Risk factors for death from hand–foot–mouth disease: a meta-analysis

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Abstract
In recent years, outbreaks of hand–foot–mouth disease (HFMD) in China, Singapore and other Western Pacific Region, involving millions of children, have become a big threat to public health. This study aimed to quantitatively assess all qualified studies and identify the risk factors for HFMD death. A systematic search of the databases PubMed, Medline, Embase and the Cochrane Library was performed. Study heterogeneity and publication bias were estimated. Seven case–control studies involving 1641 participants (634 died and 1007 survived) were included in the meta-analysis. Human enterovirus 71 infection, male, age ≤3 years, vomiting, cyanosis, convulsion, duration of fever ≥3 days, atypical rashes and abdominal distention were not significantly related to HFMD death (P > 0.05). Lethargy (odds ratio (OR) = 6.62; 95% CI 3.61–12.14; I² = 0%; P < 0.0001), pneumoedema/pneumorrhagia (OR = 4.09; 95% CI 2.44–6.87; I² = 0%; P < 0.0001), seizures (OR = 6.85; 95% CI 2.37–19.74; I² = 0%; P = 0.0004), dyspnoea (OR = 8.24; 95% CI 2.05–33.19; I² = 83%; P = 0.003) and coma (OR = 3.76; 95% CI 1.85–7.67; I² = 0%; P = 0.0003) were significantly associated with HFMD death, which were risk factors for HFMD death.

Introduction
The hand–foot–mouth disease (HFMD) is an acute communicable disease, mostly affecting children under 5 years of age [1]. It is mainly caused by human enterovirus 71 (EV-A71) and coxsackievirus A16 (CV-A16) [2]. The main manifestations are fever, maculopapular skin vesicles or exposure to a contaminated environment [3]. HFMD is generally self-limiting, and patients with no secondary cutaneous infection mostly recover in 2 weeks.

In recent years, several widespread outbreaks of HFMD in China [4], Singapore [5] and other Western Pacific Regions [6, 7], involving millions of children, have become a big threat to public health and a substantial economic burden. In March 2008, the largest pandemic in Asia occurred in China, causing more than 20 000 cases and dozens of deaths. Then, HFMD was defined as a C-class notifiable disease in Mainland China on 2 May 2008. The reported mortality rate of HFMD was 0.03/100 000 during 2008–2017 in China [8]. Moreover, some patients with neurologic and cardiorespiratory complications, delayed manifestations and poor sanitary conditions were more likely to die [9]. Numerous studies reported that EV-A71, coma and seizures were responsible for HFMD deaths [10, 11]. However, the risk factors for HFMD deaths have still not been completely identified. Thus, this study aimed to quantitatively assess all these qualified studies and identify the risk factors for HFMD deaths.

Materials and methods

Search strategy
A systematic search of the databases PubMed, Medline, Embase, the Cochrane Library, China National Knowledge Infrastructure and Wanfang (Chinese) were performed for relevant studies published before May 2019 using the key words ‘hand foot mouth disease OR HFMD’, ‘death OR fatal OR fatality OR mortality’ and ‘risk factors’. The reference lists of all retrieved studies were screened and checked for potential additional studies. No language restrictions were applied.

Selection criteria and exclusion criteria
The inclusion criteria were as follows: (1) All patients included had HFMD according to the Guidelines on the Diagnosis and Treatment of HFMD [12]; (2) the study had a case–control design including death and survival groups; (3) the studies investigated the association between risk factors and death of HFMD; and (4) the survival data were extracted from the
Data were obtained from each eligible study independently by two reviewers. Disagreements were discussed between the reviewers until consensus was reached. The main characteristics recorded for the selected study included the first author, publication year, country, study period, sample size, age and sex. The proportion of dead patients who had developed HFMD in the presence and absence of each given risk factor was recorded.

Two reviewers independently used the Newcastle–Ottawa quality assessment scale (NOS) to assess the quality of the selected studies [13, 14]. A third reviewer was consulted if a disagreement arose. The NOS was used to score the studies on three criteria: selection, comparability and outcome. The total score ranged from 0 to 9. High-quality studies had an overall score of ≥5. The details of the quality assessment are presented in Table S1.

Statistical analysis
Odds ratio (OR) with 95% confidence intervals (CI) were used to pool the outcome data. The $I^2$ test was used to test for statistical heterogeneity. For outcomes with low heterogeneity ($I^2 < 50\%$ and $P > 0.1$), a fixed-effects model (the Mantel–Haenszel method) was used for secondary analysis; otherwise ($I^2 \geq 50\%$ or $P \leq 0.1$), a random-effects model (the DerSimonian and Laird method) was used [15]. A sensitivity analysis was further conducted in which one study was removed and the rest were analysed to evaluate whether the results were affected statistically significantly. Thus, a sensitivity analysis was performed to assess the EV-A71 infection, vomiting and convulsion and to evaluate potential sources of heterogeneity in the analyses. Publication bias was evaluated via the visual analysis of funnel plots and conducting Egger’s and Begg’s tests [16, 17]. All tests were considered statistically significant for $P$ values <0.05. Statistical analyses were performed using Review Manager 5.3 (Cochrane Collaboration, UK) and Stata version 14.0 (Stata Corporation LP, TX, USA).

Results
Study selection and description
A total of 889 studies were obtained through database searches, of which 81 were excluded due to duplication. Another 878 studies were excluded after reviewing the title and abstract information was reviewed. Four studies were excluded after assessing the full text; two studies included repeated samples and two studies lacked control groups. The final meta-analysis included seven studies [18–24]. Six studies were from China and one was from Singapore. A flowchart depicting the study selection process is shown in Figure 1. A total of 1641 patients with HFMD were enrolled in the included trials; of these, 634 died and 1007 survived. All studies were case–control studies. The main characteristics of the included trials are summarised in Table 1.

Risk factors for HFMD deaths

EV-A71
Five studies [18–20, 23, 24] reported the association between EV-A71 infection and the risk of HFMD deaths, involving 433 deaths and 674 survivals. EV-A71 infection was not associated with HFMD death (OR = 2.12; 95% CI 0.87–5.18; $I^2 = 81\%$; $P = 0.10$) (Fig. 2).

Demographic characteristics
Demographic characteristics of fatal HFMD were also analysed in this study. In seven studies [18–24], with 634 deaths and 1007 survivors, the pooled OR for male was 0.89 (95% CI 0.71–1.11; $I^2 = 37\%$; $P = 0.29$) (Fig. 3). Three studies [20, 22, 24] investigated the association between age ≤3 years and the risk of fatal HFMD, involving 183 deaths and 424 survivors. The pooled OR for age ≤3 years was 1.55 (95% CI 0.86–2.82; $I^2 = 28\%$; $P = 0.15$) (Fig. S1). Male sex and age ≤3 years were not significantly related to HFMD deaths.

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Manifestations
Six studies [18–21, 23, 24] investigated the association between vomiting and the risk of HFMD death and included a total of 529 deaths and 797 survivors. The pooled OR for vomiting was 2.22 (95% CI 0.99–4.99; $I^2 = 85\%$; $P = 0.05$) (Fig. S2). Three studies [18–20] investigated the association between cyanosis and the risk of fatal HFMD involving 433 deaths and 460 survivors. The pooled OR for cyanosis was 3.65 (95% CI 1.00–13.33; $I^2 = 91\%$; $P = 0.05$) (Fig. S3). Convulsion was analysed in four studies [18, 19, 21, 24], and 489 deaths and 535 survivors were enrolled. The pooled OR for convulsion was 0.92 (95% CI 0.31–2.74; $I^2 = 88\%$; $P = 0.88$) (Fig. S4). Two studies [18, 20] investigated the association between duration of fever ≥3 days and the risk of HFMD death and included a total of 309 deaths and 408 survivors. The pooled OR for duration of fever ≥3 days was 0.81 (95% CI 0.20–3.24; $I^2 = 87\%$; $P = 0.76$) (Fig. S5). Four studies investigated the association between atypical rashes and the risk of fatal HFMD involving 443 deaths and 591 survivors [18–20, 23]. The pooled OR for atypical rashes was 1.07 (95% CI 0.80–1.44; $I^2 = 28\%$; $P = 0.65$) (Fig. S6). Moreover, abdominal distention was analysed in two studies [18, 20], and 309 deaths and 408 survivors were enrolled. The pooled OR for abdominal distention was 1.68 (95% CI 0.74–3.85; $I^2 = 0\%$; $P = 0.22$) (Fig. S7). Thus, vomiting, cyanosis, convulsion, duration of fever ≥3 days, atypical rashes and abdominal distention were not significantly related to HFMD deaths (Table 2).

Three studies [18, 21, 24] were included in the analysis between lethargy and the risk of fatal HFMD involving 362 deaths and 483 survivors. The pooled OR for lethargy was 6.62 (95% CI 3.61–12.14; $I^2 = 0\%$; $P < 0.0001$) (Fig. 4). Furthermore, pneumoedema/pneumorrhagia was analysed in two studies [19, 20], and 160 deaths and 183 survivors were enrolled. The pooled OR for pneumoedema/pneumorrhagia was 4.09 (95% CI 2.44–6.87; $I^2 = 0\%$; $P < 0.0001$) (Fig. S8). Two studies [20, 21] investigated the association between seizures and the risk of HFMD death and included a total of 74 deaths and 254 survivors. The pooled OR for seizures was 8.24 (95% CI 2.05–33.19; $I^2 = 83\%$; $P = 0.003$) (Fig. S10). Coma was analysed in two studies [19, 20], and 160 deaths and 183 survivors were enrolled. The pooled OR for coma was 7.67 (95% CI 2.03–27.87; $I^2 = 0\%$; $P = 0.003$) (Fig. S9). Therefore, lethargy, pneumoedema/pneumorrhagia, seizures, dyspnoea and coma were found to be significantly associated with HFMD death (Table 2).
Publication bias
The Egger’s and Begg’s tests provided no evidence of publication bias for males ($P = 0.909$ and 0.368, respectively). The funnel plot appeared generally symmetrical (Fig. S12).

Sensitivity analysis
In sensitivity analyses in which one study was excluded at a time from each analysis, the summary estimates were not substantially altered for EV-A71 infection, vomiting and convulsion (Figs S13–S15).

Discussion
Several outbreaks of HFMD have been reported from regions such as Malaysia [25, 26], Singapore [27], Mainland China [28], Brunei [29], Western Australia [30], the USA [31] and Germany [32]. Although most HFMD episodes are usually mild and self-
limiting, a minority of patients may rapidly progress to severe complications and lose their lives [9]. Between 2008 and 2014, more than 10 million cases, including 3000 deaths from HFMD, were reported to the countrywide disease-reporting system in Mainland China [33]. Factors associated with severe or fatal HFMD include EV-A71, a high fever of over 39 °C for more than 3 days, a raised WBC count >10.8 × 10⁹/L, vomiting, tachycardia, lethargy, hyperglycaemia and leucocytosis [10, 34, 35]. However, the risk factors for HFMD death are not completely clear. Thus, it is significant to determine the risk factors associated with the occurrence of death for patients with HFMD. This meta-analysis was designed to explore the risk factors for HFMD death and help distinguish patients at risk of developing fatal HFMD at an early stage.

Seven case–control studies were included in this meta-analysis. In this study, the risk factors for fatalities were lethargy, pneumoedema/pneumorrhagia, seizures, dyspnoea and coma. Moreover, EV-A71, male, age, vomiting, cyanosis, convulsion, duration of fever ≥3 days, atypical rashes and abdominal distension were not associated with fatal HFMD.

Several clinical studies indicated that EV-A71-related HFMD was always associated with more severe symptoms, such as acute flaccid paralysis, brainstem encephalitis, rapid fatal pulmonary oedema and haemorrhage, whereas CA6 and CA10 caused only mild symptoms [36]. In addition, some researchers held the belief that EV-A71-infected patients faced an increased risk of mortality compared with other viruses. In this study, EV-A71 infection alone was not responsible for HFMD-related deaths. EV-A71 is a neuronophagic virus that mainly affects the brainstem, causing encephalitis, aseptic meningitis and other neurological disorders characterised by vomiting, coma, startle, frequent convulsions and light-reflex insensitivity. This inconsistency may be attributed to missing virology investigation data in the survival group.

Lethargy was reported in lots of studies as a predictor of severe HFMD and confirmed in this study. It was found that lethargy was a useful clinical symptom of the nervous system involved in early disease [37]. Several studies indicated that pneumoedema/pneumorrhagia was the major cause of death for critical and severe HFMD [38, 39]. Fulminant neurogenic pneumoedema was found in HFMD-related deaths in Malaysia [40, 41], and it may be caused by the damage to certain areas of the brainstem or an increase in pulmonary vascular pressure and pulmonary endothelial permeability [42]. This study showed that pneumoedema/pneumorrhagia was associated with the mortality of patients with HFMD. Chan et al. thought that seizures were the main symptom of neurological involvement and associated with severe disease, consistent with the results [40]. Deng et al. held that dyspnoea was not a significant risk factor for the development of fatal HFMD [24]; however, Long et al. suggested that dyspnoea significantly increased the risk of HFMD death [18], which was confirmed in the present study. Coma, a neurological
disorder, was a significant symptom of HFMD death, similar to the results of the present study [19].

The male patients were more susceptible to HFMD death than female patients [43]. In this study, the male-to-female ratio of fatal cases was 1.63:1 and no significant association was found between male sex and HFMD death. The results of this study showed that the ratio of children aged >3 years was 91.2%, which was higher than the rate reported in Vietnam [44]. Moreover, it was found that age was not a risk factor for HFMD death probably because the immune system of children reaches a stable state at about 5 years of age and low immunity makes no significant difference between fatal and non-fatal cases. The manifestations of digestive system, abdominal distention and vomiting were not the risk factors for HFMD death in this study, as confirmed by some previous studies [18]. Long et al. found that children with the symptom of cyanosis were at risk of severe HFMD [18]. However, the present study showed that cyanosis was not associated with HFMD death. Furthermore, convulsion was reported to increase the risk of death from severe HFMD; four included studies confirmed the conclusion [18, 19, 21, 24]. In the present meta-analysis, convulsion was not related to the mortality of HFMD. Although boys enjoy outdoor activities, they often pay no attention to hygiene, but it is not the key to increase the mortality of HFMD. Some previous studies regarded duration of fever \( \geq 3 \) days as a neurological manifestation and found that it increased the risk of death. However, the results indicated that the duration of fever \( \geq 3 \) days was not associated with HFMD death. The occurrence of atypical rashes may be associated with mortality [23], contrary to the results of this study. However, only seven fatal cases were included in this study, reducing the strength of the results.

This study had several inevitable limitations. First, three of seven included studies were published in Chinese and the quality of studies might differ from the ones in English. Second, most of the included studies focused on the patients in China; only one study reported the epidemiology of fatal cases in Singapore. After excluding the study from Singapore, the summary estimates were not substantially altered. Therefore, including the study from Singapore into this meta-analysis was reasonable. Third, several studies might have a selection bias. It was ideal that the

**Table 2. Meta-analysis of risk factors of HFMD death in seven separate studies**

| Risk factors                        | No. of studies | Death | Survival | OR (95% CI) | Test of heterogeneity |
|-------------------------------------|----------------|-------|----------|-------------|-----------------------|
| EV71 infection                      | 5              | 372   | 61       | 2.12 (0.87–5.18) | R 20.67 0.10 81 |
| Male                                | 7              | 393   | 241      | 0.89 (0.71–1.11) | F 9.56 0.29 37 |
| Age \( \leq 3 \) year               | 3              | 167   | 16       | 1.55 (0.86–2.82) | F 2.76 0.15 28 |
| Vomiting                            | 6              | 329   | 200      | 2.22 (0.99–4.99) | R 33.65 0.05 85 |
| Cyanosis                            | 3              | 188   | 248      | 3.65 (1.00–13.33) | R 21.11 0.05 91 |
| Convulsion                          | 4              | 177   | 312      | 0.92 (0.31–2.74) | R 24.59 0.88 88 |
| Duration of fever \( \geq 3 \) days | 2              | 225   | 84       | 0.81 (0.20–3.24) | R 7.85 0.76 87 |
| Atypical rashes                     | 4              | 187   | 256      | 1.07 (0.80–1.44) | F 4.15 0.65 28 |
| Abdominal distention                | 2              | 13    | 296      | 1.68 (0.74–3.85) | F 0.35 0.22 0 |
| Lethargy                            | 3              | 98    | 264      | 6.62 (3.61–12.14) | F 1.55 <0.0001 0 |
| Pneumonoeudema/pneumorrhagia        | 2              | 112   | 48       | 4.09 (2.44–6.87) | F 0.06 <0.0001 0 |
| Seizures                            | 2              | 10    | 64       | 6.85 (2.37–19.74) | F 0 0.0004 0 |
| Dypnoea                             | 2              | 166   | 188      | 8.24 (2.05–33.19) | R 11.94 0.003 83 |
| Coma                                | 2              | 116   | 44       | 3.76 (1.85–7.67) | F 0.06 0.0003 0 |

Model: R, random; F, fixed.

![Fig. 4. Forest plots showing the results of the meta-analysis regarding lethargy.](image)

**Table 2.** Meta-analysis of risk factors of HFMD death in seven separate studies
patients were randomly enrolled in the survival group, but not all patients had undergone aetiological examinations. Thus, only patients with aetiological examinations were enrolled in the control group. Moreover, several manifestations, such as coma and abdominal distention, were reported in two included studies. Finally, though detailed sensitivity analyses were undertaken, given the heterogeneity in the study protocols, clinically relevant differences could have been missed and might be better assessed in a meta-analysis of individual patient data. There could be additional confounders not accounted for in the analysis. Also, not all of the trials reported each of the outcomes analysed.

This study aimed to identify the manifestations of fatal HFMD, so that early treatment might be started to reduce the mortality. The results of the present study revealed that neurological manifestations, such as lethargy and seizures, increased the mortality of patients with HFMD. Screening patients with HFMD for these abnormal clinical presentations was beneficial in predicting fatal nervous system disorder, allowing timely initiation of appropriate interventions. Due to no effective antiviral therapies up to now, vaccines have become the most effective solution in preventing EV-related HFMD. Moreover, two inactivated EV-A71 vaccines were available in China, showing good efficacy and safety in HFMD prevention [45–48]. In addition, hand washing, disinfecting common areas and limiting exposure by keeping ill children out of school are also very effective prevention measures for HFMD. Therefore, multivalent EV and coxsackievirus vaccines should be developed and recommended to protect children from HFMD.

In conclusion, the results suggested that lethargy, pneumonoe-dema/pneumorrhagia, seizures, dyspnoea and coma increased with HFMD deaths. EV-A71 infection, male, vomiting, cyanosis, convulsion, duration of fever ≥ 3 days, age, atypical rashes and abdominal distention were not associated with HFMD death. It is vital to screen patients with HFMD for these abnormal clinical presentations, allowing timely initiation of appropriate interventions to reduce the mortality.

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