Hand, Foot and Mouth Disease: A Single Centre Retrospective Study of 403 New Cases and Brief Review of Relevant Indian Literature to Understand Clinical, Epidemiological, and Virological Attributes of a Long-Lasting Indian Epidemic

Abstract

Background: There have been sporadic and periodic large-scale epidemics of hand, foot, and mouth disease (HFMD) with cases at risk for significant morbidity and mortality particularly in Southeast Asia since 1997 and in India since early 2003. Method: We retrospectively studied 403 cases recorded from 2009 to 2019 and reviewed relevant Indian literature published between 2004 and 2019 to understand clinical, epidemiological, and virological attributes of this long-lasting Indian epidemic. Result: There were 96.8% children and adolescents (M:F 1.6:1) aged 2 months to 18 years and 84% were aged <5 years. Adult family contacts comprised 3.2%. Only 12 sporadic cases occurred during 2009-2011 followed by increased number from 2012 to 2015 peaking with 30.8% cases in 2013 and declining slowly until the year 2019 with small resurgence in 2018. The major peaks occurred during summers with small peaks in autumns. Literature review showed 3332 cases presenting between 2004 and 2019 across Indian states with similar epidemiological trends whereas serotyping identified Coxsackievirus A16 (CV A16) in 83%, Coxsackievirus A6 (CV A6) in 17%, Enterovirus 71 in 4.1%, and multiple strains in 11.7% samples, respectively. Conclusion: The overall features of this long-lasting HFMD epidemic; affecting children aged <5 years more often than adults, none or minimum neurological or pulmonary complications in few patients, peaks occurring during summer and autumn months, and identity of the pathogenic virus coincide with global trends. However, the continuous spread of the disease across the country appears in sync with pre-epidemic periods of China and Taiwan. It calls for a continuous surveillance and making HFMD a notifiable disease in India.

Keywords: Coxsackievirus A16, coxsackievirus A6, epidemic, HFMD, human enterovirus 71, India, onychomadesis, Southeast Asia, viral infection

Introduction

Hand, foot and mouth disease (HFMD) primarily affects infants and children or occasionally adults. The diagnosis is mainly clinical from characteristic mucocutaneous lesions distributed over hands, feet and oral cavity, and a prodrome of fever, malaise and upper respiratory symptoms. Laboratory diagnosis is from isolation and molecular identification of the virus in culture or from throat swabs, stool or vesicular fluid samples. The disease is caused by Picornaviridae family comprising more than 100 types including poliovirus, coxsackievirus (CV) A and B, echoviruses, and human enteroviruses (EVs). Its clinical course is mainly mild and self-limiting but a severe form of infection and systemic complications have been attributed to EV 71. Its transmission is via oro‑fecoal route from infected patients, or contact with contaminated material surfaces, vesicle fluid, food, or water. EV71 has been identified in throat swabs or feces of patients even weeks before symptom onset, thus, an asymptomatic person may remain infectious during incubation period and even after symptoms have resolved in a symptomatic patient. The disease occurs worldwide in epidemic or sporadic form. For about 3 decades following its discovery, only small scale outbreaks limited to small geographic areas were reported. HFMD epidemics have been usually because of CV A16 and CV A6 and A10 lead to sporadic cases. As no

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specific treatment or vaccine that is effective, safe and can be used routinely in clinical practice is available, HFMD has become a significant public health problem because of frequent outbreaks and rise in its incidence, severity, and fatal complications in Southeast Asian countries in last few decades with a gradual shift towards EV 71 infection.[7-11]

The first report of disease outbreak in India was from Calicut in 2004 that was followed by a large scale outbreak in 2007 in and around Kolkata, the capital of West Bengal.[12,13] Since then, many sporadic cases and epidemic outbreaks have been reported almost from the entire country [Figure 1] until it became sporadic in nature later. However, HFMD is not a mandatory reportable disease in India.

This study presents characteristics of 403 new cases registered in a single center between 2009 and 2019 and briefly reviews relevant Indian literature to understand overall clinical, epidemiological, and virological attributes of this long-lasting Indian epidemic in the context of world trends.

Materials and Methods

Clinical data

The medical records of all patients with HFMD attending the dermatology outpatient clinic between 2009 and 2019 were analyzed retrospectively after approval from institutional ethics committee. This tertiary care institute is located in the Western Himalayas at 2.1°N 76.27°E and an average altitude of 733 meters above sea level. The terrain is mostly semi-foothills or plains with semi-temperate to subtropical weather round the year. This hospital is visited by the patients from the whole district and adjoining areas on their own or they are referred by primary health care centers for specialized services.

The data was analyzed for clinico-epidemiological characteristics such as demographic profile, personal and family history, evolution of skin lesions and systemic features. The diagnosis of HFMD was mainly from clinical presentation of a prodrome of mild to moderate fever and other symptoms and eruption of characteristic 3-7 mm erythematous macules evolving rapidly into pale white, thin-walled oval vesicles with a red areola characteristically distributed over acral areas with or without involvement of oral cavity. These would fade over 2-3 days and heal without crusting or scarring in about 7-10 days. Virological studies were not performed because of financial constraints and for want of in house facility.

Review of literature

Online search using PubMed, MEDLINE, IndMed and Google was performed for English language medical literature, news reports, and state notifications on HFMD published between 2004 and 2019 describing case reports, case series, and epidemiological studies from India. The search terms included “hand, foot and mouth disease”, “coxackievirus” and “human enteroviruses” in combination with “India” using comma-separated value function.

Statistical analysis

The MS Office™ Excel® software was used to tabulate and analyze the data pertaining to our patients. The continuous data are presented as means and categorical variables are presented as frequencies and percentages.

The data obtained from review of published reports were tabulated and analyzed for number of the affected individuals, state/region of the outbreak, year and time of the outbreak, and available results of the viral studies.

Results

Clinical, demographic and epidemiological features of new cases

We recorded 403 new cases of HFMD between 2009 and 2019. Only 12 sporadic cases occurred between 2009 and 2011. Thereafter, number of cases increased between 2012 and 2015 peaking with 124 (30.8%) cases in 2013 followed by a slow decline until the year 2019 with small resurgence in 2018 [Figure 2]. One hundred and fifty (37.2%) cases presented during summers (May-June) with other small peaks in autumn (October-November) of 2016 and 2017 [Figure 3]. Table 1 depicts their clinicoepidemiological characteristics. There were 390 (96.8%) children comprising 241 (61.8%) males and 149 (38.2%) females (M:F 1.6:1) aged between 2 months and 18 years. The majority, 339 (84.1%) patients were aged 2 months to 5 years. Forty (9.9%) children were from a single kindergarten. Thirty six (8.9%) patients were family members of the affected cases. Thirteen (3.2%) patients were adults and included 5 (38.5%) males and 8 (61.5%) females (M:F 1:1.6) aged between 21 and 37 years. They were mostly family members of the affected children and had minimal symptoms and skin lesions.

Mild fever and constitutional symptoms were first noted 1-2 days prior to characteristic skin and oral lesion in 75 (18.6%) cases. The oral (labial, buccal, tongue, palate) involvement was present in 135 (33.5%) patients and characteristic vesicular lesions involving hands [Figure 4a], feet [Figure 4b], elbows, perioral skin/oral mucosae [Figure 4c], buttocks [Figure 4d] and knees [Figure 4e] occurred variably in all patients. Pruritus/ burning sensation (in 15.4%), upper respiratory catarrh (in 12.7%), difficulty in swallowing (in 5.9%), and irritability in 2.9% cases, respectively, were other major symptoms noted in children. Only 12 (2.9%) children needed hospitalization for mild dehydration or fever (>38°C). The manifestations in adults were of minimal skin lesions without systemic symptoms [Figure 5]. Serology report for IgM CV A6 and CV A16 antibody available in three children from our center was negative.
All patients recovered completely after supportive management and none developed neurological or pulmonary complications. A 6-year-old child was brought back after 2 months with onychomadesis, leukonychia and mild nail dystrophy [Figure 6].

Figure 1: Geographic distribution of hand, foot and mouth disease cases in India (The 403 cases from this study are also shown here together with those reported from Himachal Pradesh in the literature). Note: News papers also reported small outbreaks of HFMD cases between the year 2012 and 2014 from - Delhi, Goa, Srinagar, Arunachal, Meghalaya, Nagaland, Manipur, Tripura, Mizoram, Daman and Diu, Lakshadweep, Punjab, Chandigarh, Haryana, Uttar Pradesh, Bihar, Madhya Pradesh, Chhattisgarh, and Jharkhand.
Results of reviewed literature

Three studies could not be retrieved and were excluded from review.\[^{[14-16]}\] The combined review of other 48 publications/reports in the English-language literature which could be retrieved and considered relevant for clinicoepidemiological and virological features of HFMD are tabulated in Supplementary Table S1.\[^{[12,13,17-62]}\]

Ten studies did not mention period of outbreak. Ten publications reported only single cases, 4 studies reported 2-5 cases and 34 were original or hospital based studies, and government reports. Overall, there were 3332 patients with HFMD reported between 2003 and 2019 across the entire country [Figure 1]. There were 3162 (94.9%) children (aged 4 months to 12 years), 132 (4%) adolescents (aged >12-18 years) and 38 (1.1) adults above 18 years of age.

Major outbreaks were during summers and rainy seasons (April to September) of the years 2013, 2015 and 2016 reporting 370 (11.1%), 533 (16%), and 465 (14%) cases, respectively [Figure 7]. However, small outbreaks had also occurred during autumn (October-November) and winter (December-January) months.

Fever and characteristic skin lesions starting on 2\(^{nd}\) to 3\(^{rd}\) day were the presenting features and were mild in most of the reviewed cases. Severe diarrhea and vomiting,
aseptic meningitis, and viral pneumonia in one case each were serious manifestations observed.\[58,59\] While one child had febrile convulsions, palmoplantar skin exfoliation in 12.6%, and onychomadesis and other nail changes were other complications in 34% children in a study.\[34\] Delayed nail changes were also noted in 26.4% and 48.2% patients in two separate studies.\[28,60\] All cases had improved spontaneously requiring only symptomatic treatment and parental counselling. Relapses had occurred within 3-4 weeks and as late as 1-5 years.\[27,29,35,36\]

**Causative viruses**

Only 17 publications had included virus identification studies by reverse transcriptase polymerase chain reaction (RT-PCR) for viral RNA isolation, compliment fixation test (CFT), or serology for IgM antibodies [Table 2]. Overall 1142 clinical samples from 1064 patients were tested for causative virus and mainly included throat swabs, vesicular fluid, and stool samples in majority and/or serum for antibody testing (19 samples). Viral RNA or IgM antibody results were positive in 617 (54%) of 1142 clinical samples. Serotyping identified CV A16 in 512 (83%), CV A6 in 105 (17%), and EV 71 in 25 (4.1%) samples, respectively. Multiple strains were identified in 72 (11.7%) samples. Untyped enteroviruses or EV A&B were identified in 3 (0.5%) samples.

**Discussion**

In our study, HFMD affected children mostly between 3 and 5 years of age but infection in older children and adults also occurred. Low grade fever and prodrome of upper respiratory catarrh remained the presenting symptoms followed by appearance of oral and skin lesions after 2-3 days with characteristic distribution in its typical form. The majority of the patients in the study had skin involvement starting after 2\(^{nd}\) to 3\(^{rd}\) day of prodrome while only 77% of our patients had oral lesions. No neurological or pulmonary manifestations were detected and all the cases improved spontaneously after symptomatic treatment in 7-10 days. Similar clinical presentation was also observed in the reviewed cases but with exception of severe diarrhea and vomiting, aseptic meningitis, and viral pneumonia in few cases. The long term complications such as palmoplantar exfoliation and nail changes too were
minimal. Overall, all these features in Indian patients with HFMD are in sync with its reported general clinical course.

India has weather conditions varying from arctic to temperate climates in outer Himalayas in the north to subtropical and tropical climates in the sub hills and plains, and coastal regions in the south. Both in terms of proximity and population India is only next to its neighboring China which had been the worst affected country in Southeast Asia. There was no evidence of the disease in India prior to first report of disease outbreak in 2003-2004 from Calicut followed by many outbreaks across regions with more or less similar epidemiological trends.\cite{12,13} A rising trend in new cases in our center was also seen from 2009 onwards that peaked in 2013 before declining in subsequent years coinciding with general trends of small scale outbreaks reported across almost all Indian states/union territories. The number of cases had peaked during summer months (May-June) of the years 2013, 2015 and 2016 at majority of the places. Small peaks in number of cases were also reported during autumn months from southern most states of Karnataka, Kerala, Tamil Nadu, Telangana, Andhra Pradesh, and West Bengal mostly

| Table 1: Baseline clinical features of our HFMD patients |
|--------------------|----------------|
| **Features** | **Number of patients (%) n=403** |
| **Children** | |
| n=390 | |
| Age (Mean) | 2 mo-18 y (3.1 years) |
| Range (Mean) | 2 mo-3 y |
| **Adults** | |
| n=13 | |
| Age (Mean) | 21-30 y |
| **Months with peak of HFMD cases** | May -June (summers) |
| **Family history of HFMD** | Present |
| **Clinical symptoms/signs** | Characteristic lesions |
| | Oral lesions |
| | Fever |
| | Pruritus/Burning |
| | Upper respiratory catarrh |
| | Feeding difficulties |
| | Irritability with or without disturbed sleep |
| | Hospitalization |
| | Delayed nail changes |

mo, months; y, years. * most patients had multiple symptoms

**Figure 5:** Characteristic skin lesions of hand, foot and mouth disease in mother of an affected child involving palms (a) and feet (b) only. Oral lesions were not perceptible in her.

**Figure 6:** Nail changes of onychomadesis of index fingernail, leukonychia and mild dystrophy of other nails in a 5-yr-old child seen 2 months after HFMD.
with subtropical/tropical climatic conditions year round. In comparison, review of epidemic trends in affected countries across the world showed that its first occurrence was in 1957 as a CV A16-associated mild febrile summer illness in Toronto (Canada) affecting about 60 individuals. Later epidemics had been sporadic and limited to small geographic areas in Australia, Brazil, Europe, Japan and United States.\textsuperscript{[63-68]} However with more frequent outbreaks and significantly high mortality among children over the years, it became a substantial public health problem in most Southeast Asian countries (Malaysia, Taiwan, China, Japan, Vietnam, Singapore, South Korea) between 1997 and

| Serial no. | Number of cases tested* | Number of positive results | Identity of virus=number of positive samples** | Laboratory Methods | Reference number |
|------------|-------------------------|---------------------------|----------------------------------------------|--------------------|-------------------|
| 1          | 11                      | 6                         | CVA-16=6                                    | CFT                | 28                |
| 2          | 62 (40 samples)         | 6                         | CVA 6=3                                     | RT-PCR             | 29                |
| 3          | 10                      | 6                         | CVA 16=1                                    | EV 71=2            | 31                |
| 4          | 78                      | 7                         | CVA 16=7                                    | RT-PCR             | 33                |
| 5          | 15                      | 13                        | CVA 16=2                                    | RT-PCR             | 35                |
| 6          | 7                       | 7                         | CVA 16=2                                    | RT-PCR             | 36                |
| 7          | 64 (158 samples)        | 158 samples               | CVA 4=44                                    | RT-PCR             | 37                |
| 8          | 68 (93 samples)         | 93 samples                | CVA 16=38                                   | RT-PCR             | 39                |
| 9          | 7                       | 2                         | CV A16=2                                    | RT-PCR             | 42                |
| 10         | 222                     | 189                       | CV A16=189                                  | RT-PCR             | 43                |
| 11         | 81                      | 19                        | IgM EV 71=19                                | Serology           | 13                |
| 12         | 60                      | 10                        | CVA 6=3                                     | RT-PCR             | 53                |
| 13         | 1                       | 1                         | CV A16=1                                    | RT-PCR             | 54                |
| 14         | 30 (78 Samples from vesicle fluid) | 18 | CV A16=78                                  | RT-PCR             | 55                |
| 15         | 101 (34 Samples)        | 18                        | CV A16=18                                   | RT-PCR             | 58                |
| 16         | 246                     | 63                        | CV A16=63                                   | RT-PCR             | 62                |
| 17         | 1                       | 1                         | CV A6=1                                     | RT-PCR             | 54                |
| Total      | 1064 (1142 samples)     | 617 (54.02%)              | CV A16=512 (83.0%)                          | -                  | -                 |

*Samples were mainly from throat swabs and stools swabs when not specified. **Some samples were positive for more than one virus type. ***Includes positive results by RT-PCR (6 samples) and Serology for IgM (19 samples). CFT, complement fixation test; CV, Coxsackie virus; EV, Enteroviruses; Ig, immunoglobulin; RT-PCR, reverse transcriptase polymerase chain reaction
except for few cases requiring hospitalization due to moderately severe presentation/complications. With some exceptions even patients infected with EV 71 showed no significant complications or fatal outcome, and recurrences were noted in only fewer cases. However, the true outcome perhaps remains unascertained from small number of virology reports.

Limitations

The study is limited by its retrospective design, lack of virological studies at our center, and the reviewed literature by itself may have missed some cases/reports. To delineate role played by global warming in general and changing climatic conditions in prolonging the duration of outbreaks during peak seasons or increased likelihood of its transmission throughout the year and aggravating the problem from public health point was not a part of study.

Conclusion and the Way Forward

The overall features of this long lasting HFMD epidemic affecting children, mild or no neurological or pulmonary manifestations in most patients, peaks of the disease happening during summer months and the causative virus strains appear in sync with global trends of the disease. However, it remains distinctly possible that mild nature of the disease in majority, and low level of awareness both among general public and primary health care providers might have led to continuation of ongoing epidemic at least in some regions. Nevertheless, the continuous spread of the disease across the country reminds of pre-epidemic periods of China and Taiwan. Thus, clinical and molecular research relevant to the disease, development of a safe and effective multivalent vaccine, monitoring of the disease outbreaks, and mass awareness programs for general public are urgent needs. The measures such as provision for clean drinking water, improved sanitation, and stringent hygiene practices remain important preventive measures for this growing public health problem. Strict surveillance for HFMD and making it a notifiable disease in India will perhaps be the first step in that direction.

Declaration

All authors declare that they have no competing interest and therefore nothing else to declare, and have contributed significantly and take full responsibility for the manuscript. The authors of the paper are obliged to confirm that it has not been previously published. The study was not funded by any agency.

Contributors’ statement

AS obtained compiled, tabulated all data and helped in literature search and preparing of the initial draft. VKM conceptualized, analyzed and interpreted data, drafted, and critically evaluated the manuscript for important intellectual content. KSM, PSC, SM, AC, MC, YRV, SH, and JS
helped in obtaining, compiling and interpretation of clinical data and literature search. All authors were involved in the preparation and revision of the draft manuscript and have agreed to the final contents.

**Statement of ethics**

All procedures followed were in accordance with the ethical standards of the responsible committee on human experimentation (institutional and national) and with the Helsinki Declaration of 1975, as revised in 2013.

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**Conflicts of interest**

There are no conflicts of interest.

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| Ref No. | Reference Citation | State & UTs (number of outbreaks) | Regions (number of cases) | Year(s) of outbreak | Number of cases | Results of viral studies (if any) | Remarks |
|---------|--------------------|----------------------------------|---------------------------|-------------------|----------------|----------------------------------|---------|
| 18      | IDSP–NCDC 2018     | Jammu & Kashmir (300)            | Baramulla (300)           | May 2018          | 300 (approx)  | -                                | ND      |
| 19      | IDSP–NCDC 2018     | Ladakh (537)                     | Leh (76)                  | Mar 2016          | 76            | -                                | ND      |
| 20      | Kadi et al 2019    | Leh (461)                        | April–June 2016           | 104               | 354           | -                                | 3       |
| 21      | Mohin et al 2010   | Himachal Pradesh (51)            | Kangra (1)                | June 2009         | -             | 1                                | 0       |
| 22      | Kashyap and Verma 2014 | Kashmir (300)                  | Srinagar                  | July–August 2013  | -             | 36                               | 6       |
| 23      | Sharma and Mansi 2018 | Ladakh (537)                    | Leh (104)                 | Aug–Nov 2017      | 354           | -                                | ND      |
| 24      | Agarwal et al 2015 | Rajasthan (38)                  | Udaipur (38)              | July–Sept 2012    | -             | 38                               | -       |
| 25      | Nanda, et al 2015  | Uttarakhand (330)                | Dehradun (33)             | Aug–Oct 2013      | 89            | 171                             | 30      |
| 26      | Kadri et al 2019   | Leh (461)                        | April–June 2016           | 104               | 354           | -                                | ND      |
| 27      | Kashyap and Verma 2014 | Jammu & Kashmir (300)            | Srinagar                  | July–August 2013  | -             | 36                               | 6       |
| 28      | Nanda, et al 2015  | Uttarakhand (330)                | Dehradun (33)             | Aug–Oct 2013      | 89            | 171                             | 30      |
| 29      | Sarma et al 2009   | West Bengal (338)                | Kolkata (89)              | June–Aug 2013, 2014 | 60            | 2                                | ND      |
| 30      | Nanda, et al 2015  | Uttarakhand (330)                | Dehradun (33)             | Aug–Oct 2013      | 89            | 171                             | 30      |
| 31      | Sarmi et al 2009   | West Bengal (338)                | Kolkata (89)              | June–Aug 2013, 2014 | 60            | 2                                | ND      |
| 32      | Nanda, et al 2015  | Uttarakhand (330)                | Dehradun (33)             | Aug–Oct 2013      | 89            | 171                             | 30      |
| 33      | Nanda, et al 2015  | Uttarakhand (330)                | Dehradun (33)             | Aug–Oct 2013      | 89            | 171                             | 30      |
| 34      | Nanda, et al 2015  | Uttarakhand (330)                | Dehradun (33)             | Aug–Oct 2013      | 89            | 171                             | 30      |
| 35      | Nanda, et al 2015  | Uttarakhand (330)                | Dehradun (33)             | Aug–Oct 2013      | 89            | 171                             | 30      |
| 36      | Nanda, et al 2015  | Uttarakhand (330)                | Dehradun (33)             | Aug–Oct 2013      | 89            | 171                             | 30      |
| 37      | Nanda, et al 2015  | Uttarakhand (330)                | Dehradun (33)             | Aug–Oct 2013      | 89            | 171                             | 30      |
| 38      | Nanda, et al 2015  | Uttarakhand (330)                | Dehradun (33)             | Aug–Oct 2013      | 89            | 171                             | 30      |

**Supplementary Table S1:** HFMD cases reported from various states and regions (arranged from north to south) of India.
| No. | Reference | Location | Year(s) | Age Group | Diagnosis | Details |
|-----|-----------|----------|---------|-----------|-----------|---------|
| 39  | Gopakrishna and Guinarkar 2020 (epidemiological and Virology study) | Pune and Kolhapur (68) | 2017-2018 | 1 mo-10 yr = 68 | - | CVA/A6, CVA/A6, Echo1 types (EVA-A and EV-B species) strains identified by RT-PCR studies for viral RNA/98 throat swabs, vesicular fluid and stool samples. These CV-A16 and CV-A6 strains exhibited 96-99% sequence identity with Indian strains. |
| 40  | Sreeji 2008, (case series) | Nagpur (4) | Sept 2006-April 2006 | - | 4 | - | ND |
| 41  | Kumar et al 2015 (observational study) | Karnataka (515) | Sept 2011 | - | - | - | ND |
| 42  | Sinha et al 2014 (virology study letter) | Bangalore (7) | Sept-Nov 2013 | - | 7 | - | - | CVA/A6 identified in 2 cases by RT-PCR studies for viral RNA. |
| 43  | Durga et al 2017 (virology study) | Bangalore (229) | 2013-2015 | 153 | 76 | - | - | CVA/A6 identified in 189 cases by RT-PCR studies for viral RNA. These CV-A16 strains exhibited 98-99% sequence identity with those reported in France and China. |
| 44  | Rao et al 2012 (case report) | Mangalore (1) | Not mentioned | - | 1 | - | - | ND |
| 45  | Rao et al, 2012 (case report) | Mangalore (1) | Not mentioned | - | 1 | - | - | ND |
| 46  | Kashyap et al 2015 (case report) | Mangalore (1) | Not mentioned | - | 1 | - | - | ND |
| 47  | Sankar et al 2015 (case report) | Andhra Pradesh (71) | Guntur (1) | Not mentioned | 1 | - | - | - | ND |
| 48  | Vani et al 2019 (observational study) | Guntur (70) | Oct 2017-Apr 2018 | 15 | 51 | 4 | - | ND |
| 49  | Mappa et al 2011 (case reports) | Telangana (110) | Hyderabad (1) | Not mentioned | - | 1 | - | - | ND |
| 50  | Kumar et al 2016 (cross-sectional, observational study) | Hyderabad (30) | Aug 2013-Jun 2014 | 40 | - | 10 | - | ND |
| 51  | Nagaraju et al 2019 (cross-sectional, observational study) | Adilabad (60) | Jan–Dec 2018 | 48 | 5 | 6 | - | - | ND |
| 52  | Mathew et al 2015 (case series) | Trivandrum (3) | Not mentioned | 0 | - | - | 3 | ND |
| 53  | Subitha et al 2018 (epidemiological and Virology study) | Kozhikode (60) | Sept 2015 | 38 | 15 | 1 | 5 | CVA/A-16 in 4, CVA/A6 in 31, EV (un typed) strains identified in 3 cases by RT-PCR studies for viral RNA. Constitutional symptoms were pronounced in adults. |
| 54  | Nagaraju et al 2019 (case report) | Kochi (1) | Aug 2015 | - | - | - | 1 | CVA-6 identified by RT-PCR study. |
| 56  | Vijayanaghrivas et al 2012 (virology investigative study) | Tamil Nadu (255) | Vellore (30) | Nov-Dec 2005, Jan-Feb 2008 | 30 | - | 1 | - | CVA/A6 identified by nested PCR in 30% of stools sample. |
| 56  | Thmena 2014 (case series) | Chennai (27) | Oct-Nov 2013 | 8 | 15 | 4 | - | ND |
| 57  | Sivakumar et al 2014 (case report) | Chennai (1) | Not Mentioned | - | 1 | - | - | ND |
| 58  | Kumar et al 2015 (original hospital based and community survey study) | Coonoor (101) (Wellington) | 2010 | < 5 yr = 83 | 18 | - | - | CVA/A6 identified in 18 of 101 samples by Aseptic meningitis and Viral pneumonia occurred in one case each. |
| 59  | Gargi 2017 | Kumbakonam (33) | Oct-Nov 2015 | 18 | 3 | 1 | 1 | - | One child had severe diarrhea and vomiting |
| 60  | Pichakacheri et al 2020 (observational study) | North Chennai (23) | Apr-Jun 2018 | 9mo-12 yr = 73 | ND | Onychomadesis and nail shedding occurred in 48.2% children |
| 61  | Harihar et al 2014(case report) | Puducherry (1) | Pillariyappam (1) | Not mentioned | 1 | - | - | - | ND |
| 62  | Palani, et al 2018 (hospital based study) | Andaman Nicobar (247) | Port Blair (246) | May 2013-Jun 2014 | <5 yr = 246 | - | - | - | CVA/A6 identified in 63 cases by RT-PCR study. These CV-A16 and CV-A6 strains exhibited sequence identity with strains in mainland India and Malaysia. |
| 54  | Nagarajan et al 2019 (case report) | Port Blair (1) | Sept 2011 | - | - | - | 1 | CVA/A6 identified by RT-PCR study. |