tourniquet test. However, the World Health Organization (WHO) has published guidelines stating that positivity of a tourniquet test may be included in the clinical case definition of dengue haemorrhagic fever [38], and an altered level of consciousness such as lethargy should be paid extra attention [2]. Although the results from this meta-analysis showed unexpected absence of associations, relaxing vigilance over the patients with these symptoms/signs is not recommended, because the results were generated from a random-effect model that tends to be overconservative.

Furthermore, headache was a protective factor against SDD after DENV infection (OR: 0.555; 95% CI: 0.455, 0.676), implying that dengue patients with headache had a lower probability to develop into SDD. The protective effect has been proved by a retrospective cohort study [25]. However, in another study, BALB/c mice were infected with different strains of DENV which were isolated from DSS or DF patients, respectively, and in the mice infected with the strain from DSS patients, DENV-1 isolates appeared to be primarily neurotropic, whereas in the cases of other strains the virus turned to mainly infect lung and liver [39]. Suggesting that high frequency of headache could occur in patients with SDD.

There are limitations in the present study. Firstly, the results will not apply to multicenter prospective studies, since the present meta-analysis only included retrospective and single-center prospective studies. These designs could not eliminate recall and selection biases. Hence, the true associations between these symptoms/signs and the development of SDD might have been distorted. Secondly, the definitions of DF and SDD within these studies may have varied, which brought uncertainty into determining cases. Lastly, some of the results were based on a random-effect model that might weaken the validity of the analysis. Nonetheless, this study explored a new approach to identify the correlations of the symptoms/signs after DENV infection with the risk of progression into SDD, which can greatly facilitate the prevention of SDD.
5. Conclusions

This meta-analysis identified clinical symptoms and signs that significantly predicted DF patients progressing into severe dengue. DF patients with vomiting/nausea, abdominal pain, skin rashes, bleeding (hematemesis/melena), and hepar-atomegaly were more likely to develop SDD, while patients with headache had a lower risk of progression into SDD. Other factors such as gender, lethargy, retroorbital pain, diarrhea, and positive tourniquet test are not associated with SDD. Further studies, especially ones with larger sample sizes and prospective, are warranted to confirm the findings.

Abbreviations

DENV: Dengue virus
DF: Dengue fever
SDD: Severe dengue disease
DHF: Dengue haemorrhagic fever
DSS: Dengue shock syndrome
NOS: The Newcastle-Ottawa Scale
ADE: Antibody-dependent enhancement of infection
WHO: The World Health Organization.

Conflict of Interests

On behalf of all authors, the corresponding author states that there is no conflict of interests.

Authors’ Contribution

All authors were involved in the study design, including developing the search strategies and project protocol. H. Zhang, H. J. Peng, X. H. Zhang, F. Y. Zhou, Z. H. Liu, and X. G. Chen were responsible for supervising the project and performing the literature search and data extraction. H. J. Peng, F. Y. Zhou, and Z. H. Liu assessed the quality of studies. H. Zhang performed data analysis and drafted the paper. X. H. Zhang was responsible for the language editorial. Y. P. Zhou and X. G. Chen revised the paper.

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