## National Institute for Health and Care Excellence

Final

## Fever in under 5s

[A] Evidence review for signs and symptoms predicting Kawasaki disease

NICE guideline NG143
Evidence review
November 2019

These evidence reviews were developed by the NICE Guideline Updates Team

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## Signs and symptoms predicting Kawasaki disease

## Review question

In children with fever, what symptoms and signs or combinations of symptoms and signs are predictive of Kawasaki disease?

## Introduction

Kawasaki disease is a rare but potentially serious cause of fever in children under 5. The NICE guideline on fever in under 5 s covers the assessment and early management of fever with no obvious cause in children aged under 5 . The aim of the guideline update was to review the evidence and update recommendations on identifying children who may go on to be diagnosed with Kawasaki disease.

Typical Kawasaki disease is diagnosed when fever is present for 5 days or longer and at least 4 out of 5 principal features are present (American Heart Association criteria, McCrindle 2017). For the purpose of this guideline, we use the term 'incomplete' Kawasaki disease for a clinical diagnosis of Kawasaki disease when fewer than 4 principal features are present. The term 'atypical' Kawasaki disease is also used in the evidence base, and this term is retained in the evidence tables (appendix E) when it was used in the primary studies.

## PICO table

| Population | The included population depends on study design: <br> - Cohort studies: Studies on children presenting with fever will be included. <br> - Case-control studies: Studies on children diagnosed with Kawasaki <br> disease (cases) vs control children not diagnosed with Kawasaki disease <br> will be included (irrespective of whether children explicitly do or do not <br> have fever on presentation). <br> - Case series: Case series of children diagnosed with Kawasaki disease <br> (irrespective of whether children explicitly do or do not have fever on <br> presentation). |
| :--- | :--- |
|  | Studies on children under the age of 5 will be included in the review. Studies <br> including older children (under 18) will also be included if these include <br> some children under the age of 5 : these studies will be considered as <br> indirect evidence. |
| Index test (signs | Clinical signs/symptoms at presentation consistent with the pathophysiology <br> of Kawasaki disease. Including but not limited to: |
| - Irritability |  |



## Outcomes

- Anterior uveitis
- Headache
- Neck stiffness
- Cracked/fissured, red lips
- Strawberry tongue
- Injected lips
- Injected pharynx
- Cough
- Cervical lymphadenopathy (swollen on one or both sides and tender and/or $>1.5 \mathrm{~cm}$ in diameter)
- Polymorphous rash:
- Perineal erythema and desquamation
- Macular, morbilliform or targetoid skin lesions of the trunk and extremities
- Psoriasiform eruption
- Inflammation and/or crust formation around the Bacille Calmette-Guérin (BCG) scar
- Gallop rhythm
- Added heart sounds
- Muffled heart tones
- Fusiform aneurysms of the brachial arteries palpable or visible in the axillae
- Abdominal pain
- Diarrhoea
- Vomiting
- Decreased intake
- Joint pain
- Arthritis
- Indurated oedema of hands and/or feet
- Diffuse erythema of palms and/or soles
- Sheet-like desquamation that begins in the periungual region of the hands and/or feet
- Linear nail creases (Beau's lines)
- Diagnosis of classic Kawasaki disease using the diagnostic criteria described by the American Heart Association in McCrindle 2017.
- Diagnosis of incomplete (atypical) Kawasaki disease using the diagnostic criteria described by the American Heart Association in McCrindle 2017.
- Diagnosis of Kawasaki disease with coronary artery aneurysm confirmed by imaging
Accuracy of risk factors at detecting Kawasaki disease:
- Diagnostic yield (percentage of cases with sign/symptom) and the mean time at which these signs/symptoms appeared relative to the onset of illness and then disappeared
- Sensitivity of each sign/symptom and sensitivity of specified combinations of signs/symptoms
- Specificity of each sign/symptom and specificity of specified combinations of signs/symptoms
- Positive likelihood ratio of each sign/symptom and positive likelihood ratio of specified combinations of signs/symptoms


## Methods and process

This evidence review was developed using the methods and process described in Developing NICE guidelines: the manual (2018). Methods specific to this review question are described in the review protocol in appendix A and the methods appendix (appendix B).

Declarations of interest were recorded according to NICE's 2018 declarations of interest policy.

## Clinical evidence

A systematic search was carried out for this review question to identify diagnostic accuracy studies, case series and systematic reviews of these studies, which found 2,701 references (see appendix C for literature search strategy). Based on title and abstract, 2,569 references were excluded and 132 references were ordered for screening based on their full texts.

Of the 132 references screened as full texts, 65 references were included based on their meeting the inclusion criteria specified in the review protocol (appendix A). All 65 included studies were case series. One study (Loh 2019) also presented case-control data. In addition to these studies, we added one of two studies that was used in the previous 2013 version of the guidelines (Huang 2006). This study met our inclusion criteria for this current update but the other did not as it was a case-series from outside Europe with fewer than 100 participants. Therefore, the total number of studies included in this evidence review was 66. The clinical evidence study selection is presented as a diagram in appendix D .

Meta-analysis was not appropriate for the case series included in this review question because there was a very high degree of heterogeneity across studies; the committee noted that the percentage of people with a particular sign or symptom was likely to be affected by geographical location, type of setting and the methods used to collect data (which was largely retrospective and identified from case notes). Therefore, the raw data was used to inform committee discussions and is presented in the GRADE profiles in appendix F. Medians and interquartile ranges were calculated to give an overall impression of the data for each symptom and these data are presented in Table 3: Principal symptoms. The number of signs and symptoms present was also included, as the committee agreed that this provided some useful information on combinations of symptoms that might be predictive of an eventual diagnosis, and combinations of symptoms were included in the review protocol.

We created a separate table of data for each sign or symptom. If data were available, we presented it separately for typical Kawasaki disease and incomplete Kawasaki disease. If a study did not have this data, we presented the data for all cases of Kawasaki disease together (typical and incomplete). With a view to reducing heterogeneity and assessing for indirectness caused by regional variation in clinical practice and genetics, data was further subdivided by region: UK data, Europe (not UK) and outside Europe.

Applications of GRADE methodology has not been developed for use with non-comparative studies; therefore a modified approach was applied using the GRADE framework (for details see Appendix B).

For the full evidence tables and GRADE profiles for included studies, please see appendix E and appendix F .

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## Excluded studies

See Appendix G for a list of references for excluded studies, with reasons for exclusion.

## Summary of clinical studies included in the evidence review

Table 1: Case series

| Study | Sample size | Setting | Location | Definition of Kawasaki disease |
| :---: | :---: | :---: | :---: | :---: |
| Advani 2019 | 542 | Hospital | Indonesia | Typical and atypical Kawasaki disease using AHA criteria |
| Bai 2017 | 383 | Hospital | China | Diagnosis of Kawasaki disease according to the Japanese Circulation Society Joint Working Group (Japan) |
| Baker 2009 | 198 | Hospital | USA and Canada | Typical and incomplete Kawasaki disease using AHA criteria |
| Bal 2014 | 106 | Hospital | USA | Typical and incomplete Kawasaki disease using AHA criteria |
| Behmadi 2019 | 176 | Hospital | Iran | Diagnosis of Kawasaki disease. No diagnostic criteria provided. |
| Boudiaf 2016 | 133 | Hospital | Algeria | Typical and incomplete Kawasaki disease using AHA criteria |
| Chang 2014 | 226 | Hospital | Taiwan | Typical (complete) Kawasaki disease: fever for at least 5 days and at least 4 of the 5 principal criteria |
| Chen 2016 | 351 | Hospital | Taiwan | Diagnosis of Kawasaki disease. No diagnostic criteria provided. |
| Ebbeson 2004 | 124 | Hospital | Canada | Typical and incomplete Kawasaki disease using the criteria as described by Han et al. 2000 (Canada) |
| Fabi 2018 | 302 | Hospital | Italy | Typical and incomplete Kawasaki disease using AHA criteria |
| Falcini 2007 | 266 | Hospital | Italy | Diagnosis of Kawasaki disease. No diagnostic criteria provided. |
| Falcini 2012 | 228 | Hospital | Turkey, Brazil and Italy | Diagnosis of Kawasaki disease. No diagnostic criteria provided. |
| Gamez- <br> Gonzalez $2013$ | 214 | Hospital | Mexico | Typical and incomplete Kawasaki disease using AHA criteria |
| Garrido-Garcia 2017 | 399 | Hospital | Mexico | Typical (complete) Kawasaki disease: fever for at least 5 days and at least 4 of the 5 principal criteria as described by the AHA <br> Atypical (incomplete) Kawasaki disease: fever for at least 5 days and 2 to 3 principal criteria + coronary involvement as described by the AHA |

[^0]| Study | Sample <br> size | Setting | Location | Definition of Kawasaki disease |
| :--- | :--- | :--- | :--- | :--- |
| Generini 1997 | 73 | Hospital | Italy | Diagnosis of Kawasaki disease according to the Kawasaki Disease Research <br> Committee (Japan) |
| Ghelani 2012 | 203 | Hospital | USA | Typical and incomplete Kawasaki disease using AHA criteria |
| Giannouli <br> 2013 | 86 | Hospital | Greece | Typical and incomplete Kawasaki disease using AHA criteria |
| Gorrab 2016 | 146 | Hospital | Canada (North African origin) | Diagnosis of Kawasaki disease. No diagnostic criteria provided. |
| Hu 2019 | 293 | Hospital | Taiwan | Typical and incomplete Kawasaki disease using AHA criteria |
| Huang 20061 | 768 | Hospital | Japan | Diagnosis of Kawasaki disease according to the Kawasaki Disease Research <br> Committee (Japan) |
| Jaggi 2018 | 135 | Hospital | USA | Typical and incomplete Kawasaki disease using AHA criteria |
| Jun 2015 | 355 | Hospital | South Korea | Typical and incomplete Kawasaki disease using AHA criteria |
| Jun 2017 | 146 | Hospital | South Korea | Typical and incomplete Kawasaki disease using AHA criteria |
| Kil 2017 | 615 | Hospital | South Korea | Typical and incomplete Kawasaki disease using AHA criteria |
| Kim 2017 | 14916 | Hospital | South Korea | Typical and incomplete Kawasaki disease using AHA criteria |

[^1]| Study | Sample size | Setting | Location | Definition of Kawasaki disease |
| :---: | :---: | :---: | :---: | :---: |
| Maric 2015 | 111 | Hospital | Croatia | Typical and incomplete Kawasaki disease using AHA criteria |
| Martins 2018 | 63 | Hospital | Portugal | Typical and incomplete Kawasaki disease using AHA criteria |
| Minich 2007 | 562 | Hospital | Canada and USA | Typical and incomplete Kawasaki disease using AHA criteria |
| Moore 2014 | 104 | Primary care | UK | Diagnosis of Kawasaki disease. No diagnostic criteria provided. |
| Nomura 2012 | 207 | Hospital | Japan | Diagnosis of Kawasaki disease according to Japanese Circulation Society Joint Working Group. |
| Patel 2013 | 314 | Hospital | Denmark | Discharge diagnosis of Kawasaki disease according to ICD-10. |
| Peng 2019 | 1420 | Hospital | China | Typical and incomplete Kawasaki disease using AHA criteria |
| Perrin 2009 | 59 | Hospital | France | Typical and incomplete Kawasaki disease using AHA criteria |
| Piao 2010 | 735 | Hospital | China | Diagnostic criteria for Kawasaki disease in the 7th International Kawasaki Disease Symposium (Japan) - further information not provided. |
| Ruan 2013 | 1209 | Hospital | China | Typical and incomplete Kawasaki disease using AHA criteria |
| Sanchez- <br> Maubens 2016 | 399 | Hospital | Spain | Typical and incomplete Kawasaki disease using AHA criteria |
| Sehgal 2015 | 312 | Hospital | USA | Typical and incomplete Kawasaki disease using AHA criteria |
| Shamsizadeh $2014$ | 104 | Hospital | Iran | Typical and incomplete Kawasaki disease using AHA criteria and American Academy of Paeditrics guideline. |
| Shiozawa $2014$ | 100 | Hospital | Japan | Diagnosis of Kawasaki disease according to the Kawasaki Disease Research Committee (Japan) |
| Sittiwangkul $2011$ | 170 | Hospital | Thailand | Typical and incomplete Kawasaki disease using AHA criteria |
| Sittiwangkul 2013 | 208 | Hospital | Thailand | Typical and incomplete Kawasaki disease using AHA criteria |
| Sonobe 2007 | 15857 | Hospital | Japan | Diagnosis of Kawasaki disease. No diagnostic criteria provided. |
| Stemberger $2018$ | 110 | Hospital | Croatia | Typical and incomplete Kawasaki disease using AHA criteria |

[^2]| Study | Sample <br> size | Setting | Location | Definition of Kawasaki disease |
| :--- | :--- | :--- | :--- | :--- |
| Sun 2018 | 1008 | Hospital | China | Typical and incomplete Kawasaki disease using AHA criteria |
| Tacke 2014 | 319 | Hospital | The Netherlands | Typical and incomplete Kawasaki disease using AHA criteria |
| Tajima 2015 | 100 | Hospital | Japan | Typical and incomplete Kawasaki disease using AHA criteria |
| Tang 2016 | 1016 | Hospital | China | Typical and incomplete Kawasaki disease using AHA criteria |
| Teng 2012 | 351 | Hospital | Taiwan | Diagnosis of Kawasaki disease. No diagnostic criteria provided. |
| Tewelde 2014 | 105 | Hospital | USA | Typical and incomplete Kawasaki disease using ICD-9 codes. |
| Uehara 2010 | 15524 | Hospital | Japan | Diagnosis of Kawasaki disease according to the Kawasaki Disease Research <br> Committee (Japan) |
| Wang 2009 | 243 | Hospital | USA | Typical and incomplete Kawasaki disease using AHA criteria |
| Yellen 2010 | 195 | Hospital | USA | Typical and incomplete Kawasaki disease using AHA criteria |
| Yoon 2016 | 239 | Hospital | South Korea | Typical and incomplete Kawasaki disease using AHA criteria |
| Yun 2011 | 121 | Hospital | South Korea | Diagnosis of Kawasaki disease according to the Kawasaki Disease Research <br> Committee (Japan) |
| Zhang 2016 | 518 | Hospital | Mongolia | Diagnosis of Kawasaki disease according to the Kawasaki Disease Research <br> Committee (Japan) |
| Zhang 2012 | 577 | Hospital | China | Diagnosis of Kawasaki disease according to the Kawasaki Disease Research <br> Committee (Japan) |
| Zhu 2015 | 231 | Hospital | China |  |
| Abbreviations | Heart Association, ICD: International Classification of Disease |  |  |  |
| AHA: American |  |  |  |  |
| 1.This study |  |  |  |  |

[^3]Table 2: Case-control studies

| Study | Sample size | Setting | Location | Definition of Kawasaki <br> disease |
| :--- | :--- | :--- | :--- | :--- |
| Loh 2019 | 279 | Hospital | Singapore | Typical and incomplete <br> Kawasaki disease using <br> AHA criteria |

See appendix E for full evidence tables.

## Summary of findings and quality assessment

The table below summarises the results for the principal symptoms of Kawasaki disease, as identified in the American Heart Association diagnostic criteria.

Data for other signs and symptoms, or symptoms that are sub-categorisations of the principal symptoms (e.g. oral changes - strawberry tongue) are presented in the GRADE profiles in appendix F .

Table 3: Principal symptoms

| Symptom | Kawasaki disease definition | Location | Age | Number of studies | \% with Symptom <br> Median (IQR) across studies | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Conjunctival injection (During course of illness) | Typical | Europe (not UK) | All | 3 | 92.8\% (91.6 to 93.1) | Very low |
|  |  | Outside Europe | All | 12 | 96.1\% (95.0 to 97.0) | Very low |
|  | Incomplete | Europe (not UK) | All | 3 | 65.0\% (46.1 to 71.9) | Very low |
|  |  | Outside Europe | All | 12 | 74.1\% (67.5 to 85.2) | Very low |
|  | Typical + Incomplete | Europe (not UK) | All | 6 | 83.4\% (71.3 to 92.1) | Very low |
|  |  | Outside Europe | <1 year | 7 | 84.4\% (55.3 to 89.9) | Very low |
|  |  |  | >1 year | 7 | 81.4\% (77.4 to 88.2) | Very low |
|  |  |  | All | 28 | 88.7\% (83.1 to 91.8) | Very low |
| Conjunctival injection (Primary care) | Typical + Incomplete | UK | All | 1 | 31.1\% (-) | Very low |

[^4]| Symptom | Kawasaki disease definition | Location | Age | Number of studies | \% with Symptom <br> Median (IQR) across studies | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Conjunctival injection (at time of emergence of symptoms other than fever) |  | Outside Europe | <2 year | 1 | 45.1\% (-) | Very low |
|  |  |  | >2 year | 1 | 16.3\% (-) | Very low |
| Conjunctival injection (day 5 of fever) |  | Outside Europe | <2 year | 1 | 84.3\% (-) | Very low |
|  |  |  | >2 year | 1 | 85.7\% (-) | Very low |
| Oral changes (During course of illness) | Typical | Europe (not UK) | All | 3 | 98.4\% (96.7 to 99.2) | Very low |
|  |  | Outside Europe | All | 12 | 96.1\% (93.2 to 97.6) | Very low |
|  | Incomplete | Europe (not UK) | All | 3 | 65.0\% (64.3 to 73.4) | Very low |
|  |  | Outside Europe | All | 10 | 63.6\% (57.9 to 67.9) | Very low |
|  | Typical + Incomplete | Europe (not UK) | All | 5 | 94.5\% (87.5 to 96.6) | Very low |
|  |  | Outside Europe | <1 year | 6 | 65.2\% (47.4 to 81.8) | Very low |
|  |  |  | >1 year | 6 | 77.0\% (76.2 to 78.3) | Very low |
|  |  |  | All | 27 | 89.7\% (83.8 to 94.3) | Very low |
| Oral changes (at time of emergence of symptoms other than fever) | Typical + Incomplete | Outside Europe | <2 year | 1 | 29.4\% (-) | Very low |
|  |  | Outside Europe | >2 year | 1 | 12.2\% (-) | Very low |
| Oral changes (day 5 of fever) | Typical + Incomplete | Outside Europe | <2 year | 1 | 64.7\% (-) | Very low |
|  |  | Outside Europe | >2 year | 1 | 73.5\% (-) | Very low |
| Changes in extremities (During course of illness) | Typical | Europe (not UK) | All | 3 | 78.2\% (71.9 to 87.8) | Very low |
|  |  | Outside Europe | All | 10 | 88.3\% (84.3 to 90.8) | Very low |
|  | Incomplete | Europe (not UK) | All | 3 | 30.0\% (21.8 to 42.2) | Very low |
|  |  | Outside Europe | All | 11 | 33.3\% (22.4 to 40.0) | Very low |
|  | Typical + Incomplete | Europe (not UK) | All | 5 | 82.0\% (77.4 to 85.7) | Very low |
|  |  | Outside Europe | <1 year | 5 | 36.4\% (26.9 to 46.2) | Very low |

[^5]| Symptom | Kawasaki disease definition | Location | Age | Number of studies | \% with Symptom <br> Median (IQR) across studies | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | >1 year | 5 | 61.5\% (50.0 to 73.8) | Very low |
|  |  |  | All | 24 | 72.6\% (65.9 to 83.4) | Very low |
| Changes in Extremities (at time of emergence of symptoms other than fever) | Typical + Incomplete | Outside Europe | <2 year | 1 | 19.6\% (-) | Very low |
|  |  | Outside Europe | >2 year | 1 | 6.1\% (-) | Very low |
| Changes in Extremities (day 5 of fever) | Typical + Incomplete | Outside Europe | <2 year | 1 | 62.7\% (-) | Very low |
|  |  | Outside Europe | >2 year | 1 | 61.2\% (-) | Very low |
| Polymorphous rash (During course of illness) | Typical | Europe (not UK) | All | 3 | 93.6\% (87.1 to 94.3) | Very low |
|  |  | Outside Europe | All | 12 | 93.5\% (89.4 to 97.5) | Very low |
|  | Incomplete | Europe (not UK) | All | 3 | 65.0\% (64.3 to 70.4) | Very low |
|  |  | Outside Europe | All | 13 | 66.2\% (48.3 to 68.6) | Very low |
|  | Typical + Incomplete | Europe (not UK) | All | 6 | 88.0\% (84.7 to 92.9) | Very low |
|  |  | Outside Europe | <1 year | 13 | 86.3\% (74.5 to 89.7) | Very low |
|  |  |  | >1 year | 6 | 81.8\% (79.1 to 89.1) | Very low |
|  |  |  | All | 26 | 85.5\% (79.4 to 93.3) | Very low |
| Polymorphous rash (Primary care) | Typical + Incomplete | UK | All | 1 | 63.5\% (-) | Very low |
| Polymorphous rash (at time of emergence of symptoms other than fever) |  | Outside Europe | <2 year | 1 | 78.4\% (-) | Very low |
|  |  |  | >2 year | 1 | 24.5\% (-) | Very low |
| Polymorphous rash (day 5 of fever) |  | Outside Europe | <2 year | 1 | 92.2\% (-) | Very low |
|  |  |  | >2 year | 1 | 79.6\% (-) | Very low |
| Cervical lymphadenopathy | Typical | Europe (not UK) | All | 3 | 43.6\% (41.1 to 58.2) | Very low |
|  |  | Outside Europe | All | 11 | 63.0\% (41.6 to 72.7) | Very low |

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| Symptom | Kawasaki disease definition | Location | Age | Number of studies | \% with Symptom <br> Median (IQR) across studies | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| (During course of illness) | Incomplete | Europe (not UK) | All | 3 | $30.0 \%$ (28.7 to 44.6) | Very low |
|  |  | Outside Europe | All | 12 | 31.4\% (20.8 to 38.8) | Very low |
|  | Typical + Incomplete | Europe (not UK) | All | 6 | 53.0\% (31.7 to 71.8) | Very low |
|  |  | Outside Europe | <1 year | 7 | 24.6\% (14.9 to 30.9) | Very low |
|  |  |  | >1 year | 7 | $57.3 \%$ (33.7 to 63.1) | Very low |
|  |  |  | All | 27 | $59.4 \%$ (38.2 to 68.3) | Very low |
| Cervical lymphadenopathy (primary care) | Typical + Incomplete | UK | All | 1 | 35.1\% (-) | Very low |
| Cervical <br> lymphadenopathy (at time of emergence of symptoms other than fever) | Typical + Incomplete | Outside Europe | <2 year | 1 | 35.3\% (-) | Very low |
|  |  |  | >2 year | 1 | 75.5\% (-) | Very low |
| Cervical lymphadenopathy (day 5 of fever) | Typical + Incomplete | Outside Europe | <2 year | 1 | 64.7\% (-) | Very low |
|  |  |  | $>2$ year | 1 | 93.9\% (-) | Very low |
| 2 or more principal symptoms (at time of emergence of symptoms other than fever) | Typical + Incomplete | Outside Europe | <2 year | 1 | 54.9\% (-) | Very low |
|  |  |  | >2 year | 1 | 16.3\% (-) | Very low |

Table 4: Numbers of symptoms

| Symptom | Kawasaki disease <br> definition | Location | Age | Number of <br> studies | \% with Symptom <br> $(95 \% \mathrm{CI})$ |  |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| 0 symptoms ${ }^{1}$ (at first <br> presentation | Typical + Incomplete | UK | All | 1 (Moore <br> $2014)$ | $28.4 \%(19.4$ to 38.5) | Very low |

[^6]| Symptom | Kawasaki disease definition | Location | Age | Number of studies | \% with Symptom (95\% CI) | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 symptom ${ }^{1}$ (at first presentation) | Typical + Incomplete | UK | All | $\begin{aligned} & 1 \text { (Moore } \\ & 2014 \text { ) } \end{aligned}$ | 39.2\% (28.9 to 50.6) | Very low |
| 2 symptoms ${ }^{1}$ (at first presentation) | Typical + Incomplete | UK | All | $\begin{aligned} & 1 \text { (Moore } \\ & 2014 \text { ) } \end{aligned}$ | 9.5\% (4.7 to 18.3) | Very low |
| 2 or more principal symptoms ${ }^{2}$ (at time of emergence of symptoms other than fever) | Typical + Incomplete | Outside Europe | <1 year | 1 (Shiozawa 2014) | 54.9\% (41.4 to 67.7) | Very low |
|  |  |  | >1 year | 1 (Shiozawa 2014) | 16.3\% (8.5 to 29.0) | Very low |
| 3 symptoms ${ }^{1}$ (at first presentation) | Typical + Incomplete | UK | All | $\begin{aligned} & 1 \text { (Moore } \\ & \text { 2014) } \end{aligned}$ | $6.8 \%$ (2.9 to 14.9) | Very low |
| 4 symptoms $^{1}$ (at first presentation) | Typical + Incomplete | UK | All | $\begin{aligned} & 1 \text { (Moore } \\ & 2014 \text { ) } \end{aligned}$ | 5.4\% (2.1 to 13.1) | Very low |
| 5 symptoms ${ }^{1}$ (at first presentation) | Typical + Incomplete | UK | All | $\begin{aligned} & 1 \text { (Moore } \\ & 2014 \text { ) } \end{aligned}$ | 5.4\% (2.1 to 13.1) | Very low |
| 6 symptoms ${ }^{1}$ (at first presentation) | Typical + Incomplete | UK | All | $\begin{aligned} & 1 \text { (Moore } \\ & 2014 \text { ) } \end{aligned}$ | 1.4\% (0.2 to 7.3) | Very low |
| 7 symptoms ${ }^{1}$ (at first presentation) | Typical + Incomplete | UK | All | $\begin{aligned} & 1 \text { (Moore } \\ & 2014 \text { ) } \end{aligned}$ | 1.4\% (0.2 to 7.3) | Very low |
| 8 symptoms ${ }^{1}$ (at first presentation) | Typical + Incomplete | UK | All | $\begin{aligned} & 1 \text { (Moore } \\ & 2014) \end{aligned}$ | 2.7\% (0.7 to 9.3) | Very low |
| 1.Symptoms included Rash, Lymphadenopathy, Conjunctivitis, Red, dry, or cracked lips, Strawberry tongue, Redness in mouth, Peeling skin, Red palms/soles, Oedema |  |  |  |  |  |  |

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Table 5: Case-control evidence

| No. of studies | Study design | Sample <br> size | Sensitivity (95\%CI) | Specificity (95\%CI) | Effect size (95\%CI) | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| BCG scar activation <br> Typical + incomplete Kawasaki disease, Outside Europe, all ages ${ }^{3}$ |  |  |  |  |  |  |
| $\begin{aligned} & 1 \text { (Loh } \\ & 2019) \end{aligned}$ | Case control | 370 | 0.43 (0.37, 0.49) | 0.90 (0.82, 0.95) | $\begin{aligned} & \text { LR }+4.31(2.29,8.14) \\ & \text { LR- } 0.64(0.56,0.72) \end{aligned}$ | Very low Very low |

See appendix F for full GRADE profiles.

## Economic evidence

Standard health economics filters were applied to the clinical search strategy to identify relevant cost-utility analyses. In total, 129 references were returned; all could be confidently excluded on screening of titles and abstracts. No original health economic analysis was undertaken for this guideline update, as it was agreed that any recommendations made were highly unlikely to result in a substantial resource impact.

## The committee's discussion of the evidence

## Interpreting the evidence

## The outcomes that matter most

The question focused on identifying signs and symptoms that could predict which children would eventually receive a diagnosis of Kawasaki disease. Children suspected as having a possible diagnosis of Kawasaki disease are referred to secondary care for further assessment, which may include blood tests and cardiac imaging. A definitive diagnosis of Kawasaki disease is made by a paediatrician in secondary care and is outside the scope of this review, because the NICE guideline on Fever in under 5 s is limited to the initial and assessment and management of children presenting with fever.

The consequences of missing a possible diagnosis early in the course of disease (a 'false negative' result) are potentially serious. If clinicians in primary care fail to identify possible cases this is likely to delay or prevent onward referral to secondary care for definitive diagnosis and treatment. Untreated disease or disease treated late in the course of the illness may result in coronary artery abnormalities, which are associated with long-term morbidity and mortality. Correctly identifying children with Kawasaki disease (a 'true positive' result) early in the course of disease would lead to prompt referral and treatment, which would reduce the risk of long-term complications. Children incorrectly identified as potentially having Kawasaki disease when in fact they do not (a 'false positive' result) may receive unnecessary referral and testing, which might be invasive and distressing for children and their parents and carers and may also delay correct diagnosis and treatment. However, given the serious consequences of missing cases of Kawasaki disease, the committee agreed that the sensitivity of signs and symptoms should be valued over their specificity, and that it was important to identify cases early, even at the expense of referring children without the disease for further assessment.

## The quality of the evidence

The evidence identified for this review was all very low quality. The case series identified were exclusively retrospective, and based on reviews of case notes, or in some cases, questionnaires asking about previous signs and symptoms. This was a major limitation as recall of signs and symptoms may be poor and subject to bias and recording of signs and symptoms in case notes is likely to be variable and incomplete. The majority of evidence was reported during the whole time-course of the disease, and so was of limited applicability to the review question, which focused on signs and symptoms early in the course of the illness which predicted an eventual diagnosis. The committee noted that data for typical and incomplete cases together was in fact most applicable to the review question, as complications of Kawasaki disease occur in both typical and incomplete disease, and the treatment is the same irrespective of presentation.

Two case series provided evidence on signs and symptoms earlier in the illness. Shiozawa et al. (2013) was a Japanese case series providing data on signs and symptoms relative to
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the onset of fever based on information from parental interviews, family doctors and hospital records. There were limited details on how the data had been collected. The committee also noted that Kawasaki disease was much more common in Japan and other areas of Asia than in the UK, and was likely to be diagnosed earlier. This study was therefore of limited applicability to the UK population and was rated down for indirectness.

Moore et al. (2014) reported a case series from UK primary care and so was more directly applicable to the review question. However, the committee was concerned that the method of selection of cases might be unrepresentative of typical cases of Kawasaki disease because inclusion in the study required a 'convincing diagnosis' of Kawasaki disease.

One case-control study was identified, which assessed the diagnostic accuracy of BCG scar activation. The committee agreed that the study was of very low quality, as assessment of the BCG scar differed between case and control groups. They also noted that the study was of limited applicability to the UK, as BCG vaccination is not routinely offered. The committee estimated that fewer than $20 \%$ of children with Kawasaki disease were in the groups offered BCG vaccination. No other evidence on the specificity of signs and symptoms was available, which was a major limitation of the evidence.

The target population for the review was children under 5 presenting with fever. Most of the evidence included in the review was from case-series of children with a diagnosis of Kawasaki disease, but fever was usually not an explicit inclusion criterion in the studies. However, the committee agreed that presence of fever was always required for a Kawasaki disease diagnosis and so all of the participants in the included studies were likely to have fever.

## Benefits and harms

The principal features of Kawasaki disease specified in the American Heart Association guideline for diagnosis are conjunctival injection, cervical lymphadenopathy, polymorphous rash, oral changes and changes to the hands and feet. The committee agreed that these features should be used by clinicians when assessing children for possible Kawasaki disease as they are present in the majority of patients at some point in the course of disease. Therefore, combinations of these features together with a fever lasting at least 5 days are likely to be reasonably specific, based on the committee's experience (children without Kawasaki disease are unlikely to exhibit the same symptoms). Data on other symptoms were reviewed, but these symptoms were either present in a small number of people with Kawasaki disease, were inconsistent across studies, or were thought likely to be non-specific to Kawasaki disease and so were not incorporated into recommendations.

Evidence from early in the course of disease (before a definitive diagnosis was made) showed that the number of children experiencing each principal symptom was lower. A Japanese study (Shiozawa et al., 2013) found that the number of children experiencing each of the principal symptoms 5 days after the onset of fever was between $60 \%$ and $95 \%$ for each of the principal symptoms in those aged under and over 2 years. The committee considered that these data might in fact overestimate the number of children experiencing symptoms at this stage of the disease in the UK because incomplete Kawasaki disease is more common in the UK population, and, based on the experience of the committee, features are less obvious.

A UK study (Moore et al., 2014) found that most children had either 0 or 1 symptoms (which could include fever) at first presentation in primary care. Consequently, the committee agreed that clinicians should think about Kawasaki disease when fever was present for 5 days or longer, even when no additional features are present. The committee decided to list the principal features of Kawasaki disease in the recommendation without specifying a particular number of features that must be present to prompt further action. This was
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because no evidence on the sensitivity and specificity of combinations of symptoms early in the disease course was available, and clinicians should use their clinical judgment when deciding on the likelihood of Kawasaki disease as a differential diagnosis and the need for onward referral. The recommendation does not distinguish between typical and incomplete Kawasaki disease because the committee felt that this distinction was not useful when assessing for possible Kawasaki disease - the actions required by clinicians is the same irrespective of whether typical or incomplete disease is suspected. Complications in the absence of prompt treatment are possible with both typical and incomplete disease.

A research recommendation was made for a prognostic diagnostic accuracy study on the signs and symptoms that are predictive of a subsequent diagnosis of Kawasaki disease, which would allow a more specific recommendation to be made in future updates of this guidance.

The committee recommended that clinicians should ask about the presence of the principal features of Kawasaki disease in children with a fever lasting 5 days or longer since the onset of fever because, based on the experience of the committee some of the features may have appeared and disappeared at the time of assessment.

When data for children with typical and incomplete disease was reported for children under the age of 1 year separately, the proportion of children with each of the principal symptoms was lower for some principal features in the under-1 age-group. This is consistent with the experience of the committee that under-1s are more often diagnosed with incomplete disease, and with a UK-based epidemiological study that the committee was aware of (see 'other factors the committee took into account'). The committee agreed that it was important that healthcare professionals were aware that under-1s typically present with fewer symptoms than older children, as this is something that the committee thought was not widely known and should increase the index of suspicion for Kawasaki disease in this group, even when few or no additional features are present.

## Cost effectiveness and resource use

The committee agreed that the new recommendations are unlikely to lead to a substantial resource impact: recommendations elsewhere in this guideline already categorise fever of 5 days or longer on its own as an 'amber' feature that should prompt non-paediatric practitioners to refer to specialist care for further assessment or provide 'safety-netting' advice. Therefore, the updated recommendations will not create an additional population of children who were not previously considered at risk, so they are unlikely to cause a dramatic increase in referrals. Moreover, even though more children may be referred to specialist care, the costs of false-positive referrals are small compared with the costs of false-negative failures to identify children with Kawasaki disease, which is more likely to be associated with mortality, morbidity and lifelong cardiological care if it is not identified promptly.

## Other factors the committee took into account

The committee was aware of a recent UK and Ireland based epidemiological study (Tulloh et al, 2018) on Kawasaki disease which did not match the criteria specified in the review protocol (as it did not report signs or symptoms) but provided useful information on the incidence of Kawasaki disease in the UK population. The study showed that the incidence of Kawasaki disease in the UK has risen in the last 10 years. The incidence of incomplete or atypical Kawasaki disease was highest in children under 1 year, and the rate of coronary artery aneurysm was highest in this group. The study also suggested, in keeping with epidemiological studies in other countries, that the incidence of Kawasaki disease is higher in children of black or Asian ethnicity. The committee highlighted this an equality issue, as clinicians might not know that Kawasaki disease is more common in these groups and so it
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might be under diagnosed. However, a specific recommendation on this was not made on this issue as the focus of the review was to identify signs and symptoms predictive of Kawasaki disease.

The 2013 version of the NICE guideline on Fever in under 5 s included a description of the features of Kawasaki disease. The committee decided to simplify the descriptions of additional features other than a fever of 5 days or longer and make them consistent with the diagnostic criteria for Kawasaki disease specified by the American heart association (McCrindle 2017).

## Appendices

## Appendix A - Review protocol

Review protocol for signs and symptoms predicting Kawasaki disease

| Field (based on PRISMA-P | Content |
| :--- | :--- |
| Review question | In children with fever, what symptoms and signs or combinations of symptoms and signs are predictive of Kawasaki <br> disease? |
| Type of review question | Diagnostic |
| Objective of the review | To determine what symptoms and signs or combination of symptoms and signs are predictive of Kawasaki disease in <br> children with fever. |
| Eligibility criteria - <br> population/disease/condition/issu <br> e/domain | The included population depends on study design: <br> - Cohort studies: Studies on children presenting with fever will be included. <br> Case-control studies: Studies on children diagnosed with Kawasaki disease (cases) vs control children not <br> diagnosed with Kawasaki disease will be included (irrespective of whether children explicitly do or do not have fever <br> on presentation). <br> Case series: Case series of children diagnosed with Kawasaki disease (irrespective of whether children explicitly <br> do or do not have fever on presentation). |

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|  | Studies on children under the age of 5 will be included in the review. Studies including older children (under 18) will <br> also be included if there include some children under the age of 5: these studies will be considered as indirect <br> evidence. |
| :--- | :--- |
| Eligibility criteria - intervention | Clinical signs/symptoms at presentation consistent with the pathophysiology of Kawasaki disease. Including but not <br> limited to: <br> $-\quad$ Appearance at the foot of the bed: <br> $>$ Irritability |
|  | Circulation and pulse: <br> $>$ Reduced perfusion: cold, pale or cyanotic digits of the hands and/or feet |


|  | $>$ Tachycardia out of proportion to the degree of fever as defined by the study <br> - Fever ( $\geq 38^{\circ} \mathrm{C}$ or $\geq 100.4 \mathrm{~F}$ ): <br> $>$ Duration of fever <br> $>$ Persistent fever ( 5 days or more) despite antibiotics <br> - Eyes: <br> > Bilateral conjunctival injection without exudate <br> > Photophobia <br> > Anterior uveitis <br> - Aseptic meningitis: <br> > Headache <br> > Neck stiffness <br> - Oral (mucositis): <br> > Cracked/fissured, red lips <br> > Strawberry tongue <br> > Injected lips <br> > Injected pharynx <br> > Cough <br> - Cervical lymphadenopathy (swollen on one or both sides and tender and/or $>1.5 \mathrm{~cm}$ in diameter) <br> - Skin changes: <br> > Polymorphous rash: <br> - Perineal erythema and desquamation <br> - Macular, morbilliform or targetoid skin lesions of the trunk and extremities <br> > Psoriasiform eruption <br> > Inflammation and/or crust formation around the Bacille Calmette-Guérin (BCG) scar <br> - Heart and large arteries: <br> > Gallop rhythm <br> > Added heart sounds <br> > Muffled heart tones <br> > Fusiform aneurysms of the brachial arteries palpable or visible in the axillae <br> - Gastrointestinal: <br> > Abdominal pain <br> > Diarrhoea <br> > Vomiting <br> $>$ Decreased intake |
| :---: | :---: |

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|  | Joint pain <br> > Arthritis <br> - Extremity changes: <br> > Indurated oedema of hands and/or feet <br> > Diffuse erythema of palms and/or soles <br> $>$ Sheet-like desquamation that begins in the periungual region of the hands and/or feet <br> $>$ Linear nail creases (Beau's lines) |
| :---: | :---: |
| Eligibility criteria - reference standard | - Diagnosis of classic Kawasaki disease using the diagnostic criteria described by the American Heart Association in McCrindle 2017. <br> - Diagnosis of incomplete (atypical) Kawasaki disease using the diagnostic criteria described by the American Heart Association in McCrindle 2017. <br> - Diagnosis of Kawasaki disease with coronary artery aneurysm confirmed by imaging |
| Outcomes | Accuracy of risk factors at detecting Kawasaki disease: <br> - Diagnostic yield of signs/symptoms and the mean time at which these signs/symptoms appeared relative to the onset of illness and then disappeared <br> - Sensitivity of each sign/symptom and sensitivity of specified combinations of signs/symptoms <br> - Specificity of each sign/symptom and specificity of specified combinations of signs/symptoms <br> - Positive likelihood ratio of each sign/symptom and positive likelihood ratio of specified combinations of signs/symptoms <br> - Negative likelihood ratio of each sign/symptom and negative likelihood ratio of specified combinations of signs/symptoms |
| Eligibility criteria - study design | - Diagnostic accuracy studies (cohort, cross sectional and case-control). <br> - Case series |
| Other inclusion/exclusion criteria | Exclusion <br> - Non-English language studies |

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|  | - Studies that do not include any children under 5 years of age. However, if we do not find any studies that include children under 5 years of age, then we will include studies with children over the age of 5 years but under the age of 18 . <br> - For case series, studies will be excluded if they do not meet the following sample size requirements: <br> - From the UK, sample size >1 patient; from Europe, sample size $\geq 50$ patients; from the rest of the world, sample size $\geq 100$ patients. |
| :---: | :---: |
| Sub-group analysis | - Children under 1 year of age. <br> - Ethnicity. |
| Selection process - duplicate screening/selection/analysis | $10 \%$ of the abstracts will be reviewed by two reviewers, with any disagreements will be resolved by discussion or, if necessary, a third independent reviewer. If meaningful disagreements are found between the different reviewers, a further $10 \%$ of the abstracts will be reviewed by two reviewers, with this process continuing until agreement is achieved between the two reviewers. From this point, the remaining abstracts will be screened by a single reviewer. |
| Data management (software) | See Appendix B. |
| Information sources - databases and dates | Sources to be searched <br> - Clinical searches - Medline, Medline in Process, Medline EPub Ahead of Print, Embase, Cochrane CDSR, CENTRAL, DARE (legacy records) and HTA. <br> - Economic searches - Medline, Medline in Process, Medline EPub Ahead of Print, Embase, NHS EED (legacy records) and HTA, with economic evaluations and quality of life filters applied where appropriate. <br> Supplementary search techniques <br> - None identified <br> Limits |

[^11]|  | - Studies reported in English <br> - No Study design limits will be set <br> - Animal studies will be excluded from the search results <br> - Conference abstracts will be excluded from the search results <br> - Date limited from January $1^{\text {st }} 2007$ to present |
| :---: | :---: |
| Identify if an update | N/A |
| Author contacts | Guideline updates team |
| Highlight if amendment to previous protocol | This is a new protocol. |
| Search strategy - for one database | For details please see appendix C of relevant chapter. |
| Data collection process forms/duplicate | A standardised evidence table format will be used and published as appendix D (clinical evidence tables) or H (economic evidence tables). |
| Data items - define all variables to be collected | For details please see evidence tables in appendix D (clinical evidence tables) or H (economic evidence tables). |
| Methods for assessing bias at outcome/study level | Study checklists were used to critically appraise individual studies. For details please see appendix H of Developing NICE guidelines: the manual <br> The following checklist will be used: <br> Risk of bias of diagnostic accuracy studies will be assessed using QUADAS-2 |

[^12]|  | $\underline{\text { Risk of bias for case series will be assessed using the Joanna Briggs institute critical appraisal checklist for case series. }}$ |
| :--- | :--- |
|  | The risk of bias across all available evidence will be evaluated for each outcome using an adaptation of the 'Grading of <br> Recommendations Assessment, Development and Evaluation (GRADE) toolbox' developed by the international <br> GRADE working group http://www.gradeworkinggroup.org/ |
| Criteria for quantitative synthesis | For details please see section 6 of Developing NICE guidelines: the manual. |
| Methods for quantitative analysis <br> - combining studies and <br> exploring (in)consistency | For details please see the methods and process section of the main file. |
| Meta-bias assessment - <br> publication bias, selective <br> reporting bias | For details please see section 6.2 of Developing NICE guidelines: the manual. |
| Confidence in cumulative <br> evidence | For details please see sections 6 and 9 of Developing NICE guidelines: the manual. |
| Rationale/context - what is <br> known | For details please see the introduction to the evidence review in the main file. |

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| Sources of funding/support | The NICE Guideline Updates Team is an internal team within NICE. |
| :--- | :--- |
| Name of sponsor | The NICE Guideline Updates Team is an internal team within NICE. |
| Roles of sponsor | The NICE Guideline Updates Team is an internal team within NICE. |
| PROSPERO registration number |  |

## Appendix B - Methods

## Priority screening

The review made use of the priority screening functionality with the EPPI-reviewer systematic reviewing software. This uses a machine learning algorithm (specifically, an SGD classifier) to take information on features (1, 2 and 3 word blocks) in the titles and abstract of papers marked as being 'includes' or 'excludes' during the title and abstract screening process, and re-orders the remaining records from most likely to least likely to be an include, based on that algorithm. This re-ordering of the remaining records occurs every time 25 additional records have been screened.

Research is currently ongoing as to what are the appropriate thresholds where reviewing of abstract can be stopped, assuming a defined threshold for the proportion of relevant papers it is acceptable to miss on primary screening. For this evidence review 'includes' were identified throughout the screening process, and therefore all records in the database were examined.

## Incorporating published systematic reviews

For all review questions where a literature search was undertaken looking for a particular study design, systematic reviews containing studies of that design were also included. All included studies from those systematic reviews were screened to identify any additional relevant primary studies not found as part of the initial search.

## Quality assessment

Individual systematic reviews were quality assessed using the ROBIS tool, with each classified into one of the following three groups:

- High quality - It is unlikely that additional relevant and important data would be identified from primary studies compared to that reported in the review, and unlikely that any relevant and important studies have been missed by the review.
- Moderate quality - It is possible that additional relevant and important data would be identified from primary studies compared to that reported in the review, but unlikely that any relevant and important studies have been missed by the review.
- Low quality - It is possible that relevant and important studies have been missed by the review.

Each individual systematic review was also classified into one of three groups for its applicability as a source of data, based on how closely the review matches the specified review protocol in the guideline. Studies were rated as follows:

- Fully applicable - The identified review fully covers the review protocol in the guideline.
- Partially applicable - The identified review fully covers a discrete subsection of the review protocol in the guideline (for example, some of the factors in the protocol only).
- Not applicable - The identified review, despite including studies relevant to the review question, does not fully cover any discrete subsection of the review protocol in the guideline.


## Using systematic reviews as a source of data

If systematic reviews were identified as being sufficiently applicable and high quality, and were identified sufficiently early in the review process (for example, from the surveillance review or early in the database search), they were used as the primary source of data, rather than extracting information from primary studies. The extent to which this was done depended on the quality and applicability of the review, as defined in Table 6. When systematic reviews were used as a source of primary data, and unpublished or additional data included in the review which is not in the primary studies was also included. Data from these systematic reviews was then quality assessed and presented in GRADE/CERQual tables as described below, in the same way as if data had been extracted from primary studies. In questions where data was extracted from both systematic reviews and primary studies, these were cross-referenced to ensure none of the data had been double counted through this process.

Table 6: Criteria for using systematic reviews as a source of data

| Quality | Applicability | Use of systematic review |
| :--- | :--- | :--- |
| High | Fully applicable | Data from the published systematic review were used instead of <br> undertaking a new literature search or data analysis. Searches <br> were only done to cover the period of time since the search date <br> of the review. |
| High | Partially applicable | Data from the published systematic review were used instead of <br> undertaking a new literature search and data analysis for the <br> relevant subsection of the protocol. For this section, searches <br> were only done to cover the period of time since the search date <br> of the review. For other sections not covered by the systematic <br> review, searches were undertaken as normal. |
| Moderate | Fully applicable | Details of included studies were used instead of undertaking a <br> new literature search. Full-text papers of included studies were <br> still retrieved for the purposes of data analysis. Searches were <br> only done to cover the period of time since the search date of <br> the review. |
| Moderate | Partially applicable | Details of included studies were used instead of undertaking a <br> new literature search for the relevant subsection of the protocol. |
| For this section, searches were only done to cover the period of |  |  |
| time since the search date of the review. For other sections not |  |  |
| covered by the systematic review, searches were undertaken as |  |  |
| normal. |  |  |

## Diagnostic test accuracy evidence

Diagnostic test accuracy (DTA) data are classified as any data in which a feature - be it a symptom, a risk factor, a test result or the output of some algorithm that combines many such features - is observed in some people who have the condition of interest at the time of the test and some people who do not. Such data either explicitly provide, or can be manipulated to generate, a $2 \times 2$ classification of true positives and false negatives (in people who, according to the reference standard, truly have the condition) and false positives and true negatives (in people who, according to the reference standard, do not).

The 'raw' $2 \times 2$ data can be summarised in a variety of ways. Those that were used for decision making in this guideline are as follows:

- Positive likelihood ratios describe how many times more likely positive features are in people with the condition compared to people without the condition. Values greater than 1 indicate that a positive result makes the condition more likely.
- $\mathrm{LR}^{+}=(\mathrm{TP} /[\mathrm{TP}+\mathrm{FN}]) /(\mathrm{FP} /[\mathrm{FP}+\mathrm{TN}])$
- Negative likelihood ratios describe how many times less likely negative features are in people with the condition compared to people without the condition. Values less than 1 indicate that a negative result makes the condition less likely.
- $\mathrm{LR}{ }^{-}=(\mathrm{FN} /[\mathrm{TP}+\mathrm{FN}]) /(\mathrm{TN} /[\mathrm{FP}+\mathrm{TN}])$
- Sensitivity is the probability that the feature will be positive in a person with the condition.
- sensitivity $=$ TP/(TP+FN)
- Specificity is the probability that the feature will be negative in a person without the condition.
- specificity $=$ TN/(FP+TN)
- Positive predictive values describe the probability that a person with a positive screening test has the disease.
- PPV = TP/ (TP+FP)
- Negative predictive values describe probability that a person with a negative screening test doesn't have the disease.
- NPV $=T N /(T N+F N)$

The following schema, adapted from the suggestions of Jaeschke et al. (1994), was used to interpret the likelihood ratio findings from diagnostic test accuracy reviews.

Table 7: Interpretation of likelihood ratios

| Value of likelihood ratio | Interpretation |
| :--- | :--- |
| $\mathrm{LR} \leq 0.1$ | Very large decrease in probability of disease |
| $0.1<\mathrm{LR} \leq 0.2$ | Large decrease in probability of disease |
| $0.2<\mathrm{LR} \leq 0.5$ | Moderate decrease in probability of disease |
| $0.5<\mathrm{LR} \leq 1.0$ | Slight decrease in probability of disease |
| $1.0<\mathrm{LR}<2.0$ | Slight increase in probability of disease |
| $2.0 \leq \mathrm{LR}<5.0$ | Moderate increase in probability of disease |
| $5.0 \leq \mathrm{LR}<10.0$ | Large increase in probability of disease |
| $\mathrm{LR} \geq 10.0$ | Very large increase in probability of disease |

The schema above has the effect of setting a minimal important difference for positive likelihoods ratio at 2, and a corresponding minimal important difference for negative likelihood ratios at 0.5 . Likelihood ratios (whether positive or negative) falling between these thresholds were judged to indicate no meaningful change in the probability of disease.

## Quality assessment

Individual studies were quality assessed using the QUADAS-2 tool, which contains four domains: patient selection, index test, reference standard, and flow and timing. Each individual study was classified into one of the following three groups:

- Low risk of bias - The true effect size for the study is likely to be close to the estimated effect size.
- Moderate risk of bias - There is a possibility the true effect size for the study is substantially different to the estimated effect size.
- High risk of bias - It is likely the true effect size for the study is substantially different to the estimated effect size.

Each individual study was also classified into one of three groups for directness, based on if there were concerns about the population, index features and/or reference standard in the study and how directly these variables could address the specified review question. Studies were rated as follows:

- Direct - No important deviations from the protocol in population, index feature and/or reference standard.
- Partially indirect - Important deviations from the protocol in one of the population, index feature and/or reference standard.
- Indirect - Important deviations from the protocol in at least two of the population, index feature and/or reference standard.


## Methods for combining diagnostic test accuracy evidence

Meta-analysis of diagnostic test accuracy data was conducted with reference to the Cochrane Handbook for Systematic Reviews of Diagnostic Test Accuracy (Deeks et al. 2010).

Where applicable, diagnostic syntheses were stratified by:

- Presenting symptomatology (features shared by all participants in the study, but not all people who could be considered for a diagnosis in clinical practice).
- The reference standard used for true diagnosis.

Where five or more studies were available for all included strata, a bivariate model was fitted using the mada package in R v3.4.0, which accounts for the correlations between positive and negative likelihood ratios, and between sensitivities and specificities. Where sufficient data were not available ( $2-4$ studies), separate independent pooling was performed for positive likelihood ratios, negative likelihood ratios, sensitivity and specificity, using Microsoft Excel. This approach is conservative as it is likely to somewhat underestimate test accuracy, due to failing to account for the correlation and trade-off between sensitivity and specificity (see Deeks 2010).

Random-effects models (der Simonian and Laird) were fitted for all syntheses, as recommended in the Cochrane Handbook for Systematic Reviews of Diagnostic Test Accuracy (Deeks et al. 2010).

In any meta-analyses where some (but not all) of the data came from studies at high risk of bias, a sensitivity analysis was conducted, excluding those studies from the analysis. Results from both the full and restricted meta-analyses are reported. Similarly, in any meta-analyses where some (but not all) of the data came from indirect studies, a sensitivity analysis was conducted, excluding those studies from the analysis.

## Modified GRADE for diagnostic test accuracy evidence

GRADE has not been developed for use with diagnostic studies; therefore a modified approach was applied using the GRADE framework.

The choice of primary outcome for decision making was determined by the committee and GRADE assessments were undertaken using the appropriate method from those listed below.

In all cases, following completion of the GRADE table, the downstream effects of these tests on patient- important outcomes were considered. This could be done explicitly during committee deliberations and reported as part of the discussion section of the review detailing the likely consequences of true positive, true negative, false positive and false negative test results. Alternatively, in reviews where a decision model is being carried (for example, as part of an economic analysis), these consequences may be incorporated here instead.

## Using likelihood ratios as the primary outcomes

GRADE assessments were only undertaken for positive and negative likelihood ratios, as the thresholds used to assess imprecision were based on these outcomes, but results for sensitivity and specificity are also presented alongside those data.

Evidence from diagnostic accuracy studies was initially rated as high-quality, and then downgraded according to the standard GRADE criteria (risk of bias, inconsistency, imprecision and indirectness) as detailed in Table 8 below.

The committee were consulted to set 2 clinical decision thresholds for each measure: the likelihood ratio above (or below for negative likelihood ratios) which a test would be recommended, and a second below (or above for negative likelihood ratios) which a test would be considered of no clinical use. These were used to judge imprecision (see below). If the committee were unsure which values to pick, then the default values of 2 for LR+ and 0.5 for LR- were used based on Error! Reference source not found., with the line of no effect a $s$ the second clinical decision line in both cases.

Table 8: Rationale for downgrading quality of evidence for diagnostic questions using likelihood ratio measures.

| GRADE criteria | Reasons for downgrading quality |
| :--- | :--- |
| Risk of bias | Not serious: If less than $33.3 \%$ of the weight in a meta-analysis came from <br> studies at moderate or high risk of bias, the overall outcome was not <br> downgraded. <br> Serious: If greater than $33.3 \%$ of the weight in a meta-analysis came from <br> studies at moderate or high risk of bias, the outcome was downgraded one <br> level. <br> Very serious: If greater than 33.3\% of the weight in a meta-analysis came from <br> studies at high risk of bias, the outcome was downgraded two levels. <br> Outcomes meeting the criteria for downgrading above were not downgraded if <br> there was evidence the effect size was not meaningfully different between <br> studies at high and low risk of bias. |
| Indirectness <br> Not serious: If less than 33.3\% of the weight in a meta-analysis came from <br> partially indirect or indirect studies, the overall outcome was not downgraded. <br> Serious: If greater than 33.3\% of the weight in a meta-analysis came from <br> partially indirect or indirect studies, the outcome was downgraded one level. <br> Very serious: If greater than 33.3\% of the weight in a meta-analysis came from <br> indirect studies, the outcome was downgraded two levels. |  |
| Outcomes meeting the criteria for downgrading above were not downgraded if <br> there was evidence the effect size was not meaningfully different between <br> direct and indirect studies. |  |


| GRADE criteria | Reasons for downgrading quality |
| :---: | :---: |
| Inconsistency | Concerns about inconsistency of effects across studies, occurring when there is unexplained variability in the treatment effect demonstrated across studies (heterogeneity), after appropriate pre-specified subgroup analyses have been conducted. This was assessed using the $I^{2}$ statistic. <br> N/A: Inconsistency was marked as not applicable if data on the outcome was only available from one study. <br> Not serious: If the $\mathrm{I}^{2}$ was less than $33.3 \%$, the outcome was not downgraded. Serious: If the $I^{2}$ was between $33.3 \%$ and $66.7 \%$, the outcome was downgraded one level. <br> Very serious: If the $\mathrm{I}^{2}$ was greater than $66.7 \%$, the outcome was downgraded two levels. <br> Outcomes meeting the criteria for downgrading above were not downgraded if there was evidence the effect size was not meaningfully different between studies with the smallest and largest effect sizes. |
| Imprecision | If the 95\% confidence interval for a positive likelihood ratio spanned a single LR+ clinical decision threshold (e.g. 2), the outcome was downgraded one level, as the data were deemed to be consistent with a meaningful increase in risk and no meaningful predictive value. Similarly, negative likelihood ratios that spanned a single LR- decision threshold (e.g. 0.5) led to downgrading for serious imprecision. Any likelihood ratios that spanned both the LR specific clinical decision threshold and the line of no effect were downgraded twice, as suffering from very serious imprecision. <br> Outcomes meeting the criteria for downgrading above were not downgraded if the confidence interval was sufficiently narrow that the upper and lower bounds would correspond to clinically equivalent scenarios. |

The quality of evidence for each outcome was upgraded if either of the following conditions were met:

- Data showed an effect size sufficiently large that it could not be explained by confounding alone.
- All plausible residual confounding is likely to increase our confidence in the effect estimate. Publication bias

Publication bias was assessed in two ways. First, if evidence of conducted but unpublished studies was identified during the review (e.g. conference abstracts or protocols without accompanying published data), available information on these unpublished studies was reported as part of the review. Secondly, where 10 or more studies were included as part of a single meta-analysis, a funnel plot was produced to graphically assess the potential for publication bias.

## Case series for diagnostic accuracy questions

For the purpose of this review 'diagnostic' case series are defined as studies reporting the proportion of participants with a diagnosis of the condition of interest (as defined by the reference standard) who had a positive test result (or who exhibited a particular sign or symptom).

## Quality assessment

The Joanna Briggs Institute checklists were used for diagnostic case series. Studies were assessed on the methods of participant recruitment, retention and outcome measurement (as appropriate), with each individual study classified into one of the following three groups:
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- Low risk of bias - The true result for the study is likely to be close to the estimated result
- Moderate risk of bias - There is a possibility the true result for the study is substantially different to the estimated result.
- High risk of bias - It is likely the true result for the study is substantially different to the estimated result.

Each individual study was also classified into one of three groups for directness, based on if there were concerns about the population or outcomes in the study and how directly these variables could address the specified review question. Studies were rated as follows:

- Direct - No important deviations from the protocol in population, intervention, comparator and/or outcomes.
- Partially indirect - Important deviations from the protocol in one of the population, intervention, comparator and/or outcomes.
- Indirect - Important deviations from the protocol in at least two of the population, intervention, comparator and/or outcomes.


## Methods for combining case series

Meta-analysis was considered inappropriate for the case series presented in this review due to the degree of expected heterogeneity between studies. No data was available for construction of a 2 by 2 table to enable a bivariate diagnostic accuracy meta-analysis to be performed.

## Modified GRADE for diagnostic case series

Applications of GRADE methodology has not been developed for use with non-comparative studies; therefore a modified approach was applied using the GRADE framework.

Evidence from case series was not combined in a meta-analysis, but given the large volume of data, it was considered helpful to group the data in the GRADE profile and provide an overall GRADE rating for each sign/symptom for each subgroup or sub analysis specified by the committee.

Consistent with the approach for diagnostic accuracy evidence, the quality of evidence was initially rated as 'high' and then downgraded as follows:

Table 9: Rationale for downgrading quality of evidence for diagnostic case series

| GRADE criteria | Reasons for downgrading quality |
| :--- | :--- |
| Risk of bias | Not serious: If less than $33.3 \%$ of the total participants came from studies at <br> moderate or high risk of bias <br> Serious: If greater than $33.3 \%$ of the total participants came from studies at <br> moderate or high risk of bias <br> Very serious: If greater than $33.3 \%$ of the total participants came from studies <br> at high risk of bias |
| Indirectness | Not serious: If less than $33.3 \%$ of the total participants came from partially <br> indirect or indirect studies <br> Serious: If greater than $33.3 \%$ of the total participants came from partially <br> indirect or indirect studies <br> Very serious: If greater than $33.3 \%$ of the total participants came from indirect <br> studies |


| GRADE criteria | Reasons for downgrading quality |
| :--- | :--- |
| Inconsistency | Concerns about inconsistency occurred when there is unexplained variability in <br> the effect demonstrated across studies (heterogeneity). |
|  | N/A: Inconsistency was marked as not applicable if data was only available <br> from one study. <br> Not serious: Confidence intervals across studies were overlapping <br> Serious: Confidence intervals across studies were not overlapping |
| Imprecision | Not serious: If $>33.3 \%$ of the studies had $>300$ participants <br> Serious: If $<33.3 \%$ of the studies had $>300$ participants <br> Very serious: If $<33.3 \%$ of the studies had $>100$ participants |

The quality of evidence for each outcome was upgraded if either of the following conditions were met:

- If an outcome was downgraded for risk of bias or indirectness, this outcome was upgraded if there was evidence the effect size was not meaningfully different between studies at high and low risk of bias, or between direct and indirect studies.
- If an outcome was downgraded for inconsistency, this outcome was upgraded if there was evidence the effect size was not meaningfully different between studies with the smallest and largest effect sizes.
- If an outcome was downgraded for imprecision, this outcome was upgraded if the confidence interval is sufficiently narrow that the upper and lower bounds would correspond to clinically equivalent scenarios.


## Publication bias

If evidence of conducted but unpublished studies was identified during the review (e.g. conference abstracts or protocols without accompanying published data), available information on these unpublished studies was reported as part of the review.

## Appendix C- Literature search strategies

A single systematic search was conducted for the question within this review on $16^{\text {th }}$ May 2019. The following databases were searched MEDLINE, MEDLINE in Process, MEDLINE e pub Ahead of print, and Embase, (all via the Ovid platform), the Cochrane Library databases (Wiley) and the DARE database (CRD platform).

The MEDLINE strategy is presented below. This was translated for the other databases.

1 Mucocutaneous Lymph Node Syndrome/
2 MCLS.tw.
3 (mucocutan* adj4 lymph*).tw.
4 kawasaki*.tw.
5 or/1-4
6 exp Infant/
7 (infan* or neonat* or newborn* or baby or babies).tw.
8 exp Child/
9 (child* or toddler* or boy* or girl*).tw.
10 (preschool* or pre-school*).tw.
11 exp Pediatrics/
12 (pediatric* or paediatric* or peadiatric*).tw.
13 or/6-12
14 exp "signs and symptoms"/ or Symptom Assessment/
15 (sign* or symptom* or complain* or indicator* or predict*).tw.
16 (clinical adj4 (manifestation* or feature* or finding* or aspect* or marker* or recogni* or identif*)).tw.
17 (presenting adj4 (feature* or finding* or factor* or aspect* or marker*)).tw.
18 presentation*.tw.
19 (physical adj4 (manifestation* or characteristic* or feature* or finding* or aspect* or marker*) ).tw.
20 or/14-19
21 (sensitiv: or predictive value:).mp. or accurac:.tw.
2220 or 21
235 and 13 and 22
24 animals/ not humans/
$25 \quad 23$ not 24
26 limit 25 to english language
27 limit 26 to ed=20070101-20190516

Searches to identify economic evidence were run on $17^{\text {th }}$ May 2019 in MEDLINE, MEDLINE in Process, MEDLINE e pub Ahead of print, Econlit and Embase (all via the Ovid platform), NHS EED and the Health Technology Assessment Database (via the CRD platform). NICE inhouse Economic Evaluation and Quality of Life filters were attached to lines 1 to 27 of the core strategy (lines 1 to 27 of the MEDLINE version shown above) in the MEDLINE and Embase databases. The MEDLINE version of the filters is displayed below.

```
28 Economics/
29 exp "Costs and Cost Analysis"/
30 Economics,Dental/
31 exp Economics, Hospital/
32 exp Economics, Medical/
33 Economics, Nursing/
34 Economics, Pharmaceutical/
35 Budgets/
36 exp Models, Economic/
37 Markov Chains/
38 Monte Carlo Method/
39 Decision Trees/
40 econom$.tw.
41 cba.tw.
42 cea.tw.
43 cua.tw.
44 markov$.tw.
45 (monte adj carlo).tw.
46 (decision adj3 (tree$ or analys$)).tw.
47 (cost or costs or costing$ or costly or costed).tw.
48 (price$ or pricing$).tw.
49 budget$.tw.
50 expenditure$.tw.
51 (value adj3 (money or monetary)).tw.
52 (pharmacoeconomic$ or (pharmaco adj economic$)).tw.
53 or/28-52
```

54 "Quality of Life"/
55 quality of life.tw.
56 "Value of Life"/
57 Quality-Adjusted Life Years/
58 quality adjusted life.tw.
59 (qaly\$ or qald\$ or qale\$ or qtime\$).tw.
60 disability adjusted life.tw.
61 daly\$.tw.
62 Health Status Indicators/
63 (sf36 or sf 36 or short form 36 or shortform 36 or sf thirtysix or sf thirty six or shortform thirtysix or shortform thirty six or short form thirtysix or short form thirty six).tw.
64 (sf6 or sf 6 or short form 6 or shortform 6 or sf six or sfsix or shortform six or short form six).tw.
65 (sf12 or sf 12 or short form 12 or shortform 12 or sf twelve or sftwelve or shortform twelve or short form twelve).tw.
66 (sf16 or sf 16 or short form 16 or shortform 16 or sf sixteen or sfsixteen or shortform sixteen or short form sixteen).tw.
67 (sf20 or sf 20 or short form 20 or shortform 20 or sf twenty or sftwenty or shortform twenty or short form twenty).tw.
68 (euroqol or euro qol or eq5d or eq 5d).tw.
69 (qol or hql or hqol or hrqol).tw.
70 (hye or hyes).tw.
71 health\$ year\$ equivalent\$.tw.
72 utilit\$.tw.
73 (hui or hui1 or hui2 or hui3).tw.
74 disutili\$.tw.
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75 rosser.tw.
76 quality of wellbeing.tw.
77 quality of well-being.tw.
78 qwb.tw.
79 willingness to pay.tw.
80 standard gamble\$.tw.
81 time trade off.tw.
82 time tradeoff.tw.
83 tto.tw.
84 or/54-83

## Appendix D - Clinical evidence study selection



66 included studies

## Appendix E - Clinical evidence tables

Advani 2019

| Advani 2019 |  |
| :---: | :---: |
| Bibliographic Reference | Advani, Najib; Santoso, Lucyana Alim; Sastroasmoro, Sudigdo; Profile of Kawasaki Disease in Adolescents: Is It Different? Acta medica Indonesiana; 2019; vol. 51 (no. 1); 42-46 |
| Study details |  |
| Study type | Patient records audit |
| Study details | Study location <br> Indonesia <br> Study setting <br> Hospital <br> Study dates <br> 2003 to 2016 <br> Exclusions <br> Hundreds of cases were excluded because of incomplete information <br> Sources of funding <br> Not mentioned. |
| Inclusion criteria | Typical and incomplete Kawasaki disease using the criteria as described by the AHA |
| Exclusion criteria | None or not meeting the inclusion criteria <br> Incomplete records or different sources of information that have conflicting information. |
| Sample characteristics | Sample size <br> 542 <br> \% Female <br> 39\% <br> Average age (variance) <br> Mean 27 months (SD 16) |
| Outcome(s) | Rates of occurrence of the principal criteria for Kawasaki disease according to the AHA |
| Study arm |  |
|  | Patients with Kawasaki disease ( $\mathrm{N}=542$ ) |

[^14]Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Joanna Briggs critical appraisal checklist for case series

Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
No
(Hundreds of patients were excluded because of incomplete data.)
Did the case series have complete inclusion of participants?
No
(Hundreds of patients were excluded because of incomplete data.)
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study. Hundreds of patients were excluded because of incomplete data.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Bai 2017

Bai 2017

| Bibliographic | Bai, L.; Feng, T.; Yang, L.; Zhang, Y.; Jiang, X.; Liao, J.; Chen, L.; Feng, X.; |
| :--- | :--- |
| Reference | Rong, Y.; Li, Y.; Qin, Z.; Qiao, J.; Retrospective analysis of risk factors |
| associated with Kawasaki disease in China; Oncotarget; 2017; vol. 8 (no. 33); |  |
|  | $54357-54363$ |

## Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> China |
| Study setting <br> Hospital <br> Study dates <br> 1998 to 2008 |  |
| Exclusions |  |

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| Study type | Patient records audit |
| :--- | :--- |
|  | None <br> Sources of funding <br> Life Consortium Corporation in Japan |
| Inclusion criteria | Diagnosis of Kawasaki disease according to the Japanese Circulation <br> Society Joint Working Group (Japan) |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 383 <br> \% Female <br> 40\% |
|  | Average age (variance) <br> "Incomplete KD patients' average age was $2.87 \pm 2.23$ and typical KD patients' average age was 3.01 <br> $\pm 2.35 . "$ |
| Clinical features |  |
| Outcome(s) |  |

Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=383$ )

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
This question has not yet been answered.
Were valid methods used for identification of the condition for all participants included in the case series?
This question has not yet been answered.
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study)

## Joanna Briggs critical appraisal checklist for case series

Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

Baker 2009
Baker 2009

| Bibliographic | Baker, Annette L.; Lu, Minmin; Minich, L. LuAnn; Atz, Andrew M.; Klein, Gloria L.; |
| :--- | :--- |
| Reference | Korsin, Rosalind; Lambert, Linda; Li, Jennifer S.; Mason, Wilbert; Radojewski, |
|  | Elizabeth; Vetter, Victoria L.; Newburger, Jane W.; Pediatric Heart Network, |
|  | Investigators; Associated symptoms in the ten days before diagnosis of Kawasaki |
|  | disease; The Journal of pediatrics; 2009; vol. 154 (no. 4); 592-595.e2 |

Study details

| Study type | Prospective cohort study |
| :---: | :---: |
| Study details | Study location <br> USA and Canada <br> Study setting <br> Hospital <br> Study dates <br> 2002 to 2004 <br> Exclusions <br> Sources of funding <br> National Institutes of Health, Ciarnanello Family Fund. |
| Inclusion criteria | Typical and incomplete Kawasaki disease using the criteria as described by the AHA |
| Exclusion criteria | None or not meeting the inclusion criteria Fever $>10$ days |
| Sample characteristics | Sample size <br> 198 <br> \% Female <br> Not provided <br> Average age (variance) <br> Mean approx 3.2 years (SD 2.2) |
| Outcome(s) | Rates of occurrence of associated symptoms and signs <br> Within 10 days prior to diagnosis |

## Study arm

Patients with Kawasaki disease $(\mathbf{N}=198)$

## Joanna Briggs critical appraisal checklist for case series <br> Were there clear criteria for inclusion in the case series?

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## Joanna Briggs critical appraisal checklist for case series

Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
Low
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Bal 2014

## Bal 2014

Bibliographic
Reference

Bal, Aswine K.; Prasad, Deepa; Umali Pamintuan, Maria Angela; MammenPrasad, Elizabeth; Petrova, Anna; Timing of intravenous immunoglobulin treatment and risk of coronary artery abnormalities in children with Kawasaki disease; Pediatrics and neonatology; 2014; vol. 55 (no. 5); 387-92

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> USA |
|  | Study setting <br> Hospital <br> Study dates <br> 1999 to 2011 |
|  | Exclusions <br> None <br> Sources of funding |


| Study type | Patient records audit |
| :--- | :--- |
| Inclusion criteria | Typical and incomplete Kawasaki disease using the criteria as described by <br> the AHA |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 106 |
| \% Female |  |
| 37\% |  |

## Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=106$ )

```
Joanna Briggs critical appraisal checklist for case series
Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case
series?
Yes
Were valid methods used for identification of the condition for all participants included in the case
series?
Yes
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable
```

FINAL

## Joanna Briggs critical appraisal checklist for case series

(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Behmadi 2019

Behmadi 2019

| Bibliographic | Behmadi, Maryam; Alizadeh, Behzad; Malek, Abdolreza; Comparison of Clinical |
| :--- | :--- |
| Reference | Symptoms and Cardiac Lesions in Children with Typical and Atypical Kawasaki |
| Disease; Medical sciences (Basel, Switzerland); 2019; vol. 7 (no. 4) |  |

## Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> Iran <br> Study setting <br> Hospital <br> Study dates <br> 2015 to 2018 <br> Exclusions <br> Some records were excluded because they were incomplete <br> Sources of funding <br> Research Vice Chancellor of Mashhad University of Medical Sciences |
| Inclusion criteria | Diagnosis of Kawasaki disease. No diagnostic criteria provided. |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 176 <br> \% Female <br> 36\% |
| Average age (variance) |  |
| Mean 32.43 (range 2-114) |  |

Study arm
Patients with Kawasaki disease $(\mathrm{N}=176)$

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
No
Was the condition measured in a standard, reliable way for all participants included in the case series?
No
(Diagnostic criteria not provided.)
Were valid methods used for identification of the condition for all participants included in the case series?

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```
Joanna Briggs critical appraisal checklist for case series
Yes
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
No
(Some records were excluded because they were incomplete.)
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital
admission. Some patients were over 5 years of age.)
```


## Boudiaf 2016

## Boudiaf 2016

| Bibliographic | Boudiaf, Houda; Achir, Moussa; The Clinical Profile of Kawasaki Disease in |
| :--- | :--- |
| Reference | Algerian Children: A Single Institution Experience; Journal of tropical pediatrics; |
|  | 2016; vol. 62 (no. 2); 139-43 |

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> Algeria <br> Study setting <br> Hospital |
|  | Study dates <br> 2005 to 2014 |
|  | Exclusions <br> None |
|  | Sources of funding <br> Not mentioned |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the <br> AHA |
| Exclusion criteria | None or not meeting the inclusion criteria |


| Study type | Patient records audit |
| :--- | :--- |
| Sample <br> characteristics | Sample size <br> 133 <br> \% Female <br> 38\% |
|  | Average age (variance) <br> Median 31 months (range 5 to 132 months) |
| Outcome(s) | Rates of occurrence of the principle criteria for Kawasaki disease according <br> to the AHA <br> Clinical features |

## Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=133$ )

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?

Yes
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(High (retrospective))
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Chang 2014

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## Chang 2014

Bibliographic Reference

Chang, Luan-Yin; Lu, Chun-Yi; Shao, Pei-Lan; Lee, Ping-Ing; Lin, Ming-Tai; Fan, Tsui-Yien; Cheng, Ai-Ling; Lee, Wan-Ling; Hu, Jen-Jan; Yeh, Shu-Jen; Chang, Chien-Chih; Chiang, Bor-Luen; Wu, Mei-Hwan; Huang, Li-Min; Viral infections associated with Kawasaki disease; Journal of the Formosan Medical Association = Taiwan yi zhi; 2014; vol. 113 (no. 3); 148-54

Study details

| Study type | Patient records audit |
| :---: | :---: |
| Study details | Study location Taiwan |
|  | Study setting Hospital |
|  | Study dates <br> 2004 to 2010 |
|  | Exclusions None |
|  | Sources of funding <br> Not mentioned |
| Inclusion criteria | Typical (complete) Kawasaki disease: fever for at least 5 days and at least 4 of the 5 principal criteria |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample characteristics | Sample size <br> 226 |
|  | $\begin{aligned} & \text { \% Female } \\ & 41 \% \end{aligned}$ |
|  | Average age (variance) |
|  | Mean 2.07 years (SD 1.76), Median 1.57 years (range 0.12 to 9.43 ) |
| Outcome(s) | Clinical features |

## Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=\mathbf{2 2 6}$ )

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Yes

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```
Joanna Briggs critical appraisal checklist for case series
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital
admission. Some patients were over 5 years of age.)
```


## Chen 2016

Chen 2016

| Bibliographic | Chen, J. J.; Ma, X. J.; Liu, F.; Yan, W. L.; Huang, M. R.; Huang, M.; Huang, G. Y.; |
| :--- | :--- |
| Reference | Epidemiologic features of Kawasaki disease in Shanghai from 2008 Through |
|  | 2012; Pediatric Infectious Disease Journal; 2016; vol. 35 (no. 1); 7-12 |

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> Taiwan <br> Study setting <br> Hospital |
|  | Study dates <br> 1997 to 2007 |
|  | Exclusions <br> None |
|  | Sources of funding <br> Not mentioned |
| Inclusion criteria | Diagnosis of Kawasaki disease. No diagnostic criteria provided. |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 351 |
| \% Female |  |
| Not provided |  |


| Study type | Patient records audit |
| :--- | :--- |
|  | Average age (variance) <br> Not provided |
| Outcome(s) | Clinical features |
| Study arm | Patients with Kawasaki disease $(\mathbf{N}=\mathbf{3 5 1})$ |

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Ebbeson 2004

## Ebbeson 2004

| Bibliographic | Ebbeson, Regan L.; Riley, Mark R.; Potts, Jim E.; Human, Derek G.; Malleson, |
| :--- | :--- |
| Reference | Peter N.; Kawasaki disease at British Columbia's Children's Hospital; Paediatrics |
|  | \& child health; 2004; vol. 9 (no. 7); 466-70 |

## Study details

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)
\(\left.$$
\begin{array}{|l|l|}\hline \text { Study type } & \text { Patient records audit } \\
\hline \text { Study details } & \begin{array}{l}\text { Study location } \\
\text { Canada }\end{array} \\
& \begin{array}{l}\text { Study setting } \\
\text { Hospital } \\
\text { Study dates } \\
\text { 1992 to 2000 }\end{array} \\
\hline & \begin{array}{l}\text { Exclusions } \\
\text { None }\end{array}
$$ <br>
\hline Sources of funding <br>

Not provided\end{array}\right]\)| Typical and atypical Kawasaki disease using the criteria as described by Han |
| :--- |
| et al. 2000 (Canada) |

## Study arm

Patients with Kawasaki disease $(\mathrm{N}=124)$

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Joanna Briggs critical appraisal checklist for case series

Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. )

## Fabi 2018

| Fabi 2018 |  |
| :--- | :--- |
| Bibliographic | Fabi, Marianna; Corinaldesi, Elena; Pierantoni, Luca; Mazzoni, Elisa; Landini, |
| Reference | Chiara; Bigucci, Barbara; Ancora, Gina; Malaigia, Laura; Bodnar, Tetyana; Di |
|  | Fazzio, Giorgia; Lami, Francesca; Valletta, Enrico; Cicero, Cristina; Biasucci, <br> Giacomo; Iughetti, Lorenzo; Marchetti, Federico; Sogno Valin, Paola; Amarri, |
|  | Sergio; Brusa, Sandra; Sprocati, Monica; Maggiore, Giuseppe; Dormi, Ada; <br> Lanzoni, Paolo; Donti, Andrea; Lanari, Marcello; Gastrointestinal presentation of <br> Kawasaki disease: A red flag for severe disease?; PloS one; 2018; vol. 13 (no. <br> 9); e0202658 |

## Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> Italy <br> Study setting <br> Hospitals |
|  | Study dates <br> 2000 to 2015 <br> Exclusions <br> None |
|  | Sources of funding <br> There was no specific funding for this study. |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the <br> AHA |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 302 <br> \% Female <br> 40\% |

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

FINAL

| Study type | Patient records audit |
| :--- | :--- |
|  | Average age (variance) <br> Of those who did not have abdominal manifestations: median 38.8 months (SD 31.6). Of those who <br> did: median 28.4 months (SD 31.7) |
| Outcome(s) | Rate of occurrence of abdominal manifestations |
|  | The following gastrointestinal manifestations were considered: vomiting, diarrhoea, abdominal pain, <br> abdominal distension, paralytic ileus, jaundice, pancreatitis and pseudo-obstruction. The presence of <br> vomiting and diarrhea was documented based on standard definition if reported by caregivers and/or <br> directly observed during the acute phase of the hospital stay. Abdominal pain was defined on physical <br> examination using a pain assessment scale appropriate for age [the Face, Legs, Activity, Cry, <br> Consolability scale (FLACC scale), Wong Baker FACES pain rating scale]. Paralytic ileus, obstruction, <br> jaundice and pancreatitis were clinically suspected and confirmed by imaging and laboratory findings, <br> when appropriate. They excluded patients with positive fecal cultures, anatomical malformations and <br> etiologies other than Kawasaki disease from the data analysis. |

Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=302$ )

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?

## Yes

Was the condition measured in a standard, reliable way for all participants included in the case series?

Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Study includes children over the age of 5 years.)

Falcini 2007

## Falcini 2007

Bibliographic Reference

Falcini, F.; Calabri, G. B.; Ricci, L.; Simonini, G.; Capannini, S.; Giani, T.; De Martino, M.; Update on Kawasaki disease: The 25 year experience at the "A. Mayer" Children's Hospital, Florence; Italian Journal of Pediatrics; 2007; vol. 33 (no. 1); 32-40

## Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> Hospital <br> Study setting <br> Italy <br> Study dates <br> 1980 to 2007 <br> Exclusions <br> Not mentioned <br> Sources of funding <br> Not mentioned |
| Inclusion criteria | Diagnosis of Kawasaki disease. No diagnostic criteria provided. <br> There were no diagnostic criteria mentioned in the methods section. |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 266 <br> \% Female <br> $39 \%$ |
| Autcome(s) | Average age (variance) <br> Median 26 months (range 2 to 293) |
| Rates of occurance of the principal criteria for Kawasaki disease |  |

## Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=\mathbf{2 6 6}$ )

```
Joanna Briggs critical appraisal checklist for case series
Were there clear criteria for inclusion in the case series?
No
(No diagnostic criteria mentioned in the methods section.)
Was the condition measured in a standard, reliable way for all participants included in the case
series?
Yes
Were valid methods used for identification of the condition for all participants included in the case
series?
Yes
Did the case series have consecutive inclusion of participants?
```

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

FINAL

```
Joanna Briggs critical appraisal checklist for case series
Unclear
Did the case series have complete inclusion of participants?
Unclear
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(No diagnostic criteria mentioned in the methods section. Retrospective study.)
Applicability as a source of data
Indirectly applicable
(Data on signs and symptoms was not collected at first presentation; data was collected during the
hospital stay. Some patients were over 5 years of age.)
```


## Falcini 2012

## Falcini 2012

Bibliographic
Reference

Falcini, Fernanda; Ozen, Seza; Magni-Manzoni, Silvia; Candelli, Marco; Ricci, Laura; Martini, Giorgia; Cuttica, Ruben J.; Oliveira, Sheila; Calabri, Giovanni Battista; Zulian, Francesco; Pistorio, Angela; La Torre, Francesco; Rigante, Donato; Discrimination between incomplete and atypical Kawasaki syndrome versus other febrile diseases in childhood: results from an international registrybased study; Clinical and experimental rheumatology; 2012; vol. 30 (no. 5); 799804

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> Turkey, Brazi and Italy |
|  | Study setting <br> Hospital |
| Study dates <br> 2005 to 2007 |  |
|  | Exclusions <br> There were 1466 Kawasaki, disease cases in the registry. 13 patients were excluded due to insufficient <br> data. 1,225 were excluded due to unexplained reason(s). <br> Sources of funding <br> Not mentioned. |
| Inclusion criteria | Diagnosis of Kawasaki disease. No diagnostic criteria provided. |

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

FINAL

| Study type | Patient records audit |
| :--- | :--- |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 228 <br> \% Female <br> 39\% <br> Average age (variance) <br> Typical KS, mean 29.6 months (SD 29). Atypical KD, mean 44.6 months (SD 38) |
| Clinical features |  |
| Rates of occurance of the principal criteria for Kawasaki disease |  |$|$| Study arm | Patients with Kawasaki disease (N=228) |
| :--- | :--- |

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
No
(There were 1466 Kawasaki disease cases in the registry. 13 patients were excluded due to insufficient data. 1,225 were excluded due to unexplained reason(s).)
Did the case series have complete inclusion of participants?
No
(13 patients were excluded due to insufficient data.)
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
No
(All the data from the 3 sites was grouped together.)
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Joanna Briggs critical appraisal checklist for case series

Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Gamez-Gonzalez 2013

## Gamez-Gonzalez 2013

| Bibliographic | Gamez-Gonzalez, Luisa Berenise; Murata, Chiharu; Munoz-Ramirez, Mireya; |
| :--- | :--- |
| Reference | Yamazaki-Nakashimada, Marco; Clinical manifestations associated with |
| Kawasaki disease shock syndrome in Mexican children; European journal of |  |
| pediatrics; 2013; vol. 172 (no. 3); 337-42 |  |

## Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> Mexico |
|  | Study setting <br> Hospital <br> Study dates <br> 2000 to 2012 <br> Exclusions <br> None |
| Sources of funding |  |
| Not mentioned |  |, | Typical and atypical Kawasaki disease using the criteria as described by the |
| :--- |
| AHA |

## Study arm

Patients with Kawasaki disease $(\mathbf{N}=212)$

```
Joanna Briggs critical appraisal checklist for case series
Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case
series?
Yes
```

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Joanna Briggs critical appraisal checklist for case series

Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Garrido-Garcia 2017

## Garrido-Garcia 2017

| Bibliographic | Garrido-Garcia, Luis Martin; Castillo-Moguel, Ariel; Vazquez-Rivera, Mirella; |
| :--- | :--- |
| Reference | Cravioto, Patricia; Fernando, Galvan; Reaction of the BCG Scar in the Acute |
|  | Phase of Kawasaki Disease in Mexican Children; The Pediatric infectious |
| disease journal; 2017; vol. 36 (no. 10); e237-e241 |  |

## Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> Mexico |
|  | Study setting <br> Hospital <br> Study dates <br> 1995 to 2015 |
|  | Exclusions <br> None |
|  | Sources of funding <br> Not mentioned |

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| Study type | Patient records audit |
| :--- | :--- |
| Inclusion criteria | Typical (complete) Kawasaki disease: fever for at least 5 days and at least 4 <br> of the 5 principal criteria as described by the AHA <br> Atypical (incomplete) Kawasaki disease: fever for at least 5 days and 2 to 3 <br> principal criteria + coronary involvement as described by the AHA |
| Exclusion criteria | No information about BCG status |
| Sample <br> characteristics | Sample size <br> 399 <br> \% Female <br> 35\% |
| Average age (variance) <br> BGC reaction: mean 19.09 months (range 3 to 131 months). No BCG reaction: mean 44.92 months <br> (range 2 to 200 months) |  |
| Rates of occurrence of the principal criteria for Kawasaki disease according <br> to the AHA <br> Rate of occurrence of BCG scar reaction |  |

## Study arm

$$
\text { Patients with Kawasaki disease ( } \mathrm{N}=399 \text { ) }
$$

```
Joanna Briggs critical appraisal checklist for case series
Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case
series?
No
(Retrospective study.)
Were valid methods used for identification of the condition for all participants included in the case
series?
Yes
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
No
(Some records did not have BCG vaccination status)
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
```

```
Joanna Briggs critical appraisal checklist for case series
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital
admission. Some patients were over 5 years of age.)
```


## Generini 1997

## Generini 1997

Bibliographic Reference

Generini, S.; Ermini, M.; Taccetti, G.; Trapani, S.; Cerinic, M. M.; Falcini, F.; Clinical and laboratory features and disease outcome of kawasaki disease: the analysis of our experience and literature review; Journal of clinical rheumatology : practical reports on rheumatic \& musculoskeletal diseases; 1997; vol. 3 (no. 5); 241-7

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> Italy <br> Study setting <br> Hospital <br> Study dates <br> 1980 to 1996 <br> Exclusions <br> None |
| Sources of funding |  |
| Not mentioned. |  |\(\left|\begin{array}{ll}Diagnosis of Kawasaki disease according to the Kawasaki Disease Research <br>

Committee (Japan)\end{array}\right|\)

## Study arm

Patients with Kawasaki disease ( $\mathrm{N}=73$ )

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Ghelani 2012

## Ghelani 2012

Bibliographic Reference

Ghelani, Sunil J.; Sable, Craig; Wiedermann, Bernhard L.; Spurney, Christopher F.; Increased incidence of incomplete Kawasaki disease at a pediatric hospital after publication of the 2004 American Heart Association guidelines; Pediatric cardiology; 2012; vol. 33 (no. 7); 1097-103

## Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> USA |
| Study setting <br> Hospital |  |
| Study dates <br> 2000 to 2002 and 2007 to 2009 <br> Exclusions |  |


| Study type | Patient records audit |
| :---: | :---: |
|  | Unknown number who had incomplete records or who did not receive intravenous gammaimmunoglobulin <br> Sources of funding <br> Not mentioned |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the AHA |
| Exclusion criteria | Incomplete records or different sources of information that have conflicting information. <br> Patients who did not receive intravenous gamma-immunoglobulin |
| Sample characteristics | Sample size <br> 203 <br> \% Female <br> 38\% <br> Average age (variance) <br> Median of earlier group 26 months (IQR 12.5 to 52 ). Median of later group 38.5 months (IQR 18-63) |
| Outcome(s) | Rates of occurrence of the principal criteria for Kawasaki disease according to the AHA |

## Study arm

Patients with Kawasaki disease $(\mathbf{N}=203)$

```
Joanna Briggs critical appraisal checklist for case series
Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case
series?
No
(AHA 2001 and 2004 guidelines were used.)
Were valid methods used for identification of the condition for all participants included in the case
series?
Yes
Did the case series have consecutive inclusion of participants?
No
(Some patients were excluded because of incomplete records and because they did not receive
standard treatment.)
Did the case series have complete inclusion of participants?
No
(Some patients were excluded because of incomplete records and because they did not receive
standard treatment.)
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
```

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Joanna Briggs critical appraisal checklist for case series

Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(High (retrospective). Study used a combination of the 2001 and 2004 AHA guidelines to diagnose KD. Some patients were excluded because of incomplete records and not receiving standard treatment.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Giannouli 2013

Giannouli 2013

| Bibliographic | Giannouli, Georgia; Tzoumaka-Bakoula, Chryssa; Kopsidas, Ioannis; |
| :--- | :--- |
| Reference | Papadogeorgou, Paraskevi; Chrousos, George P.; Michos, Athanasios; |
|  | Epidemiology and risk factors for coronary artery abnormalities in children with |
| complete and incomplete Kawasaki disease during a 10-year period; Pediatric |  |
| cardiology; 2013; vol. 34 (no. 6); 1476-81 |  |

## Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> Greece <br> Study setting <br> Hospital <br> Study dates <br> 2001 to 2010 |
|  | Exclusions <br> Incomplete data was included on 2 or 3 patients <br> Sources of funding <br> State Scholarship Foundation of Greece |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the <br> AHA |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample |  |
| characteristics | Sample size <br> 86 |
| \% Female |  |
| 39\% |  |

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

FINAL

| Study type | Patient records audit |
| :--- | :--- |
| Outcome(s) | Rates of occurrence of the principal criteria for Kawasaki disease according <br> to the AHA |
| Study arm | Patients with Kawasaki disease $\mathbf{( N = 8 6 )}$ |

```
Joanna Briggs critical appraisal checklist for case series
Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case
series?
Yes
Were valid methods used for identification of the condition for all participants included in the case
series?
Yes
Did the case series have consecutive inclusion of participants?
No
(9 were excluded because of missing data)
Did the case series have complete inclusion of participants?
No
(Incomplete data on 2 or 3 patients was used.)
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study. Incomplete data for 2 or 3 patients was used.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital
admission. Some patients were over 5 years of age.)
```


## Gorrab 2016

## Gorrab 2016

| Bibliographic | Gorrab, Arbia Abir; Fournier, Anne; Bouaziz, Asma Abed; Spigelblatt, Linda; |
| :--- | :--- |
| Reference | Scuccimarri, Rosie; Mrabet, Ali; Dahdah, Nagib; Incidence Rate and |
|  | Epidemiological and Clinical Aspects of Kawasaki Disease in Children of |
|  | Maghrebi Origin in the Province of Quebec, Canada, Compared to the Country of |
|  | Origin; Global pediatric health; 2016; vol. 3; 2333794x16630670 |

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> Canada (North African origin) <br> Study setting <br> Hospital <br> Study dates <br> 1996 to 2013 <br> Exclusions <br> None <br> Sources of funding <br> Not mentioned |
| Inclusion criteria | Diagnosis of Kawasaki disease. No diagnostic criteria provided. <br> North African origin |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 146 |
| \% Female |  |
| Not provided |  |$\quad$| Average age (variance) |
| :--- |
| Not provided |

## Study arm

## Patients with Kawasaki disease $(\mathrm{N}=146)$

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
No
(Diagnostic criteria was not provided)
Was the condition measured in a standard, reliable way for all participants included in the case series?
No
(Diagnostic criteria was not provided)
Were valid methods used for identification of the condition for all participants included in the case series?
No
(Diagnostic criteria was not provided)
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
Unclear
Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Joanna Briggs critical appraisal checklist for case series

Was there clear reporting of the demographics of the participants in the study?
No
(No demographic information)
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
No
(No demographic information)
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Diagnostic criteria not provided. No demographic data. Retrospective study.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Hu 2019

Hu 2019

| Bibliographic | Hu, Ya-Chiao; Liu, Hsin-Min; Lin, Ming-Tai; Chen, Chun-An; Chiu, Shuenn-Nan; |
| :--- | :--- |
| Reference | Lu, Chun-Wei; Chang, Luan-Yin; Wang, Jou-Kou; Wu, Mei-Hwan; Outcomes of |
|  | Kawasaki Disease Children With Spontaneous Defervescence Within 10 Days; |
|  | Frontiers in pediatrics; 2019; vol. 7; 158 |

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> Taiwan <br> Study setting <br> Hospital <br> Study dates <br> 2008 to 2015 |
|  | Exclusions <br> An unnown number of patients were excluded if they had received more than 1 dose of intravenous <br> gamma-immnoglobulin <br> Sources of funding <br> Not mentioned |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the <br> AHA |
| Exclusion criteria | None or not meeting the inclusion criteria <br> Patients receiving more than 1 dose of Intravenous gamma-immunoglobulin |

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

| Study type | Patient records audit |
| :--- | :--- |
|  | Patients lost to follow-up <br> Fever >10 days |
| Sample <br> characteristics | Sample size <br> 293 <br> \% Female <br> 45\% |
| Average age (variance) <br> Mean 1.8 years (SD 1.6) |  |
| Outcome(s) | Rates of occurrence of the principal criteria for Kawasaki disease according <br> to the AHA |

## Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=\mathbf{2 9 3 \text { ) }}$

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?

Yes
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable

## Joanna Briggs critical appraisal checklist for case series

(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were excluded if they received more than 1 dose of intravenous gammaimmunoglobulin)

## Huang 2006

## Huang 2006

Bibliographic Huang GY, Ma XJ, Huang M, et al. Epidemiologic pictures of Kawasaki disease Reference in Shanghai from 1998 through 2002. Journal of Epidemiology 2006;16(1):9-14.

## Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> Japan <br> Study setting <br> Hospital <br> Study dates <br> 1998 to 2002 |
| Exclusions <br> Cases were excluded if the questionnaire form was not completed correctly <br> Sources of funding <br> This study was supported by Japan Kawasaki Disease Research Center and Japan Monbu-kagakusho <br> Research Foundation |  |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the <br> Kawasaki Disease Research Committee in Japan |
| Exclusion criteria | None or not meeting the inclusion criteria <br> Cases were excluded if the questionnaire form was not completed correctly |
| Sample | Sample size <br> characteristics |
| \% Female |  |
| $35 \%$ |  |$\quad$| Average age (variance) |
| :--- |
| Mean 1.8 years (range 1 month to 18.8 years) |

## Study arm

Patients with Kawasaki disease ( $\mathrm{N}=768$ )

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Joanna Briggs critical appraisal checklist for case series

Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
Uncear
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Jaggi 2018

Jaggi 2018

| Bibliographic | Jaggi, Preeti; Grcic, Michelle; Kovalchin, John; Wilhelm, Carolyn M.; Yildirim- |
| :--- | :--- |
| Reference | Toruner, Cagri; Texter, Karen; Using the Electronic Medical Record to Correlate |
|  | Kawasaki Disease Phenotypes With Clinical Outcomes; Journal of the Pediatric |
|  | Infectious Diseases Society; 2018; vol. 7 (no. 2); 119-123 |

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> USA |
| Study setting <br> Hospital |  |
| Study dates <br> 2012 to 2015 <br> Exclusions <br> None <br> Sources of funding <br> Not mentioned |  |

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

| Study type | Patient records audit |
| :---: | :---: |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the AHA |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample characteristics | Sample size <br> 135 |
|  | \% Female Not provided |
|  | Average age (variance) |
|  | Typical group: median, 2.6 years (IQR 1.6 to 4.8 ). Atypical group with coronary artery abnormalities median 5.0 years (IQR 2.2 to 8.4). Atypical group without coronary artery abnormalities, median 3.6 years (IQR 1.9 to 5.4) |
| Outcome(s) | Rates of occurrence of the principal criteria for Kawasaki disease according to the AHA |
| Study arm |  |
|  | Patients with Kawasaki disease ( $\mathrm{N}=135$ ) |

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
Unclear
Was there clear reporting of the demographics of the participants in the study?
No
(No mention of gender)
Was there clear reporting of clinical information of the participants?
No
(Data on neck swelling by history is missing)
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High

```
Joanna Briggs critical appraisal checklist for case series
(Retrospective study. Data on history of neck swelling is absent.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital
admission.)
```


## Jun 2015

Jun 2015

| Bibliographic | Jun, Hyun Ok; Yu, Jeong Jin; Kang, So Yeon; Seo, Chang Deok; Baek, Jae Suk; |
| :--- | :--- |
| Reference | Kim, Young-Hwue; Ko, Jae-Kon; Diagnostic characteristics of supplemental |
| laboratory criteria for incomplete Kawasaki disease in children with complete |  |
|  | Kawasaki disease; Korean journal of pediatrics; 2015; vol. 58 (no. 10); 369-73 |

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> South Korea |
|  | Study setting <br> Hospital <br> Study dates <br> 2006 to 2012 |
|  | Exclusions <br> Following patients were excluded from the study: 19 patients who were transferred from other institutes <br> after initial IIG treatment, 64 patients showing incomplete presentation, 10 patients that were admitted <br> after fever lasting for more than 10 days and 63 patients in whom fever spontaneously subsided before <br> initial IVIG administration. |
| Sources of funding <br> Not mentioned |  |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the <br> AHA |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Fever >10 days |  |
| Patients who did not receive intravenous gamma-immunoglobulin |  |
| Patients transferred to other hospitals |  |

## Study arm

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Patients with Kawasaki disease ( $\mathrm{N}=355$ )

```
Joanna Briggs critical appraisal checklist for case series
Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case
series?
Yes
Were valid methods used for identification of the condition for all participants included in the case
series?
Yes
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
No
(Following patients were excluded from the study: }19\mathrm{ patients who were transferred from other
institutes after initial IVIG treatment, 64 patients showing incomplete presentation, 10 patients that
were admitted after fever lasting for more than }10\mathrm{ days and 63 patients in whom fever spontaneously
subsided before initial IVIG administration.)
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital
admission. Some patients were over 5 years of age.)
```


## Jun 2017

## Jun 2017

| Bibliographic | Jun, Woo Young; Ann, Yu Kyung; Kim, Ja Yeong; Son, Jae Sung; Kim, Soo-Jin; |
| :--- | :--- |
| Reference | Yang, Hyun Suk; Bae, Sun Hwan; Chung, Sochung; Kim, Kyo Sun; Kawasaki |
|  | Disease with Fever and Cervical Lymphadenopathy as the Sole Initial |
|  | Presentation; Korean circulation journal; 2017; vol. 47 (no. 1); 107-114 |

## Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location |

## Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

| Study type | Patient records audit |
| :--- | :--- |
|  | South Korea <br> Study setting <br> South Korea <br> Study dates <br> 2009 to 2013 <br> Exclusions <br> None <br> Sources of funding <br> Not mentioned |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the <br> AHA |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 146 |
| \% Female |  |
| 40\% |  |$\quad$| Average age (variance) |
| :--- |
| Those presenting with fever and lymphadenopathy only: mean 3.9 years (SD 2.3). Others: mean 2.4 |
| years (SD 1.7) |\(\left|\begin{array}{l}Rates of occurrence of the principal criteria for Kawasaki disease according <br>

to the AHA\end{array}\right|\)

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Joanna Briggs critical appraisal checklist for case series

Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Kil 2017

Kil 2017

| Bibliographic | Kil, Hong-Ryang; Yu, Jae-Won; Lee, Sung-Churl; Rhim, Jung-Woo; Lee, Kyung- <br> Yil; Changes in clinical and laboratory features of Kawasaki disease noted over <br> Reference |
| :--- | :--- |
|  | 1ime in Daejeon, Korea; Pediatric rheumatology online journal; 2017; vol. 15 (no. |

## Study details

| Study type | Patient records audit |
| :---: | :---: |
| Study details | Study location South Korea |
|  | Study setting Hospital |
|  | Study dates <br> 2000 to 2004 |
|  | Exclusions None |
|  | Sources of funding |
|  | There was no funding |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the AHA |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample characteristics | Sample size 615 |
|  | $\begin{aligned} & \text { \% Female } \\ & 38 \% \end{aligned}$ |
|  | Average age (variance) |
|  | Mean 29.7 months (SD 21.3) |
| Outcome(s) | Rates of occurrence of the principal criteria for Kawasaki disease according to the AHA |

## Study arm

[^15]
## Patients with Kawasaki disease ( $\mathrm{N}=\mathbf{6 1 5 \text { ) }}$

```
Joanna Briggs critical appraisal checklist for case series
Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case
series?
Yes
Were valid methods used for identification of the condition for all participants included in the case
series?
Yes
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital
admission. Some patients were over 5 years of age.)
```


## Kim 2017

Kim 2017

| Bibliographic | Kim, Gi Beom; Park, Sohee; Eun, Lucy Youngmin; Han, Ji Whan; Lee, Soo |
| :--- | :--- |
| Reference | Young; Yoon, Kyung Lim; Yu, Jeong Jin; Choi, Jong-Woon; Lee, Kyung-Yil; |
|  | Epidemiology and Clinical Features of Kawasaki Disease in South Korea, 2012- |
|  | $2014 ;$ The Pediatric infectious disease journal; 2017; vol. 36 (no. 5); 482-485 |

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> South Korea |
| Study setting <br> Hospital |  |

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

| Study type | Patient records audit |
| :--- | :--- |
|  | Study dates <br> 2012 to 2014 |
|  | Exclusions <br> None <br> Sources of funding <br> Not mentioned |
| Inclusion criteria | Diagnosis of Kawasaki disease. No diagnostic criteria provided. |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 14916 <br> \% Female <br> 42\% |
| Average age (variance) |  |
| Mean 32.9 months (SD 24.0) |  |

## Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=14916$ )

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
No
(Criteria were not mentioned.)
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Unclear
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
Unclear
(Result of a questionnaire sent to hospitals)
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?

## Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease

 FINAL (November 2019)```
Joanna Briggs critical appraisal checklist for case series
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital
admission. Some patients were over 5 years of age.)
```


## Kim 2018

## Kim 2018

Bibliographic Reference

Kim, S. H.; Lee, H. J.; Lee, J. S.; Clinical aspects of periungual desquamation in Kawasaki disease; Iranian Journal of Pediatrics; 2018; vol. 28 (no. 3); e59262

Study details
$\left.\left.\begin{array}{|l|l|}\hline \text { Study type } & \text { Patient records audit } \\ \hline \text { Study details } \\ & \begin{array}{l}\text { South Korea } \\ \text { Study setting } \\ \text { Hospital }\end{array} \\ & \begin{array}{l}\text { Study dates } \\ \text { 2011 to 2016 }\end{array} \\ \hline & \begin{array}{l}\text { Exclusions } \\ \text { Follow-up loss of 18 patients } \\ \text { Sources of funding }\end{array} \\ \hline \text { Not mentioned }\end{array} \right\rvert\, \begin{array}{ll}\text { Typical and atypical Kawasaki disease using the criteria as described by the } \\ \text { AHA }\end{array}\right]$

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=329$ )

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
No
(Follow-up loss of 18 patients.)
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Kim 2009

Kim 2009

| Bibliographic | Kim, Seong Hyun; Kim, Ki Hwan; Kim, Dong Soo; Clinical characteristics of <br> Reference |
| :--- | :--- |
| Kawasaki disease according to age at diagnosis; Indian pediatrics; 2009; vol. 46 <br> (no. 7); 585-90 |  |

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> South Korea |
| Study setting <br> Hospital |  |

[^16]| Study type | Patient records audit |
| :--- | :--- |
|  | Study dates <br> 2006 to 2007 <br> Exclusions <br> None <br> Sources of funding <br> No funding |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the <br> AHA <br> Atypical (incomplete) Kawasaki disease was defined as meeting less than 4 <br> clinical criteria, irrespective of echocardiography findings |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Samplecharacteristics Sample size <br> 153 <br> \% Female  <br> $45 \%$  |  |
| Average age (variance) |  |
| Mean age of 5 months and under group = 3.5 months. Mean age of 6 months to under 5 years group = |  |
| 27 months |  |

## Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=153$ )

```
Joanna Briggs critical appraisal checklist for case series
Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case
series?
Yes
Were valid methods used for identification of the condition for all participants included in the case
series?
Yes
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
```

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

```
Joanna Briggs critical appraisal checklist for case series
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital
admission.)
```

Kubota 2008

## Kubota 2008

Bibliographic Reference

Kubota, Masaru; Usami, Ikuya; Yamakawa, Masaru; Tomita, Yasuhiko; Haruta Tsunekazu; Kawasaki disease with lymphadenopathy and fever as sole initial manifestations; Journal of paediatrics and child health; 2008; vol. 44 (no. 6); 35962

Study details
$\left.\left.\begin{array}{|l|l|}\hline \text { Study type } & \text { Patient records audit } \\ \hline \text { Study details } & \begin{array}{l}\text { Study location } \\ \text { Japan } \\ \text { Study setting } \\ \text { Hospital } \\ \text { Study dates } \\ \text { 2000 to 2006 }\end{array} \\ \hline & \begin{array}{l}\text { Exclusions } \\ 8 \text { because this was a study about lymphadenopathy in Kawasaki disease and these patients had } \\ \text { concomitant conditions affecting lymph node size. }\end{array} \\ \hline \text { Sources of funding } \\ \text { Not mentioned }\end{array} \right\rvert\, \begin{array}{ll}\text { Diagnosis of Kawasaki disease according to the Kawasaki Disease Research } \\ \text { Committee (Japan) }\end{array}\right\}$

FINAL

| Study type | Patient records audit |
| :--- | :--- |
| Outcome(s) | Rates of occurrence of the principal criteria for Kawasaki disease according <br> to the Kawasaki Disease Research Committee (Japan) |
| Study arm | Patients with Kawasaki disease $\mathbf{( N = 1 3 6 )}$ |

```
Joanna Briggs critical appraisal checklist for case series
Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case
series?
Yes
Were valid methods used for identification of the condition for all participants included in the case
series?
Yes
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
No
(This was a study about lymphadenopathy in Kawasaki disease so concomitant conditions that
affected lymph gland size were excluded (8 patients).)
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital
admission. This was a study about cervical lymphadenopathy in Kawasaki disease so concomitant
conditions that affected lymph gland size were excluded. Study includes children over the age of 5
years.)
```


## Lee 2016

## Lee 2016

| Bibliographic | Lee, K. J.; Kim, H. J.; Kim, M. J.; Yoon, J. H.; Lee, E. J.; Lee, J. Y.; Oh, J. H.; |
| :--- | :--- |
| Reference | Lee, S. J.; Lee, K. Y.; Han, J. W.; Usefulness of anterior uveitis as an additional |

Lee 2016
tool for diagnosing incomplete Kawasaki disease; Korean Journal of Pediatrics; 2016; vol. 59 (no. 4); 174-177

Study details

| Study type | Prospective cohort study |
| :---: | :---: |
| Study details | Study location <br> Taiwan <br> Study setting <br> Hospital <br> Study dates <br> 1993 to 2008 <br> Exclusions <br> None <br> Sources of funding <br> Not provided |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the AHA |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample characteristics | Sample size <br> 145 <br> \% Female <br> 34\% <br> Average age (variance) <br> Not provided |
| Outcome(s) | Rates of occurrence of the principal criteria for Kawasaki disease according to the AHA <br> Rates of occurrence of associated symptoms and signs |

Study arm
Patients with Kawasaki disease ( $\mathrm{N}=145$ )

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
No
(Diagnostic criteria were not provided.)
Was the condition measured in a standard, reliable way for all participants included in the case series?
Unclear
(Diagnostic criteria were not provided.)
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

```
Joanna Briggs critical appraisal checklist for case series
Yes
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Diagnostic criteria were not provided.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital
admission. Some patients were over 5 years of age.)
```


## Li 2019

Li 2019

| Bibliographic | Li, Wei; Zhang, Li; Huang, Ping; Zhang, Zhiwei; Clinical features and mid-term |
| :--- | :--- |
| Reference | follow-up in infants younger than 3 months with Kawasaki disease in a Chinese |
| population; Journal of paediatrics and child health; 2019; vol. 55 (no. 5); 523-527 |  |

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> China <br> Study setting <br> Hospital <br> Study dates <br> 2009 to 2013 |
|  | Exclusions <br> None |
|  | Sources of funding <br> Not mentioned |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the <br> AHA |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 200 |

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

| Study type | Patient records audit |
| :---: | :---: |
|  | $\begin{aligned} & \text { \% Female } \\ & 31 \% \end{aligned}$ |
|  | Average age (variance) |
|  | $<3$ month old group, mean 2.4 months (range 1 to 3 months). >3 month old group, mean 25 months (range 4 to 108 months) |
| Outcome(s) | Rates of occurrence of the principal criteria for Kawasaki disease according to the AHA |
|  | Clinical features |

## Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=\mathbf{2 0 0 )}$

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable
(Data on signs and symptoms was not collected at first presentation; data was collected during the
hospital stay. Some patients were over 5 years of age.)
Liu 2012

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)


Study arm
Patients with Kawasaki disease ( $\mathrm{N}=145$ )

```
Joanna Briggs critical appraisal checklist for case series
Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case
series?
Yes
Were valid methods used for identification of the condition for all participants included in the case
series?
Yes
Did the case series have consecutive inclusion of participants?
Unclear
```

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

```
Joanna Briggs critical appraisal checklist for case series
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital
admission. Some patients were over 5 years of age.)
```


## Loh 2019

## Loh 2019

| Bibliographic | Loh, A. C. E.; Kua, P. H. J.; Tan, Z. L.; Erythema and induration of the bacillus <br> calmette-guerin site for diagnosing kawasaki disease; Singapore Medical Journal; <br> Reference |
| :--- | :--- |
| 2019; vol. 60 (no. 2); 89-93 |  |

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> Singapore <br> Study setting <br> Hospital <br> Study dates <br> 2007 to 2010 <br> Exclusions <br> None |
| Sources of funding |  |

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)
$\left.\begin{array}{|l|l|}\hline \text { Study type } & \text { Patient records audit } \\ \hline & \begin{array}{l}\text { \% Female } \\ 38 \%\end{array} \\ \text { Average age (variance) } \\ \text { Mean 1.6 years (SD 1.8) }\end{array}\right]$

```
Joanna Briggs critical appraisal checklist for case series
Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case
series?
Yes
Were valid methods used for identification of the condition for all participants included in the case
series?
Yes
Did the case series have consecutive inclusion of participants?
No
(Some children were excluded because details regarding any BCG vaccination was not included.)
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable
(Data on signs and symptoms was not collected at first presentation; data was collected during the
hospital stay.)
```


## QUADAS-2 checklist

```
Patient selection: risk of bias
```

Patient selection: risk of bias
Was a consecutive or random sample of patients enrolled?
Was a consecutive or random sample of patients enrolled?
No

```
No
```

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

Joanna Briggs critical appraisal checklist for case series
(Retrospective case-control design.)
Was a case-control design avoided?
No
Did the study avoid inappropriate exclusions?
Yes
Could the selection of patients have introduced bias?
High
Patient selection: applicability
Are there concerns that included patients do not match the review question?
Low
Index tests: risk of bias
Were the index test results interpreted without knowledge of the results of the reference
standard?
Yes
If a threshold was used, was it pre-specified?
Unclear
("A constellation of clinical features")
Could the conduct or interpretation of the index test have introduced bias?
Unclear
Index tests: applicability
Are there concerns that the index test, its conduct, or interpretation differ from the review question?
Unclear
(Lack of information available which details initial assessment.)
Reference standard: risk of bias
Is the reference standard likely to correctly classify the target condition?
Yes
Were the reference standard results interpreted without knowledge of the results of the index test?
Unclear
Could the reference standard, its conduct, or its interpretation have introduced bias?
Unclear
Reference standard: applicability
Is there concern that the target condition as defined by the reference standard does not match
the review question?
Low
Flow and timing: risk of bias
Was there an appropriate interval between index test(s) and reference standard?
Unclear
Did all patients receive a reference standard?
Yes
Did patients receive the same reference standard?
Yes
Were all patients included in the analysis?
Yes
Could the patient flow have introduced bias?
High
Overall risk of bias and directness

## Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

```
Joanna Briggs critical appraisal checklist for case series
Risk of Bias
High
Directness
Partially applicable
(Mixed population of KD and non-KD in control.)
```


## Manlhiot 2012

## Manlhiot 2012

Bibliographic Reference

Manlhiot, Cedric; Christie, Erin; McCrindle, Brian W.; Rosenberg, Hans; Chahal, Nita; Yeung, Rae S. M.; Complete and incomplete Kawasaki disease: two sides of the same coin; European journal of pediatrics; 2012; vol. 171 (no. 4); 657-62

Study details
\(\left.$$
\begin{array}{|l|l|}\hline \text { Study type } & \text { Patient records audit } \\
\hline \text { Study details } & \begin{array}{l}\text { Study location } \\
\text { Canada } \\
\text { Study setting } \\
\text { Hospital } \\
\text { Study dates } \\
1990 \text { to 2007 } \\
\text { Exclusions } \\
\text { None }\end{array}
$$ <br>
\hline Sources of funding <br>

Arthritis Society Investigator Award and the CIBC World Markets Children's Miracle Foundation\end{array}\right]\)| Typical and atypical Kawasaki disease using the criteria as described by the |
| :--- | :--- |
| AHA |

## Study arm

## Joanna Briggs critical appraisal checklist for case series <br> Were there clear criteria for inclusion in the case series? <br> Yes

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Joanna Briggs critical appraisal checklist for case series

Was the condition measured in a standard, reliable way for all participants included in the case series?

Yes
Were valid methods used for identification of the condition for all participants included in the case series?

Yes
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

Maric 2015

## Maric 2015

Bibliographic Reference

Maric, Lorna Stemberger; Knezovic, Ivica; Papic, Neven; Mise, Branko; Roglic, Srdan; Markovinovic, Leo; Tesovic, Goran; Risk factors for coronary artery abnormalities in children with Kawasaki disease: a 10-year experience; Rheumatology international; 2015; vol. 35 (no. 6); 1053-8

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> Croatia |
| Study setting <br> Hospital <br> Study dates <br> 2003 to 2012 |  |

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

| Study type | Patient records audit |
| :--- | :--- |
|  | Exclusions <br> Six patients were excluded from the study due to incomplete medical records. <br> Sources of funding <br> Not mentioned |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the <br> AHA |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 111 <br> \% Female <br> 37\% |
|  | Average age (variance) <br> Median 27 months (IQR 12 to 50) |
| Outcome(s) | Rates of occurrence of the principal criteria for Kawasaki disease according <br> to the AHA <br> Rate of occurrence of BCG scar reaction |

## Study arm

Patients with Kawasaki disease ( $\mathrm{N}=111$ )

```
Joanna Briggs critical appraisal checklist for case series
Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case
series?
Yes
Were valid methods used for identification of the condition for all participants included in the case
series?
Yes
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
```


## Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease

 FINAL (November 2019)```
Joanna Briggs critical appraisal checklist for case series
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital
admission. Some patients were over 5 years of age.)
```


## Martins 2018

## Martins 2018

Bibliographic Reference

Martins, Andreia; Conde, Marta; Brito, Maria; Gouveia, Catarina; Arthritis in Kawasaki disease: A poorly recognised manifestation; Journal of paediatrics and child health; 2018; vol. 54 (no. 12); 1371-1374

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> Portugal <br> Study setting <br> Hospital <br> Study dates <br> 1998 to 2013 |
| Exclusions <br> None |  |
| Sources of funding |  |
| Not mentioned |  |$|$| Typical and atypical Kawasaki disease using the criteria as described by the |
| :--- |
| AHA |

## Study arm

$$
\text { Patients with Kawasaki disease }(\mathrm{N}=63)
$$

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Minich 2007

## Minich 2007

| Bibliographic | Baker, Annette L.; Lu, Minmin; Minich, L. LuAnn; Atz, Andrew M.; Klein, Gloria L.; |
| :--- | :--- |
| Reference | Korsin, Rosalind; Lambert, Linda; Li, Jennifer S.; Mason, Wilbert; Radojewski, |
|  | Elizabeth; Vetter, Victoria L.; Newburger, Jane W.; Pediatric Heart Network, |
|  | Investigators; Associated symptoms in the ten days before diagnosis of Kawasaki |
|  | disease; The Journal of pediatrics; 2009; vol. 154 (no. 4); 592-595.e2 |

Study details

| Study type | Prospective cohort study |
| :--- | :--- |
| Study details | Study location <br> Canada and USA |
| Study setting <br> Hospital |  |
| Study dates <br> 2002 to 2004 |  |

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

| Study type | Prospective cohort study |
| :--- | :--- |
|  | Exclusions <br> None or not mentioned. <br> Sources of funding <br> Not mentioned |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the <br> AHA |
| Exclusion criteria | None or not meeting the inclusion criteria <br> Patients who did not receive intravenous gamma-immunoglobulin |
| Sample <br> characteristics | Sample size <br> 562 |
| \% Female |  |
| 40\% |  |$\quad$| Average age (variance) |
| :--- |
| Median 3.6 years (SD 2.9) |, | Rates of occurrence of the principal criteria for Kawasaki disease according |
| :--- |
| to the AHA |

## Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=562$ )

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?

Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes

## Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease

 FINAL (November 2019)```
Joanna Briggs critical appraisal checklist for case series
Overall Bias and Directness
Overall Risk of Bias
Low
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital
admission. Some patients were over 5 years of age.)
```


## Moore 2014

## Moore 2014

Bibliographic Reference

Moore, Abigail; Harnden, Anthony; Mayon-White, Richard; Recognising Kawasaki disease in UK primary care: a descriptive study using the Clinical Practice Research Datalink; The British journal of general practice : the journal of the Royal College of General Practitioners; 2014; vol. 64 (no. 625); e477-83

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> UK |
|  | Study setting <br> Primary care <br> Study dates <br> 2007 to 2011 |
|  | Exclusions <br> 681 out of 755 children were excluded from the study because of missing data. <br> Sources of funding |
| Inclusion criteria | Children under the age of 12 years who had a diagnosis of Kawasaki disease <br> in the Clinical Practice Research Datalink and who had a hospital discharge <br> diagnosis of Kawasaki disease |
| Diagnosis of Kawasaki disease. No diagnostic criteria provided. |  |

## Study arm

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Patients with Kawasaki disease ( $\mathrm{N}=104$ )

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
No
(No diagnostic criteria for Kawasaki disease)
Was the condition measured in a standard, reliable way for all participants included in the case series?
No
(Retrospective study.)
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
No
(227 children were excluded because the records did not correlate. A further 424 were excluded because there was missing data on signs and symptoms.)
Did the case series have complete inclusion of participants?
No
(227 children were excluded because the records did not correlate. A further 424 were excluded because there was missing data on signs and symptoms.)
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Not applicable
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(681 out of 755 children were excluded from the study because of missing data. There was no reference standard for diagnosing Kawasaki disease.)
Applicability as a source of data
Partially applicable
(Some patients were over 5 years of age.)

## Nomura 2012

Nomura 2012

Bibliographic Reference

Nomura, Yuichi; Arata, Michiko; Masuda, Kiminori; Koriyama, Chihaya; Suruki, Nobutaka; Ueno, Kentaro; Yoshikawa, Hideki; Eguchi, Taisuke; Kawano, Yoshifumi; Kawasaki disease patients with six principal symptoms have a high risk of being a non-responder; Pediatrics international : official journal of the Japan Pediatric Society; 2012; vol. 54 (no. 1); 14-8

## Study details

[^17]| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> Japan <br> Study setting <br> Hospital |
| Study dates <br> 2002 to 2009 |  |
| Exclusions <br> 20: These patients were not treated with IVIG or were admitted after 7 days of illmess. <br> Sources of funding |  |
| Inclusion criteria | Diagnosis of Kawasaki disease according to the Japanese Circulation <br> Society Joint Working Group (Japan) |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 207 <br> \% Female <br> 42\% |
| Average age (variance) |  |
| 5 symptom group, median 1.3 years (IQR 0.7 to 2.3). 6 symptom group, median 2.2 years (IQR 1.1 to |  |
| 3.8) |  |

## Study arm

$$
\text { Patients with Kawasaki disease }(\mathrm{N}=207)
$$

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?

Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
No
(20: These patients were not treated with IVIG or were admitted after 7 days of illness.)
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Joanna Briggs critical appraisal checklist for case series

Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study. 20 patients were omitted. These patients were not treated with IVIG or were admitted after 7 days of illness.)
Applicability as a source of data Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

Patel 2013

| Patel 2013 | Patel, Amy; Holman, Robert C.; Callinan, Laura S.; Sreenivasan, Nandini; |
| :--- | :--- |
| Bibliographic | Schonberger, Lawrence B.; Fischer, Thea K.; Belay, Ermias D.; Evaluation of <br> Reference <br> clinical characteristics of Kawasaki syndrome and risk factors for coronary artery <br> abnormalities among children in Denmark; Acta paediatrica (Oslo, Norway : <br> 1992); 2013; vol. 102 (no. 4); 385-90 |

Study details

| Study type | Patient records audit |
| :--- | :--- |
|  | Study location <br> Denmark <br> Study setting <br> Hospital <br> Study dates <br> 1994 to 2008 <br> Exclusions <br> 57 patients were excluded because of incomplete records. <br> Sources of funding <br> Funded through the Centers for Disease Control and Prevention. |
| Inclusion criteria | Discharge diagnosis of Kawasaki disease according to ICD-10 |

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

FINAL

| Study type | Patient records audit |
| :--- | :--- |
| Outcome(s) | Rates of occurrence of the principal criteria for Kawasaki disease |

## Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=314$ )

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Unclear
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
No
(57 patients were excluded because of incomplete records.)
Was there clear reporting of the demographics of the participants in the study?
No
(Demographic information for the signs and symptoms data is not provided.)
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
No
(No demographic information for the signs and symptoms data.)
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

Peng 2019

## Peng 2019

Bibliographic Reference

Peng, Yu; Liu, Xiaohui; Duan, Zhao; Deng, Yuhong; Cai, Sufen; Wang, Zhi; Xu, Kun; Kang, Hui; Jiang, Man; Li, Lin; Zhou, Yulan; Zou, Zheng; Prevalence and characteristics of arthritis in Kawasaki disease: a Chinese cohort study; Clinical and experimental medicine; 2019; vol. 19 (no. 2); 167-172

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> China <br> Study setting <br> Hospital <br> Study dates <br> 2014 to 2017 <br> Exclusions <br> 216 patients were excluded due to the following reasons: 56 patients received the initial treatment <br> before hospitalization, 11 patients were recurrent cases and the data were incomplete in 149 patients. <br> Sources of funding <br> Health Committee Foundation of Jiangxi Province, and Natural Science Foundation of Jiangxi Province |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the <br> AHA |
| Exclusion criteria | Incomplete records or different sources of information that have conflicting <br> information. |
| Initial treatment before hospitalisation |  |

## Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=1420$ )

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
No

## Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease

 FINAL (November 2019)```
Joanna Briggs critical appraisal checklist for case series
(There were many exclusions because of incomplete records.)
Did the case series have complete inclusion of participants?
No
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study. Many patients were excluded because of incomplete records.)
Applicability as a source of data
Indirectly applicable
(Data on signs and symptoms was not collected at first presentation; data was collected during the
hospital stay. Some patients were over 5 years of age.)
```


## Perrin 2009

## Perrin 2009

| Bibliographic | Perrin, Laurence; Letierce, Alexia; Guitton, Corinne; Tran, Tu-Anh; Lambert, |
| :--- | :--- |
| Reference | Virginie; Kone-Paut, Isabelle; Comparative study of complete versus incomplete |
|  | Kawasaki disease in 59 pediatric patients; Joint, bone, spine : revue du |
| rhumatisme; 2009; vol. 76 (no. 5); 481-5 |  |

## Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> France <br> Study setting <br> Hospital |
|  | Study dates <br> 1995 to 2006 |
|  | Exclusions <br> None |
|  | Sources of funding <br> Not mentioned |
| Inclusion criteria | Typical (complete) Kawasaki disease: fever for at least 5 days and at least 4 <br> of the 5 principal criteria as described by the AHA |
|  | Atypical (incomplete) Kawasaki disease: fever for at least 5 days and 1 to 3 <br> principal criteria as described by the AHA |

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

| Study type | Patient records audit |
| :--- | :--- |
|  | Exceptional cases: Typical (complete) Kawasaki disease: fever for less than <br> 5 days and at least 4 of the 5 principal criteria as described by the AHA + <br> coronary artery disease on echocardiography |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 59 <br> \% Female <br> 29 |
|  | Average age (variance) <br> 33 months (range: 2 to 169 months) |
| Outcome(s) | Rates of occurrence of the principal criteria for Kawasaki disease according <br> to the AHA |
| Study arm | Patients with Kawasaki disease (N=59) |

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
No
(This is a retrospective review of case records.)
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(This is a retrospective review of case records.)
Applicability as a source of data
Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Joanna Briggs critical appraisal checklist for case series

Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Piao 2010

Piao 2010

Bibliographic Reference

Piao, Jin-hua; Jin, Lian-hua; Lv, Jie; Zhou, Yan; Jin, Chun-ji; Jin, Zheng-yong; Epidemiological investigation of Kawasaki disease in Jilin province of China from 2000 to 2008; Cardiology in the young; 2010; vol. 20 (no. 4); 426-32

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> China <br> Study setting <br> Hospital <br> Study dates <br> 2000 to 2008 <br> Exclusions <br> None <br> Sources of funding <br> Not mentioned |
| Inclusion criteria | Diagnostic criteria for Kawasaki disease in the VIIth International Kawasaki <br> Disease Symposium (Japan) |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 735 |
| \% Female |  |
| 34\% |  |

## Study arm

Patients with Kawasaki disease ( $\mathrm{N}=735$ )

```
Joanna Briggs critical appraisal checklist for case series
Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case
series?
Yes
```

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

```
Joanna Briggs critical appraisal checklist for case series
Were valid methods used for identification of the condition for all participants included in the case
series?
Yes
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital
admission. Some patients were over 5 years of age.)
```


## Ruan 2013

## Ruan 2013

| Bibliographic | Ruan, Yu; Ye, Bei; Zhao, Xiaodong; Clinical characteristics of Kawasaki <br> syndrome and the risk factors for coronary artery lesions in China; The Pediatric <br> Reference |
| :--- | :--- |
| infectious disease journal; 2013; vol. 32 (no. 10); e397-402 |  |

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> China <br> Study setting <br> Hospital <br> Study dates <br> 2003 to 2009 |
|  | Exclusions <br> Four of these patients were excluded because of insufficient clinical or laboratory data. <br> Sources of funding |
| Not mentioned |  |


| Study type | Patient records audit |
| :--- | :--- |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 1209 <br> $\%$ Female <br> $36 \%$ |
|  | Average age (variance) <br> Not provided for the relevant <6 months age and over 6 months to 60 months ages groups. |
| Outcome(s) | Clinical features |

Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=1209$ )

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. )

## Sanchez-Maubens 2016

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Sanchez-Maubens 2016

| Bibliographic | Sanchez-Manubens, Judith; Anton, Jordi; Bou, Rosa; Iglesias, Estibaliz; Calzada- |
| :--- | :--- |
| Reference | Hernandez, Joan; Kawasaki Disease in Catalonia Working, Group; Incidence, <br> epidemiology and clinical features of Kawasaki disease in Catalonia, Spain; <br> Clinical and experimental rheumatology; 2016; vol. 34 (no. 3suppl97); S139-44 |

## Study details

| Study type | Patient records audit |
| :---: | :---: |
| Study details | Study location Spain <br> Study setting hospital <br> Study dates <br> 2004 to 2013 <br> Exclusions <br> None <br> Sources of funding <br> Not mentioned |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the AHA |
| Exclusion criteria | None or not meeting the inclusion criteria <br> Incomplete records or different sources of information that have conflicting information. <br> Over 16 years of age <br> Children admitted for a second opinion |
| Sample characteristics | Sample size <br> 399 <br> \% Female <br> 41\% <br> Average age (variance) <br> Mean 37 months (SD 33) |
| Outcome(s) | Rates of occurrence of all symptoms |

Study arm
Patients with Kawasaki disease ( $\mathrm{N}=399$ )

```
Joanna Briggs critical appraisal checklist for case series
Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case
series?
Yes
```

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Joanna Briggs critical appraisal checklist for case series

Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Sehgal 2015

## Sehgal 2015

| Bibliographic | Sehgal, Swati; Chen, Xinguang; Ang, Jocelyn Y.; Epidemiology, Clinical <br> Reference |
| :--- | :--- |
| Presentation, and Outcomes of Kawasaki Disease Among Hospitalized Children <br> in an Inner City Hospital Before and After Publication of the American Academy <br> of Pediatrics/American Heart Association Guidelines for Treatment of Kawasaki <br> Disease: An 11-Year Period; Clinical pediatrics; 2015; vol. 54 (no. 13); 1283-9 |  |

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location |
|  | USA |
|  | Study setting <br> Hospital |
|  | Study dates <br> 2000 to 2009 |
|  | Exclusions <br> None |
|  | Sources of funding <br> There was no funding. |

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)
\(\left.$$
\begin{array}{|l|l|}\hline \text { Study type } & \text { Patient records audit } \\
\hline \text { Inclusion criteria } & \begin{array}{l}\text { Typical and atypical Kawasaki disease using the criteria as described by the } \\
\text { AHA }\end{array} \\
\hline \text { Exclusion criteria } & \text { None or not meeting the inclusion criteria } \\
\hline \begin{array}{l}\text { Sample } \\
\text { characteristics }\end{array} & \begin{array}{l}\text { Sample size } \\
312 \\
\text { \% Female } \\
39 \%\end{array}
$$ <br>
\hline Average age (variance) <br>

Mean 42 months (SD 31)\end{array}\right]\)| Rates of occurrence of the principal criteria for Kawasaki disease according |
| :--- | :--- |
| to the AHA |$\quad$| Outcome(s) |
| :--- |

## Study arm

Patients with Kawasaki disease $(\mathrm{N}=312)$

```
Joanna Briggs critical appraisal checklist for case series
Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case
series?
Yes
Were valid methods used for identification of the condition for all participants included in the case
series?
Yes
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study)
Applicability as a source of data
Indirectly applicable
```

FINAL

## Joanna Briggs critical appraisal checklist for case series

(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Shamsizadeh 2014

## Shamsizadeh 2014

| Bibliographic | Shamsizadeh, Ahmad; Ziaei Kajbaf, Tahereh; Razavi, Maryam; Cheraghian, |
| :--- | :--- |
| Reference | Bahman; Clinical and epidemiological characteristics of kawasaki disease; <br> Jundishapur journal of microbiology; 2014; vol. 7 (no. 8); e11014 |

## Study details

| Study type | Patient records audit |
| :---: | :---: |
| Study details | Study location Iran |
|  | Study setting Hospital |
|  | Study dates 2000 to 2010 |
|  | Exclusions None |
|  | Sources of funding <br> Not mentioned |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the AHA |
|  | Kawasaki disease using the criteria as described by the American Academy of Pediatrics guideline. |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample characteristics | $\underset{104}{\text { Sample size }}$ |
|  | \% Female 37\% |
|  | Average age (variance) |
|  | Mean 33.6 months (SD 24.2), range: 3 months -8 years |
| Outcome(s) | Rates of occurrence of the principal criteria for Kawasaki disease according to the AHA |
|  | Clinical features |

## Study arm

Patients with Kawasaki disease ( $\mathrm{N}=104$ )

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes

## Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Joanna Briggs critical appraisal checklist for case series

Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Shiozawa 2014

## Shiozawa 2014

Bibliographic
Reference

Shiozawa, Yusuke; Inuzuka, Ryo; Harita, Yutaka; Kagawa, Jiro; Age-related differences in the course of the acute phase symptoms of Kawasaki disease; The Pediatric infectious disease journal; 2013; vol. 32 (no. 9); e365-9

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> Japan <br> Study setting <br> Hospital |
| Study dates <br> 2006 to 2012 |  |
|  | Exclusions <br> None |
|  | Sources of funding |

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

FINAL

| Study type | Patient records audit |
| :--- | :--- |
|  | This study was supported by institutional and departmental sources at the Department of Pediatrics, <br> Fujieda Municipal General Hospital, Japan. |
| Inclusion criteria | Diagnosis of Kawasaki disease according to the Kawasaki Disease Research <br> Committee (Japan) |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 100 <br> \% Female <br> 34\% |
| Average age (variance) <br> Median 24 months (IQR 10 to 53) |  |
| Outcome(s) | Rates of occurrence of the principal criteria for Kawasaki disease according <br> to the Kawasaki Disease Research Committee (Japan) |
| Study arm | Patients with Kawasaki disease (N=100) |

```
Joanna Briggs critical appraisal checklist for case series
Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case
series?
Yes
Were valid methods used for identification of the condition for all participants included in the case
series?
Yes
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
```

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Joanna Briggs critical appraisal checklist for case series

Partially applicable
(Some patients were over 5 years of age.)

## Sittiwangkul 2011

## Sittiwangkul 2011

| Bibliographic | Sittiwangkul, R.; Pongprot, Y.; Silvilairat, S.; Phornphutkul, C.; Delayed diagnosis <br> of Kawasaki disease: risk factors and outcome of treatment; Annals of tropical <br> Reference |
| :--- | :--- |
| paediatrics; 2011; vol. 31 (no. 2); 109-14 |  |

Study details

| Study type | Patient records audit |
| :---: | :---: |
| Study details | Study location Thailand |
|  | Study setting Hospital |
|  | Study dates <br> 2000 to 2008 |
|  | Exclusions None |
|  | Sources of funding |
|  | Not mentioned |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the AHA |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample characteristics | Sample size $170$ |
|  | \% Female <br> 40\% |
|  | Average age (variance) |
|  | Median 19 months (range 2 to 80) |
| Outcome(s) | Rates of occurrence of the principal criteria for Kawasaki disease according to the AHA <br> Clinical features |

Study arm
Patients with Kawasaki disease ( $\mathrm{N}=170$ )

```
Joanna Briggs critical appraisal checklist for case series
Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case
series?
Yes
```

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Joanna Briggs critical appraisal checklist for case series

Were valid methods used for identification of the condition for all participants included in the case series?

Yes
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study)
Applicability as a source of data
Indirectly applicable
(Some patients were over 5 years of age. Data on signs and symptoms was not collected at first presentation; data was collected during the hospital stay.)

## Sittiwangkul 2013

## Sittiwangkul 2013

| Bibliographic | Sittiwangkul, Rekwan; Pongprot, Yupada; Silvilairat, Suchaya; |
| :--- | :--- |
| Reference | Makonkaewkeyoon, Krit; Clinical spectrum of incomplete Kawasaki disease in |
|  | Thailand; Paediatrics and international child health; 2013; vol. 33 (no. 3); 176-80 |

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> Thailand <br> Study setting <br> Hospital - tertiary referral centre <br> Study dates <br> 2001 to 2009 <br> Exclusions |
|  | Sources of funding <br> Not mentioned |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the <br> AHA |

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

| Study type | Patient records audit |
| :--- | :--- |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 208 <br> \% Female <br> 37\% |
| Average age (variance) <br> Typical: mean 22.3 months (SD 15.3). Atypical: mean 22.4 months (SD 17.9) |  |
| Outcome(s) | Rates of occurrence of the principal criteria for Kawasaki disease according <br> to the AHA <br> Rates of occurrence of associated symptoms and signs |

## Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=\mathbf{2 0 8 )}$

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?

Yes
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
Unclear
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. This study was conducted at a tertiary referral centre.)
Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Sonobe 2007

Sonobe 2007

Bibliographic Reference

Sonobe, Tomoyoshi; Kiyosawa, Nobuyuki; Tsuchiya, Keiji; Aso, Seijiro; Imada, Yoshio; Imai, Yoko; Yashiro, Mayumi; Nakamura, Yoshikazu; Yanagawa, Hiroshi; Prevalence of coronary artery abnormality in incomplete Kawasaki disease; Pediatrics international : official journal of the Japan Pediatric Society; 2007; vol. 49 (no. 4); 421-6

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> Japan <br> Study setting <br> Hospital <br> Study dates <br> 2001 to 2002 <br> Exclusions <br> None mentioned <br> Sources of funding |
| Inclusion criteria | Diagnosis of Kawasaki disease. No diagnostic criteria provided. |$|$| Exclusion criteria | None or not meeting the inclusion criteria <br> Incomplete records or different sources of information that have conflicting <br> information. |
| :--- | :--- |
| Sample  <br> characteristics Sample size |  |
| \% Female |  |
| $43 \%$ |  |

## Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=15857$ )

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
No
(Diagnostic criteria was not provided.)
Was the condition measured in a standard, reliable way for all participants included in the case series?
Unclear
(Diagnostic criteria was not provided.)

## Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease

 FINAL (November 2019)```
Joanna Briggs critical appraisal checklist for case series
Were valid methods used for identification of the condition for all participants included in the case
series?
Unclear
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
Unclear
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study. Diagnostic criteria was not provided.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital
admission. Some patients were over 5 years of age.)
```


## Stemberger 2018

## Stemberger 2018

| Bibliographic | Stemberger Maric, Lorna; Papic, Neven; Sestan, Mario; Knezovic, Ivica; Tesovic, <br> Reference |
| :--- | :--- |
| Goran; Challenges in early diagnosis of Kawasaki disease in the pediatric <br> emergency department: differentiation from adenoviral and invasive <br> pneumococcal disease; Wiener klinische Wochenschrift; 2018; vol. 130 (no. 78); <br> $264-272$ |  |

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> Croatia <br> Study setting <br> Hospital <br> Study dates <br> 2006 to 2015 |
|  | Exclusions <br> None <br> Sources of funding <br> Not mentioned |

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

| Study type | Patient records audit |
| :--- | :--- |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the <br> AHA |
| Exclusion criteria | None or not meeting the inclusion criteria <br> Patients under 3 months of age <br> Patients over 3 years of age |
| Sample <br> characteristics | Sample size <br> 110 <br> \% Female <br> 35\% |
| Average age (variance) <br> Mean 16 months (SD 8.7 to 25.2) |  |
| Outcome(s) | Rates of occurrence of the principal criteria for Kawasaki disease according <br> to the AHA <br> Clinical features |

## Study arm

$$
\text { Patients with Kawasaki disease ( } \mathrm{N}=110 \text { ) }
$$

```
Joanna Briggs critical appraisal checklist for case series
Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case
series?
Yes
Were valid methods used for identification of the condition for all participants included in the case
series?
Yes
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
Unclear
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
```

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Joanna Briggs critical appraisal checklist for case series

High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable
(Data on signs and symptoms was not collected at first presentation; data was collected during the hospital stay. There was a relatively narrow window of recruitment -3 months to 3 years of age.)

Sun 2018

| Sun 2018 |  |
| :--- | :--- |
| Bibliographic <br> Reference | Sun, Ling; Tang, Yunjia; Wang, Ye; Qian, Guanghui; Yan, Wenhua; Wang, Bo; Li, <br> Xuan; Lv, Haitao; Changes in Profiles of Kawasaki Disease Noted over Time in <br> Suzhou, China; Cardiology; 2018; vol. 141 (no. 1); 25-31 |

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> China |
|  | Study setting <br> Hospital <br> Study dates <br> 2006 to 2017 <br> Exclusions <br> None <br> Sources of funding |
| Inclusion criteria | National Natural Science Foundation of China, Science and Technology Support Program of Jiangsu <br> Province and Science and Technology Projects for the Youth of Suzhou |
| AHA and atypical Kawasaki disease using the criteria as described by the |  |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 1008 |
| \% Female |  |
| 36\% |  |

## Study arm

$$
\text { Patients with Kawasaki disease }(\mathrm{N}=1008)
$$

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Joanna Briggs critical appraisal checklist for case series

Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
No
(Clinical details are missing for 896 patients, leaving data for 1008)
Did the case series have complete inclusion of participants?
No
(Clinical details are missing for 896 patients, leaving data for 1008)
Was there clear reporting of the demographics of the participants in the study?
No
(Demographic details of only the patients who have clinical data is not available.)
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
No
(Clinical details are missing for 896 patients, leaving data for 1008)
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study. Clinical details are missing for 896 patients, leaving data for 1008)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Tacke 2014

## Tacke 2014

| Bibliographic | Tacke, Carline E.; Breunis, Willemijn B.; Pereira, Rob Rodrigues; Breur, <br> Reference |
| :--- | :--- |
| Johannes M.; Kuipers, Irene M.; Kuijpers, Taco W.; Five years of Kawasaki <br> disease in the Netherlands: a national surveillance study; The Pediatric infectious <br> disease journal; 2014; vol. 33 (no. 8); 793-7 |  |

## Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> The Netherlands |
| Study setting <br> Hospital <br> Study dates <br> 2008 to 2012 |  |

[^18]| Study type | Patient records audit |
| :--- | :--- |
|  | Exclusions <br> Not all clinicians returned data. <br> Sources of funding <br> Stinafo Foundation |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the <br> AHA |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 319 <br> \% Female <br> 40\% |
|  | Average age (variance) <br> Median 2.4 years (range 0.1 to 14.6) |
| Outcome(s) | Rates of occurrence of the principal criteria for Kawasaki disease according <br> to the AHA <br> Clinical features |

## Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=319$ )

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?

Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
No
(Survey but not all clinicians returned data)
Did the case series have complete inclusion of participants?
No
(Survey but not all clinicians returned data)
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes

## Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease

 FINAL (November 2019)```
Joanna Briggs critical appraisal checklist for case series
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study. Survey but not all clinicians returned data.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital
admission. Some patients were over 5 years of age.)
```


## Tajima 2015

| Tajima 2015 | Bibliographic <br> Reference |
| :--- | :--- |
| Tajima, Miyu; Shiozawa, Yusuke; Kagawa, Jiro; Early Appearance of Principal <br> Symptoms of Kawasaki Disease is a Risk Factor for Intravenous Immunoglobulin <br> Resistance; Pediatric cardiology; 2015; vol. 36 (no. 6); 1159-65 |  |
| Study type | Patient records audit |
| Study details | Study location <br> Japan <br> Study setting <br> Hospital |
| Study dates <br> 2006 |  |
|  | Exc 2012 <br> None |
| Sources of funding |  |

## Study arm

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
Unclear
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study)
Applicability as a source of data
Indirectly applicable
(Data on signs and symptoms was not collected at first presentation; data was collected during the hospital stay. Some patients were over 5 years of age.)

## Tang 2016

Tang 2016

Bibliographic Reference

Tang, Yunjia; Gao, Xiang; Shen, Jie; Sun, Ling; Yan, Wenhua; Epidemiological and Clinical Characteristics of Kawasaki Disease and Factors Associated with Coronary Artery Abnormalities in East China: Nine Years Experience; Journal of tropical pediatrics; 2016; vol. 62 (no. 2); 86-93

## Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> China |
| Study setting <br> Hospital <br> Study dates <br> 2006 to 2014 |  |
| Exclusions |  |


| Study type | Patient records audit |
| :--- | :--- |
|  | Eleven auto-discharged cases and three cases with incomplete data were excluded. <br> Sources of funding <br> Chinese Natural Science Foundation, Jiangsu Province Science Foundation and Suzhou Science and <br> Technology Bureau. |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the <br> AHA |
| Exclusion criteria | None or not meeting the inclusion criteria <br> Sample <br> characteristics <br> Sample size <br> 1016 <br> \% Female <br> 36\% |
| Average age (variance) |  |
| Median 17 months (range 2 to 129) |  |

Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=1016$ )

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?

Yes
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)

## Joanna Briggs critical appraisal checklist for case series

Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Teng 2012

## Teng 2012

| Bibliographic | Teng, Mei-Chen; Wang, Li-Chieh; Yu, Hsin-Hui; Lee, Jyh-Hong; Yang, Yao-Hsu; |
| :--- | :--- |
| Reference | Chiang, Bor-Luen; Kawasaki disease and Henoch-Schonlein purpura - 10 years' <br> experience of childhood vasculitis at a university hospital in Taiwan; Journal of <br> microbiology, immunology, and infection = Wei mian yu gan ran za zhi; 2012; vol. <br>  <br> 45 (no. 1); 22-30 |

## Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> Taiwan <br> Study setting <br> Hospital <br> Study dates <br> 1997 to 2007 <br> Exclusions <br> None <br> Sources of funding |
| Inclusion criteria | Not mentioned |

## Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=351$ )

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
No
(Diagnostic criteria was not provided)

## Joanna Briggs critical appraisal checklist for case series

Was the condition measured in a standard, reliable way for all participants included in the case series?
Unclear
Were valid methods used for identification of the condition for all participants included in the case

## series?

Unclear
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
Unclear
Was there clear reporting of the demographics of the participants in the study?
No
(No demographic details)
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Unclear
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective and diagnostic criteria not provided.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

Tewelde 2014
Tewelde 2014

| Bibliographic | Tewelde, Helen; Yoon, Jeein; Van Ittersum, Wendy; Worley, Sarah; Preminger, |
| :--- | :--- |
| Reference | Tamar; Goldfarb, Johanna; The Harada score in the US population of children |
| with Kawasaki disease; Hospital pediatrics; 2014; vol. 4 (no. 4); 233-8 |  |

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> USA |
| Study setting <br> Hospital <br> Study dates <br> 2001 to 2011 |  |
| Exclusions <br> None |  |
| Sources of funding |  |

[^19]| Study type | Patient records audit |
| :--- | :--- |
| Inclusion criteria | No external funding <br> ICD-9 codes for "Kawasaki disease", and "Kawasaki disease and <br> "mucocutaneous lymph node syndrome". |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 105 <br> \% Female <br> 32\% <br> Average age (variance) <br> Median 2.8 years (Q1 1.6, Q3 5.5) |
| Outcome(s) | Rates of occurrence of the principal criteria for Kawasaki disease according <br> to ICD-9 |
| Study arm | Patients with Kawasaki disease (N=105) |

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?

Yes
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable
Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Joanna Briggs critical appraisal checklist for case series

(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

Uehara 2010
Uehara 2010

| Bibliographic | Uehara, Ritei; Igarashi, Hiroshi; Yashiro, Mayumi; Nakamura, Yosikazu; <br> Reference |
| :--- | :--- |
|  | Yanagawa, Hiroshi; Kawasaki disease patients with redness or crust formation at |
| the Bacille Calmette-Guerin inoculation site; The Pediatric infectious disease |  |
| journal; 2010; vol. 29 (no. 5); 430-3 |  |

Study details

| Study type | Patient records audit |
| :---: | :---: |
| Study details | Study location <br> Japan <br> Study setting <br> Hospital <br> Study dates <br> 2005 to 2006 <br> Exclusions <br> None mentioned <br> Sources of funding <br> Ministry of Health, Labour, and Welfare in Japan |
| Inclusion criteria | Diagnosis of Kawasaki disease according to the Kawasaki Disease Research Committee (Japan) <br> BCG vaccination |
| Exclusion criteria | None or not meeting the inclusion criteria No BCG vaccination |
| Sample characteristics | Sample size <br> 15524 <br> \% Female <br> Not provided <br> Average age (variance) <br> Not provided |
| Outcome(s) | Rate of occurrence of BCG scar reaction |

## Study arm

Patients with Kawasaki disease who had a BCG vaccination ( $\mathrm{N}=15524$ )

## Joanna Briggs critical appraisal checklist for case series <br> Were there clear criteria for inclusion in the case series? <br> Yes

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```
Joanna Briggs critical appraisal checklist for case series
Was the condition measured in a standard, reliable way for all participants included in the case
series?
Yes
Were valid methods used for identification of the condition for all participants included in the case
series?
Yes
Did the case series have consecutive inclusion of participants?
No
(Data was returned by 71% of hospitals)
Did the case series have complete inclusion of participants?
No
(Data was returned by 71% of hospitals)
Was there clear reporting of the demographics of the participants in the study?
No
(Data on mean age and gender was not provided.)
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study. Data was only returned by 71% of hospitals. Data on age and gender was not
provided.)
Applicability as a source of data
Indirectly applicable
(Data on signs and symptoms was not collected at first presentation; data was collected during the
hospital stay. Some patients were over 5 years of age.)
```

Wang 2009
Wang 2009

Bibliographic Reference

Wang, Susan; Best, Brookie M.; Burns, Jane C.; Periungual desquamation in patients with Kawasaki disease; The Pediatric infectious disease journal; 2009; vol. 28 (no. 6); 538-9

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> USA |
| Study setting <br> Hospital |  |


$\left.$| Study type | Patient records audit |
| :--- | :--- |
|  | Study dates <br> 2003 to 2007 |
|  | Exclusions <br> None <br> Sources of funding <br> Not mentioned |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the <br> AHA |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 243 |
| \% Female |  |
| Not provided. |  |$\quad$| Average age (variance) |
| :--- |
| Not provided. | \right\rvert\, | Rates of occurrence of the principal criteria for Kawasaki disease according |
| :--- |
| to the AHA |

## Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=\mathbf{2 4 3 \text { ) }}$

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
No
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Joanna Briggs critical appraisal checklist for case series

Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable
(Some patients may have been over 5 years of age. Perianal desquamation was measured one month after the fever started)

## Yellen 2010

## Yellen 2010

| Bibliographic | Yellen, Elizabeth S.; Gauvreau, Kimberlee; Takahashi, Masato; Burns, Jane C.; |
| :--- | :--- |
| Reference | Shulman, Stanford; Baker, Annette L.; Innocentini, Nancy; Zambetti, Chiara; |
|  | Pancheri, Joan M.; Ostrow, Adam; Frazer, Jeffrey R.; Sundel, Robert P.; Fulton, |
|  | David R.; Newburger, Jane W.; Performance of 2004 American Heart Association |
|  | recommendations for treatment of Kawasaki disease; Pediatrics; 2010; vol. 125 <br>  <br>  <br> (no. 2); e234-41 |

## Study details

| Study type | Patient records audit |
| :---: | :---: |
| Study details | Study location USA <br> Study setting <br> Hospital <br> Study dates <br> 1981 to 2006 <br> Exclusions <br> 53 were excluded because of incomplete records. |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the AHA |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample characteristics | Sample size <br> 195 <br> \% Female <br> 37\% <br> Average age (variance) <br> Median 2.1 years (range 0.1 to 19.4) |
| Outcome(s) | Rates of occurrence of the principal criteria for Kawasaki disease according to the AHA |

Study arm

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
No
(53 were excluded because of incomplete records.)
Did the case series have complete inclusion of participants?
No
(53 were excluded because of incomplete records.)
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study. 53 were excluded because of incomplete records.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Yoon 2016

## Yoon 2016

| Bibliographic | Yoon, You Min; Yun, Hye Won; Kim, Sung Hye; Clinical Characteristics of |
| :--- | :--- |
| Reference | Kawasaki Disease in Infants Younger than Six Months: A Single-Center Study; |
|  | Korean circulation journal; 2016; vol. 46 (no. 4); 550-5 |

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> South Korea |
| Study setting <br> Hospital |  |
| Study dates |  |


| Study type | Patient records audit <br> 2013 to 2015 |
| :--- | :--- |
|  | Exclusions <br> None |
| Sources of funding |  |
| Not mentioned |  |$|$

## Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=\mathbf{2 3 9 \text { ) }}$

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

```
Joanna Briggs critical appraisal checklist for case series
Overall Risk of Bias
High
(Retrospective study)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital
admission. Some patients were over 5 years of age.)
```

Yun 2011

| Yun 2011 |  |
| :--- | :--- |
| Bibliographic <br> Reference | Yun, Sang Hyun; Yang, Nu Ri; Park, Sin Ae; Associated symptoms of kawasaki <br> disease; Korean circulation journal; 2011; vol. 41 (no. 7); 394-8 |

Study details

| Study type | Patient records audit |
| :---: | :---: |
| Study details | Study location South Korea |
|  | Study setting Hospital |
|  | Study dates 2005 to 2010 |
|  | Exclusions None |
|  | Sources of funding |
|  | Not mentioned |
| Inclusion criteria | Typical and atypical Kawasaki disease using the criteria as described by the AHA |
|  | Which is the same as the Japanese Ministry of Heath and Welfare's criteria. |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample characteristics | Sample size <br> 121 |
|  | \% Female 36\% |
|  | Average age (variance) |
|  | Mean 31.8 months $+1-23.8$ |
| Outcome(s) | Rates of occurrence of associated symptoms and signs |

## Study arm

Patients with Kawasaki disease ( $\mathrm{N}=121$ )

## Joanna Briggs critical appraisal checklist for case series <br> Were there clear criteria for inclusion in the case series? <br> Yes

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Joanna Briggs critical appraisal checklist for case series

Was the condition measured in a standard, reliable way for all participants included in the case series?

Yes
Were valid methods used for identification of the condition for all participants included in the case series?

Yes
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study.)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Zhang 2016

## Zhang 2016

Bibliographic
Reference

Zhang, X.; Liang, Y.; Feng, W.; Su, X.; Zhu, H.; Epidemiologic survey of Kawasaki disease in Inner Mongolia, China, between 2001 and 2013;
Experimental and Therapeutic Medicine; 2016; vol. 12 (no. 2); 1220-1224

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> Mongolia |
|  | Study setting <br> Hospital |
|  | Study dates <br> 2001 to 2013 |
|  | Exclusions <br> None |
|  | Sources of funding <br> Not mentioned |
|  |  |

[^20]| Study type | Patient records audit |
| :--- | :--- |
| Inclusion criteria | Diagnosis of Kawasaki disease according to the Kawasaki Disease Research <br> Committee (Japan) |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 518 <br> \% Female <br> 38\% |
|  | Average age (variance) <br> Median 1.42 years (range 49 days to 14 years) |
| Outcome(s) | Rates of occurrence of the principal criteria for Kawasaki disease according <br> to the Kawasaki Disease Research Committee (Japan) <br> Clinical features |

Study arm
Patients with Kawasaki disease $(\mathrm{N}=518)$

```
Joanna Briggs critical appraisal checklist for case series
Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case
series?
Yes
Were valid methods used for identification of the condition for all participants included in the case
series?
Yes
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
No
(Some patients were excluded because of incomplete records.)
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study. Some patients were excluded because of incomplete records.)
```


## Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease

 FINAL (November 2019)FINAL

## Joanna Briggs critical appraisal checklist for case series

Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Zhang 2012

Zhang 2012

Bibliographic Reference

Zhang, X.; Zhang, Z.; Liu, S.; Sun, J.; Epidemiologic survey of kawasaki disease in Jilin from 1999 through 2008; Pediatric Cardiology; 2012; vol. 33 (no. 2); 272279

Study details

| Study type | Patient records audit |
| :---: | :---: |
| Study details | Study location China |
|  | Study setting Hospital |
|  | Study dates 1999 to 2008 |
|  | Exclusions None |
|  | Sources of funding |
|  | Not mentioned |
| Inclusion criteria | Diagnosis of Kawasaki disease according to the Japanese Circulation Society Joint Working Group (Japan) |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample characteristics | Sample size 577 |
|  | $\begin{aligned} & \text { \% Female } \\ & 34 \% \end{aligned}$ |
|  | Average age (variance) |
|  | Mean 2.67 years (SD 2.37) |
| Outcome(s) | Rates of occurrence of the principal criteria for Kawasaki disease according to the Kawasaki Disease Research Committee (Japan) |

## Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=577$ )

```
Joanna Briggs critical appraisal checklist for case series
Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case
series?
```

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Joanna Briggs critical appraisal checklist for case series

Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Yes
Did the case series have complete inclusion of participants?
Yes
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)

## Zhu 2015

Zhu 2015

Bibliographic Reference

Zhu, Hua; Yu, Shao-Fei; Bai, Yu-Xin; Liang, Yan-Yan; Su, Xue-Wen; Pan, JingYing; Kawasaki disease in children: Epidemiology, clinical symptoms and diagnostics of 231 cases in 10 years; Experimental and therapeutic medicine; 2015; vol. 10 (no. 1); 357-361

Study details

| Study type | Patient records audit |
| :--- | :--- |
| Study details | Study location <br> China |
| Study setting <br> China <br> Study dates <br> 2003 to 2012 |  |
| Exclusions <br> None |  |
| Sources of funding |  |
| Not mentioned |  |

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

| Study type | Patient records audit |
| :--- | :--- |
| Inclusion criteria | Diagnosis of Kawasaki disease according to the Kawasaki Disease Research <br> Committee (Japan) |
| Exclusion criteria | None or not meeting the inclusion criteria |
| Sample <br> characteristics | Sample size <br> 231 <br> \% Female <br> $32 \%$ |
|  | Average age (variance) <br> Age range 3 months to 10 years |
| Outcome(s) | Rates of occurance of the principal criteria for Kawasaki disease |

## Study arm

## Patients with Kawasaki disease ( $\mathrm{N}=\mathbf{2 3 1 \text { ) }}$

## Joanna Briggs critical appraisal checklist for case series

Were there clear criteria for inclusion in the case series?
Yes
Was the condition measured in a standard, reliable way for all participants included in the case series?
Yes
Were valid methods used for identification of the condition for all participants included in the case series?
Yes
Did the case series have consecutive inclusion of participants?
Unclear
Did the case series have complete inclusion of participants?
Unclear
Was there clear reporting of the demographics of the participants in the study?
Yes
Was there clear reporting of clinical information of the participants?
Yes
Were the outcomes or follow up results of cases clearly reported?
Yes
Was there clear reporting of the presenting site(s)/clinic(s) demographic information?
Yes
Was statistical analysis appropriate?
Yes
Overall Bias and Directness
Overall Risk of Bias
High
(Retrospective study)
Applicability as a source of data
Indirectly applicable
(Signs and symptoms were not collected at presentation; they were collected throughout hospital admission. Some patients were over 5 years of age.)
Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

FINAL
Signs and symptoms predicting Kawasaki disease

Fever under 5 s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

## Appendix F - GRADE profiles

## Case series

The 5 principal signs and symptoms (specified by the American heart association, AHA)
During course of illness: conjunctival injection

| Studies | Location | Age | Sample size | \% with Symptom (95\% CI) | Risk of bias | Indirectn ess | Inconsi stency | Impreci sion | Qualit <br> y |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Maric 2015 <br> Giannouli 2013 <br> Perrin 2009 | Europe (not UK) | All | $\begin{aligned} & 78 \\ & 62 \\ & 39 \end{aligned}$ | $\begin{aligned} & 92.8 \%(84.2 \text { to } 96.4) \\ & 90.3 \%(80.5 \text { to } 95.5) \\ & 93.3 \%(79.7 \text { to } 97.4) \\ & \text { Median } 92.8 \% \\ & \text { IQR (91.6 to } 93.1) \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Very serious ${ }^{3}$ | Very low |
| Manlhiot 2012 <br> Tang 2016 <br> Bai 2017 <br> Behmadi 2019 <br> Sonobe 2007 <br> Kil 2017 <br> Chang 2014 <br> Sittiwangkul 2013 <br> Tewelde 2014 <br> Ghelani 2012 <br> Yellen 2010 <br> Jaggi 2018 | Outside Europe | All | $\begin{aligned} & 728 \\ & 716 \\ & 298 \\ & 105 \\ & 13301 \\ & 387 \\ & 226 \\ & 147 \\ & 67 \\ & 127 \\ & 195 \\ & 105 \end{aligned}$ | $\begin{aligned} & 95.0 \%(93.2 \text { to } 96.3) \\ & 92.0 \%(89.8 \text { to } 93.8) \\ & 95.0 \%(91.9 \text { to } 96.9) \\ & 96.2 \%(90.6 \text { to } 98.5) \\ & 96.9 \%(96.6 \text { to } 97.2) \\ & 96.9 \%(94.7 \text { to } 98.2) \\ & 96.9 \%(93.8 \text { to } 98.5) \\ & 94.6 \%(89.6 \text { to } 97.2) \\ & 95.5 \%(87.6 \text { to } 98.5) \\ & 96.1 \%(91.1 \text { to } 98.3) \\ & 100.0 \%(97.2 \text { to } 100.0) \\ & 83.8 \%(75.6 \text { to } 89.6) \\ & \text { Median } 95.8 \% \\ & \text { IQR (94.9 to } 96.9) \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Maric 2015 <br> Giannouli 2013 <br> Perrin 2009 | Europe <br> (not UK) | All | $\begin{aligned} & 33 \\ & 22 \\ & 20 \end{aligned}$ | $\begin{aligned} & 78.8 \%(62.3 \text { to } 89.3) \\ & 27.3 \%(13.2 \text { to } 48.2) \\ & 65.0 \%(43.3 \text { to } 81.9) \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Very serious ${ }^{3}$ | Very low |


| Studies | Location | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectn ess | Inconsi stency | Impreci sion | Qualit <br> y |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | Median 65.0\% IQR (46.1 to 71.9) |  |  |  |  |  |
| Manlhiot 2012 <br> Tang 2016 <br> Bai 2017 <br> Behmadi 2019 <br> Sonobe 2007 <br> Lee 2016 <br> Kil 2017 <br> Sittiwangkul 2013 <br> Tewelde 2014 <br> Ghelani 2012 <br> Yellen 2010 <br> Jaggi 2018 | Outside <br> Europe | All | 217 300 85 71 2556 111 228 61 38 76 53 30 | 70.5\% (64.1 to 76.2) <br> 65.3\% (59.8 to 70.5) <br> 74.1\% (63.9 to 82.4) <br> 62.0\% (50.3 to 72.4) <br> 75.6\% (73.9 to 77.2) <br> 93.7\% (87.6 to 96.9) <br> 85.5\% (80.4 to 89.5) <br> 63.9\% (51.4 to 74.8) <br> 92.1\% (79.2 to 97.3) <br> 69.7\% (58.7 to 78.9) <br> 84.9\% (73.0 to 92.2) <br> 60.0\% (42.3 to 75.4) <br> Median 72.3\% <br> IQR (65.0 to 85.1) | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Stemberger 2018 <br> Patel 2013 <br> Generini 1997 <br> Falcini 2007 <br> Sanchez-Maubens 2016 <br> Tacke 2014 | Europe (not UK) | All | $\begin{aligned} & 110 \\ & 314 \\ & 73 \\ & 266 \\ & 399 \\ & 319 \end{aligned}$ | 65.5\% (56.2 to 73.7) <br> 94.6\% (91.5 to 96.6) <br> 68.5\% (57.1 to 78.0) <br> 97.4\% (94.7 to 98.7) <br> 79.7\% (75.5 to 83.4) <br> 87.1\% (83.0 to 90.4) <br> Median 83.4\% <br> IQR (71.3 to 92.1) | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Not serious | Very low |
| Ebbeson 2005 <br> Ruan 2013 <br> Li 2018 <br> Kim 2009 <br> Yoon 2016 <br> Teng 2012 <br> Liu 2012 | Outside Europe | $\begin{aligned} & <1 \\ & \text { year } \end{aligned}$ | $\begin{aligned} & 32 \\ & 49 \\ & 40 \\ & 22 \\ & 26 \\ & 109 \\ & 65 \end{aligned}$ | 87.5\% (71.9 to 95.0) 93.9\% (83.5 to 97.9$)^{6}$ $35.0 \%$ (22.1 to 50.5$)^{7}$ 68.2\% (47.3 to 83.6) ${ }^{8}$ 42.3\% (25.4 to 61.1) ${ }^{9}$ 84.4\% (76.4 to 90.0) 92.3\% (83.2 to 96.7) | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Very serious ${ }^{3}$ | Very low |


| Studies | Location | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectn ess | Inconsi stency | Impreci sion | Qualit <br> y |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | Median 84.4\% IQR (55.3 to 89.9) |  |  |  |  |  |
| Ebbeson 2005 <br> Ruan 2013 <br> Li 2018 <br> Kim 2009 <br> Yoon 2016 <br> Teng 2012 <br> Liu 2012 | Outside Europe | $\begin{aligned} & \geq 1 \\ & \text { year } \end{aligned}$ | $\begin{aligned} & 92 \\ & 1160 \\ & 160 \\ & 131 \\ & 213 \\ & 242 \\ & 80 \end{aligned}$ | 82.6\% (73.6 to 89.0) <br> 95.9\% (94.6 to 96.9) ${ }^{6}$ <br> 74.4\% (67.1 to 80.5) ${ }^{7}$ <br> $72.5 \%(64.3 \text { to } 79.4)^{8}$ <br> $80.3 \%(74.4 \text { to } 85.1)^{9}$ <br> 81.4\% (76.0 to 85.8) <br> 93.8\% (86.2 to 97.3) <br> Median 81.4\% <br> IQR (77.4 to 88.2) | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Boudiaf 2016 <br> Saundankar 2014 <br> Zhang 2016 <br> Zhang 2012 <br> Zhu 2015 <br> Chen 2016 <br> Sun 2018 <br> Peng 2019 <br> Advani 2019 <br> Shamsizadeh 2014 <br> Kubota 2008 <br> Piao 2010 <br> Nomura 2012 <br> Tajima 2015 <br> Garrido-Garcia 2017 <br> Gorrab 2016 <br> Kim 2017 <br> Jun 2017 <br> Jun 2015 <br> Kim 2018 <br> Hu 2019 | Outside Europe | All | $\begin{aligned} & 133 \\ & 353 \\ & 518 \\ & 577 \\ & 231 \\ & 2304 \\ & 1008 \\ & 1420 \\ & 542 \\ & 104 \\ & 136 \\ & 735 \\ & 207 \\ & 100 \\ & 399 \\ & 146 \\ & 14916 \\ & 146 \\ & 355 \\ & 292 \\ & 293 \end{aligned}$ | 91.0\% (84.9 to 94.8) 86.7\% (82.7 to 89.8) 68.5\% (64.4 to 72.4) 76.1\% (72.4 to 79.4) 66.2\% (59.9 to 72.0) 84.3\% (82.8 to 85.7) 89.1\% (87.0 to 90.9) 95.8\% (94.6 to 96.7) 85.2\% (85.2 to 88.0) 89.4\% (82.1 to 94.0) 96.3\% (91.7 to 98.4) $77.8 \%$ (74.7 to 80.7) 98.6\% (95.8 to 99.5) 98.0\% (93.0 to 99.5) 90.2\% (86.9 to 92.8) $79.5 \%$ (72.2 to 85.2) 88.6\% (88.1 to 89.1) 86.3\% (79.8 to 91.0) 97.7\% (95.6 to 98.9) 88.8\% (84.9 to 91.7) 91.8\% (88.1 to 94.4) | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |


| Studies | Location | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectn ess | Inconsi stency | Impreci <br> sion | Qualit <br> y |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Sittiwangkul 2011 <br> Falcini 2012 <br> LuAnn Minich 2007 <br> Wang 2009 <br> Bal 2014 <br> Sehgal 2015 <br> Huang 2006 ${ }^{10}$ |  |  | $\begin{aligned} & 170 \\ & 228 \\ & 562 \\ & 243 \\ & 106 \\ & 312 \\ & 768 \end{aligned}$ | 88.2\% (82.5 to 92.3) <br> 63.6\% (57.2 to 69.6) <br> 88.1\% (85.1 to 90.5) <br> 91.8\% (87.6 to 94.6) <br> 90.6\% (90.3 to 95.8) <br> 93.6\% (90.3 to 95.8) <br> 78.4\% (75.3 to 81.2) <br> Median 88.8\% <br> IQR (84.8 to 91.8) |  |  |  |  |  |
| 1. $>33.3 \%$ of partic <br> 2. $>33.3 \%$ participa <br> 3. $<33.3 \%$ of studie <br> 4. Confidence inter <br> 5. $<33.3 \%$ of studie <br> 6. Ruan 2013 grou <br> 7. Li 2018 groups d <br> 8. Kim 2009 groups <br> 9. Yoon 2016 group <br> 10. This study is one | udies at hi ies that we participants -overlapp participants ildren <6 en $<3$ mon dren $\leq 5 \mathrm{mo}$ ildren $\leq 6 \mathrm{~m}$ s that form | risk <br> indir <br> nths <br> s old <br> ths old <br> nths <br> $d$ the | as <br> and for child for childre d for child and for child is of the $p$ | en between 6 months over 3 months old between 5 months to on over 6 months old vious 2013 recommen | years old years old |  |  |  |  |

At presentation: conjunctival injection

| Studies | Locatio n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Moore 2014 <br> (Primary care) | UK | All | 74 | 31.1\% (21.7 to 42.3) <br> Median 31.1\% IQR - | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Shiozawa 2013 <br> (at time of presentation of principal symptoms other than fever) | Outside Europe | $\begin{aligned} & <2 \\ & \text { year } \end{aligned}$ | 51 | $\begin{aligned} & 45.1 \% ~(32.3 \text { to } \\ & 58.6)^{4} \\ & \text { Median } 45.1 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |


| Studies | Locatio n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Shiozawa 2013 <br> (at time of presentation of principal symptoms other than fever) | Outside Europe | $\begin{aligned} & \geq 2 \\ & \text { year } \end{aligned}$ | 49 | $16.3 \%(8.5 \text { to } 29.0)^{4}$ <br> Median 16.3\% IQR - | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Shiozawa 2013 (day 5 of fever) | Outside Europe | $\begin{aligned} & <2 \\ & \text { year } \end{aligned}$ | 51 | $\begin{aligned} & 84.3 \%(72.0 \text { to } \\ & 91.8)^{4} \\ & \text { Median } 84.3 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Shiozawa 2013 (day 5 of fever) | Outside Europe | $\begin{aligned} & \geq 2 \\ & \text { year } \end{aligned}$ | 49 | $85.7 \text { \% (73.3 to 92.9) }$ <br> Median 85.7\% IQR - | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |

1. $>33.3 \%$ of participants from studies at high risk of bias
2. $>33.3 \%$ participants from studies that were partially direct
3. $<33.3 \%$ of studies had $>100$ participants
4. Shiozawa 2013 groups data for children $\leq 2$ years old and for children over 2 years old

During course of illness: oral changes

| Studies | Locatio <br> n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Maric 2015 <br> Giannouli 2013 <br> Perrin 2009 | Europe (not UK) | All | $\begin{aligned} & 78 \\ & 62 \\ & 39 \end{aligned}$ | 94.9\% (87.5 to 98.0) <br> 98.4\% (91.4 to 99.7) <br> 100.0\% (91.0 to 100) <br> Median 98.4\% <br> IQR 96.7 to 99.2 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Very serious ${ }^{3}$ | Very low |
| Manlhiot 2012 <br> Bai 2017 <br> Behmadi 2019 <br> Sonobe 2007 <br> Lee 2016 <br> Kil 2017 | Outside Europe | All | $\begin{aligned} & 738 \\ & 298 \\ & 105 \\ & 13301 \\ & 111 \\ & 387 \end{aligned}$ | 96.2\% (94.6 to 97.4) 81.9\% (77.1 to 85.8) 98.1\% (93.3 to 99.5) 95.7\% (95.3 to 96.0) 63.1\% (53.8 to 71.5) 96.1\% (93.7 to 97.6) | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Not serious | Very low |


| Studies | Locatio <br> n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Chang 2014 <br> Sittiwangkul 2013 <br> Tewelde 2014 <br> Ghelani 2012 <br> Yellen 2010 <br> Jaggi 2018 |  |  | $\begin{aligned} & 226 \\ & 147 \\ & 67 \\ & 127 \\ & 137 \\ & 105 \end{aligned}$ | $\begin{aligned} & 95.1 \%(91.5 \text { to } 97.3) \\ & 100.0 \%(97.5 \text { to } 100) \\ & 98.5 \%(92.0 \text { to } 99.7) \\ & 91.3 \%(85.2 \text { to } 95.1) \\ & 97.1 \%(92.7 \text { to } 98.9) \\ & 77.1 \%(68.2 \text { to } 84.1) \\ & \text { Median } 95.9 \% \\ & \text { IQR } 89.0 \text { to } 97.4 \end{aligned}$ |  |  |  |  |  |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Maric 2015 <br> Giannouli 2013 <br> Perrin 2009 | Europe (not UK) | All | $\begin{aligned} & 33 \\ & 22 \\ & 20 \end{aligned}$ | 63.6\% ( 46.6 to 77.8 ) <br> 81.8\% (61.5 to 92.7) <br> 65.0\% (43.3 to 81.9) <br> Median 65.0\% <br> IQR 64.3 to 73.4 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Very serious ${ }^{3}$ | Very low |
| Manlhiot 2012 <br> Bai 2017 <br> Behmadi 2019 <br> Sonobe 2007 <br> Kil 2017 <br> Sittiwangkul 2013 <br> Tewelde 2014 <br> Ghelani 2012 <br> Yellen 2010 <br> Jaggi 2018 | Outside Europe | All | $\begin{aligned} & 217 \\ & 85 \\ & 71 \\ & 2556 \\ & 228 \\ & 61 \\ & 38 \\ & 76 \\ & 53 \\ & 30 \end{aligned}$ | 67.3\% (60.8 to 73.2) 54.1\% (43.6 to 64.3) $56.3 \%$ ( 44.8 to 67.3 ) 62.8\% (60.9 to 64.7) 63.6\% (57.2 to 69.6) $73.8 \%$ (61.6 to 83.2) $71.1 \%$ (55.2 to 83.0) 57.9\% (46.7 to 68.4) 67.9\% (54.5 to 78.9) 43.3\% (27.4 to 60.8) Median 63.2\% IQR 56.7 to 67.8 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Very serious ${ }^{3}$ | Very low |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Stemberger Maric 2018 <br> Patel 2013 <br> Generini 1997 <br> Falcini 2007 <br> Tacke 2014 | Europe (not UK) | All | $\begin{aligned} & 110 \\ & 314 \\ & 73 \\ & 266 \\ & 319 \end{aligned}$ | 38.2\% (29.7 to 47.5) <br> 97.8\% (95.5 to 98.9) <br> 94.5\% (86.7 to 97.9) <br> 96.6\% (93.7 to 98.2) <br> 87.5\% (83.4 to 90.7) | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Not serious | Very low |


| Studies | Locatio n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | Median 94.5\% IQR 87.5 to 96.6 |  |  |  |  |  |
| Ebbeson 2004 <br> Li 2018 <br> Kim 2009 <br> Yoon 2016 <br> Teng 2012 <br> Liu 2012 | Outside Europe | $\begin{aligned} & <1 \\ & \text { year } \end{aligned}$ | $\begin{aligned} & 32 \\ & 40 \\ & 22 \\ & 26 \\ & 109 \\ & 65 \end{aligned}$ | $\begin{aligned} & 62.5 \%(45.2 \text { to } 77.1) \\ & 40.0 \%(26.4 \text { to } \\ & 55.4)^{6} \\ & 86.4 \%(66.7 \text { to } \\ & 95.3)^{7} \\ & 42.3 \%(25.5 \text { to } \\ & 61.1)^{8} \\ & 67.9 \%(58.6 \text { to } 75.9) \\ & 89.2 \%(79.4 \text { to } 94.7) \\ & \text { Median } 65.2 \% \\ & \text { IQR } 47.4 \text { to } 81.8 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Very serious ${ }^{3}$ | Very low |
| Ebbeson 2004 <br> Li 2018 <br> Kim 2009 <br> Yoon 2016 <br> Teng 2012 <br> Liu 2012 | Outside Europe | $\begin{aligned} & \geq 1 \\ & \text { year } \end{aligned}$ | $\begin{aligned} & 92 \\ & 160 \\ & 131 \\ & 213 \\ & 242 \\ & 80 \end{aligned}$ | $\begin{aligned} & 76.1 \%(66.4 \text { to } 83.6) \\ & 77.5 \%(70.4 \text { to } \\ & 83.3)^{6} \\ & 78.6 \%(70.8 \text { to } \\ & 84.8)^{7} \\ & 76.5 \%(70.4 \text { to } \\ & 81.7)^{8} \\ & 73.6 \%(67.7 \text { to } 78.7) \\ & 91.3 \%(83.0 \text { to } 95.7) \\ & \text { Median } 77.0 \% \\ & \text { IQR } 76.2 \text { to } 78.3 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Boudiaf 2016 <br> Saundankar 2014 <br> Zhang 2016 <br> Zhang 2012 <br> Zhu 2015 <br> Chen 2016 <br> Peng 2019 <br> Advani 2019 <br> Shamsizadeh 2014 | Outside Europe | All | 133 <br> 353 <br> 518 <br> 577 <br> 231 <br> 2304 <br> 1420 <br> 542 <br> 104 | 97.7\% (93.6 to 99.2) <br> 94.6\% (91.8 to 96.5) <br> $73.9 \%$ (70.0 to 77.5) <br> 90.6\% (88.0 to 92.8) <br> 87.4\% (82.6 to 91.1) <br> 84.0\% (82.4 to 85.4) <br> 89.5\% (87.8 to 91.0) <br> 93.9\% (91.6 to 95.6) <br> 86.5\% (78.7 to 91.8) | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very Iow |



1. $>33.3 \%$ of participants from studies at high risk of bias
2. $>33.3 \%$ participants from studies that were indirect
3. $<33.3 \%$ of studies had $>100$ participants
4. Confidence intervals were non-overlapping
5. $<33.3 \%$ of studies had $>300$ participants
6. Li 2018 groups data for children $<3$ months old and for children over 3 months old
7. Kim 2009 groups data for children $\leq 5$ months old and for children between 5 months to $<5$ years old
8. Yoon 2016 groups data for children $\leq 6$ months old and for children over 6 months old
9. This study is one of two studies that formed the basis of the previous 2013 recommendations

At presentation: oral changes

| Studies | Locatio <br> n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Shiozawa 2013 (at time of presentation of principal symptoms other than fever) | Outside Europe | $\begin{aligned} & <1 \\ & \text { year } \end{aligned}$ | 51 | 29.4\% (18.7 to 43.0) <br> Median 29.4\% IQR - | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Shiozawa 2013 (at time of presentation of principal symptoms other than fever) | Outside Europe | $\geq 1$ <br> year | 49 | 12.2\% (5.73 to 24.2) <br> Median 12.2\% IQR - | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Shiozawa 2013 (day 5 of fever) | Outside Europe | $\begin{aligned} & <2 \\ & \text { year } \end{aligned}$ | 51 | $\begin{aligned} & 64.7 \% ~(51.0 \text { to } \\ & 76.4)^{4} \\ & \text { Median } 64.7 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Shiozawa 2013 (day 5 of fever) | Outside Europe | $\begin{aligned} & \geq 2 \\ & \text { year } \end{aligned}$ | 49 | $\begin{aligned} & 73.5 \% ~(59.7 \text { to } \\ & 83.8)^{4} \\ & \text { Median } 73.5 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |

1. $>33.3 \%$ of participants from studies at high risk of bias
2. $>33.3 \%$ participants from studies that were partially direct
3. $<33.3 \%$ of studies had $>100$ participants
4. Shiozawa 2013 groups data for children $\leq 2$ years old and for children over 2 years old

During course of illness: changes in the extremities

| Studies | Locatio $\mathrm{n}$ | Age | Sample <br> size | $\begin{aligned} & \text { \% with Symptom } \\ & (95 \% \mathrm{CI}) \end{aligned}$ | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Maric 2015 <br> Giannouli 2013 <br> Perrin 2009 | Europe (not UK) | All | $\begin{aligned} & 78 \\ & 64 \\ & 39 \end{aligned}$ | $\begin{aligned} & 78.2 \%(67.8 \text { to } 85.9) \\ & 65.6 \%(53.4 \text { to } 76.1) \\ & 97.4 \%(86.8 \text { to } 99.6) \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Very serious ${ }^{3}$ | Very low |


| Studies | Locatio <br> n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | Median 78.2\% IQR 71.9 to 87.8 |  |  |  |  |  |
| Manlhiot 2012 <br> Behmadi 2019 <br> Sonobe 2007 <br> Kil 2017 <br> Chang 2014 <br> Sittiwangkul 2013 <br> Telwelde 2014 <br> Ghelani 2012 <br> Yellen 2010 <br> Jaggi 2018 | Outside Europe | All | $\begin{aligned} & 738 \\ & 105 \\ & 13301 \\ & 387 \\ & 226 \\ & 147 \\ & 67 \\ & 127 \\ & 137 \\ & 105 \end{aligned}$ | 87.0\% (84.4 to 89.2) 67.6\% (58.2 to 75.8) 90.8\% (90.3 to 91.3) 93.8\% (90.9 to 95.8) $71.7 \%$ (65.5 to 77.2) 90.5\% (84.7 to 94.2) 92.5\% (83.7 to 96.8) 84.3\% (76.9 to 89.6) 88.3\% (81.9 to 92.7) 68.6\% (59.2 to 76.7) Median 87.7\% IQR 74.9 to 90.8) | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Maric 2015 <br> Giannouli 2013 <br> Perrin 2009 | Europe (not UK) | All | $\begin{aligned} & 33 \\ & 22 \\ & 20 \end{aligned}$ | $\begin{aligned} & 54.5 \%(38.0 \text { to } 70.2) \\ & 13.6 \%(4.75 \text { to } 33.3) \\ & 30.0 \%(13.6 \text { to } 51.9) \\ & \text { Median } 30.0 \% \\ & \text { IQR 21.8 to } 42.2 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Very serious ${ }^{3}$ | Very low |
| Manlhiot 2012 <br> Behmadi 2019 <br> Sonobe 2007 <br> Lee 2016 <br> Kil 2017 <br> Sittiwangkul 2013 <br> Falcini 2012 <br> Tewelde 2014 <br> Ghelani 2012 <br> Yellen 2010 <br> Jaggi 2018 | Outside Europe | All | $\begin{aligned} & 217 \\ & 71 \\ & 2556 \\ & 111 \\ & 228 \\ & 61 \\ & 228 \\ & 38 \\ & 76 \\ & 53 \\ & 30 \end{aligned}$ | 40.1\% (33.8 to 46.7) <br> 15.5\% (8.88 to 25.7) <br> 44.3\% (42.4 to 46.2) <br> 18.0\% (12.0 to 26.2) <br> 59.2\% (52.7 to 65.4) <br> 37.7\% (26.6 to 50.3) <br> 27.2\% (21.8 to 33.3) <br> 28.9\% (17.0 to 44.8) <br> 39.5\% (29.3 to 50.7) <br> 20.8\% (12.0 to 33.5) <br> 40.0\% (24.6 to 57.7) | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |


| Studies | Locatio <br> n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | Median 37.7\% IQR 24.0 to 40.1 |  |  |  |  |  |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Stemberger Maric 2018 <br> Patel 2013 <br> Generini 1997 <br> Falcini 2007 <br> Tacke 2014 | Europe (not UK) | All | $\begin{aligned} & 110 \\ & 269 \\ & 73 \\ & 266 \\ & 319 \end{aligned}$ | 26.4\% (19.0 to 35.3) <br> 85.7\% (81.4 to 89.1) <br> 89.0\% (79.8 to 94.3) <br> 82.0\% (76.9 to 86.1) <br> 77.4\% (72.5 to 81.7) <br> Median 82.0\% <br> IQR 77.4 to 85.7 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Serious ${ }^{5}$ | Very low |
| Ebbeson 2005 <br> Li 2018 <br> Kim 2009 <br> Yoon 2016 <br> Liu 2012 | Outside Europe | $<1$ <br> year | $\begin{aligned} & 32 \\ & 40 \\ & 22 \\ & 26 \\ & 65 \end{aligned}$ | $\begin{aligned} & 46.9 \%(30.9 \text { to } 63.6) \\ & 12.5 \%(5.46 \text { to } \\ & 26.1)^{6} \\ & 36.4 \%(19.7 \text { to } \\ & 57.1)^{7} \\ & 26.9 \%(13.7 \text { to } \\ & 46.1)^{8} \\ & 46.2 \%(34.6 \text { to } 58.2) \\ & \text { Median } 36.4 \% \\ & \text { IQR } 26.9 \text { to } 46.2 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Very serious ${ }^{3}$ | Very low |
| Ebbeson 2005 <br> Li 2018 <br> Kim 2009 <br> Yoon 2016 <br> Liu 2012 | Outside Europe | $\geq 1$ <br> year | $\begin{aligned} & 92 \\ & 160 \\ & 58 \\ & 213 \\ & 80 \end{aligned}$ | $\begin{aligned} & 80.4 \%(71.2 \text { to } 87.3) \\ & 73.8 \%(66.4 \text { to } \\ & 80.0)^{6} \\ & 44.3 \%(36.1 \text { to } \\ & 52.8)^{7} \\ & 61.5 \%(54.8 \text { to } \\ & 67.8)^{8} \\ & 50.0 \%(39.3 \text { to } 60.7) \\ & \text { Median } 61.5 \% \\ & \text { IQR 50.0 to } 73.8 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Boudiaf 2016 <br> Saundankar 2014 <br> Zhang 2016 | Outside Europe | All | $\begin{aligned} & 133 \\ & 353 \\ & 518 \end{aligned}$ | $\begin{aligned} & 96.2 \%(91.5 \text { to } 98.4) \\ & 72.8 \%(67.9 \text { to } 77.2) \\ & 62.2 \%(57.9 \text { to } 66.2) \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |


| Studies | Locatio <br> n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Zhang 2012 <br> Gorrab 2016 <br> Zhu 2015 <br> Chen 2016 <br> Peng 2019 <br> Advani 2019 <br> Shamsizadeh 2014 <br> Kubota 2008 <br> Normura 2012 <br> Tajima 2015 <br> Garrido-Garcia 2017 <br> Kim 2017 <br> Jun 2017 <br> Jun 2018 <br> Kim 2018 <br> Hu 2019 <br> Sittiwangkul 2011 <br> LuAnn Minich 2007 <br> Wang 2009 <br> Sehgal 2015 <br> Piao 2010 |  |  | 577 146 231 2304 1420 542 104 136 207 100 399 14916 146 355 329 293 170 562 243 312 735 | 84.9\% (81.8 to 87.6) 82.9\% (75.9 to 88.1) 61.0\% (54.6 to 67.1) 67.4\% (65.4 to 69.3) 48.9\% (46.3 to 51.5) 67.7\% (63.7 to 71.5) 66.3\% (56.8 to 74.7) 87.5\% (80.9 to 92.1) 93.7\% (89.6 to 96.3) 85.0\% (76.7 to 90.7) 71.4\% (66.8 to 75.6) 64.8\% (64.0 to 65.6) $75.3 \%$ (67.8 to 81.6) 93.0\% (89.8 to 95.2) 63.2\% (57.9 to 68.3) $72.4 \%$ (67.0 to 77.2) $75.3 \%$ (68.3 to 81.2) $77.0 \%$ (73.4 to 80.3) $79.8 \%$ (74.3 to 84.4) $72.4 \%$ (67.2 to 77.1) $35.2 \%$ (31.9 to 38.8) Median 72.6\% IQR 65.9 to 83.4 |  |  |  |  |  |
| 1. $>33.3 \%$ of particip <br> 2. $>33.3 \%$ participan <br> 3. $<33.3 \%$ of studies <br> 4. Confidence interv <br> 5. $<33.3 \%$ of studies <br> 6. Li 2018 groups da <br> 7. Kim 2009 groups <br> 8. Yoon 2016 group | from stu fom studie > $>100$ pa were non> $>300$ pa or children for childr ta for child | ies at that icipan verlap icipan <3 m n $\leq 5$ en $\leq 6$ | gh risk of re indirec <br> ng <br> hs old and nths old a onths old | ias <br> for children over 3 mo d for children between and for children over 6 | s old months onths old | <5 years old |  |  |  |

At presentation: changes in the extremities

| Studies | Locatio n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Shiozawa 2013 (at time of presentation of principal symptoms other than fever) | Outside Europe | $\begin{aligned} & <1 \\ & \text { year } \end{aligned}$ | 51 | $\begin{aligned} & 19.6 \% ~(11.0 \text { to } \\ & 32.5)^{4} \\ & \text { Median } 19.6 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Shiozawa 2013 (at time of presentation of principal symptoms other than fever) | Outside Europe | $\begin{aligned} & \geq 1 \\ & \text { year } \end{aligned}$ | 49 | $6.1 \%(2.10 \text { to } 16.5)^{4}$ <br> Median 6.1\% <br> IQR - | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Shiozawa 2013 (day 5 of fever) | Outside Europe | $\begin{aligned} & <2 \\ & \text { year } \end{aligned}$ | 51 | $\begin{aligned} & 62.7 \% ~(49.0 \text { to } \\ & 74.7)^{4} \\ & \text { Median } 62.7 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Shiozawa 2013 (day 5 of fever) | Outside Europe | $\begin{aligned} & \geq 2 \\ & \text { year } \end{aligned}$ | 49 | $\begin{aligned} & 61.2 \%(47.3 \text { to } \\ & 73.6)^{4} \\ & \text { Median } 61.2 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |

1. $>33.3 \%$ of participants from studies at high risk of bias
2. $>33.3 \%$ participants from studies that were partially direct
3. $<33.3 \%$ of studies had $>100$ participants
4. Shiozawa 2013 groups data for children $\leq 2$ years old and for children over 2 years old

## During course of illness: polymorphous rash

| Studies | Locatio n | Age | Sampl <br> e size | \% with Symptom $\text { ( } 95 \% \mathrm{CI} \text { ) }$ | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Maric 2015 <br> Giannouli 2013 | Europe (not UK) | All | $\begin{aligned} & 78 \\ & 50 \end{aligned}$ | $\begin{aligned} & 93.6 \%(85.9 \text { to } 97.2) \\ & 80.6 \%(69.2 \text { to } 88.6) \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Very serious ${ }^{3}$ | Very low |


| Studies | Locatio <br> n | Age | Sampl <br> e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Perrin 2009 |  |  | 39 | 94.9\% (83.1 to 98.6) <br> Median 93.6\% <br> IQR 87.1 to 94.3 |  |  |  |  |  |
| Manlhiot 2012 <br> Tang 2016 <br> Bai 2017 <br> Behmadi 2019 <br> Sonobe 2007 <br> Kil 2017 <br> Chang 2014 <br> Sittiwangkul 2013 <br> Tewelde 2014 <br> Ghelani 2012 <br> Yellen 2010 <br> Jaggi 2018 | Outside Europe | All | $\begin{aligned} & 738 \\ & 716 \\ & 298 \\ & 105 \\ & 13301 \\ & 387 \\ & 226 \\ & 147 \\ & 67 \\ & 127 \\ & 137 \\ & 105 \end{aligned}$ | 93.5\% (91.5 to 95.1) 90.4\% (88.0 to 92.3) 88.3\% (84.1 to 91.4) 87.6\% (80.0 to 92.6) 94.0\% (93.6 to 94.4) 86.3\% (82.5 to 89.4) 92.0\% (87.8 to 94.9) 98.0\% (94.2 to 99.3) 98.5\% (92.0 to 99.7) 96.9\% (92.2 to 98.8) 100.0\% (97.3 to 100) 89.5\% (82.2 to 94.1) Median 92.8\% IQR 89.2 to 97.2 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Maric 2015 <br> Giannouli 2013 <br> Perrin 2009 | Europe (not UK) | All | $\begin{aligned} & 33 \\ & 22 \\ & 20 \end{aligned}$ | $75.8 \%$ (59.0 to 87.2) <br> 63.6\% (43.0 to 80.3) <br> 65.0\% (43.3 to 81.9) <br> Median 65.0\% <br> IQR 64.3 to 70.4 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Very serious ${ }^{3}$ | Very low |
| Manlhiot 2012 <br> Tang 2016 <br> Bai 2017 <br> Behmadi 2019 <br> Sonobe 2007 <br> Lee 2016 <br> Kil 2017 <br> Sittiwangkul 2013 <br> Falcini 2012 | Outside Europe | All | $\begin{aligned} & 217 \\ & 300 \\ & 85 \\ & 71 \\ & 2556 \\ & 111 \\ & 228 \\ & 61 \\ & 288 \end{aligned}$ | 69.1\% (62.7 to 74.9) 44.3\% (38.8 to 50.0) 49.4\% (39.0 to 59.8) 45.1\% (34.1 to 56.6) 64.9\% (63.0 to 66.7) $67.6 \%$ ( 58.4 to 75.6 ) 24.6\% (19.4 to 30.5) 62.3\% (62.3 to 73.4) 67.5\% (61.2 to 73.3) | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |


| Studies | Locatio <br> n | Age | Sampl e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Tewelde 2014 |  |  | 38 | 73.7\% (58.0 to 85.0) |  |  |  |  |  |
| Ghelani 2012 |  |  | 76 | 68.4\% (57.3 to 77.8) |  |  |  |  |  |
| Yellen 2010 |  |  | 53 | 79.2\% (66.5 to 88.0) |  |  |  |  |  |
| Jaggi 2018 |  |  | 30 | 66.7\% (48.8 to 80.8) |  |  |  |  |  |
|  |  |  |  | Median 66.7\% <br> IQR 49.4 to 68.4 |  |  |  |  |  |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Stemberger Maric 2018 | Europe (not UK) | All | 110 | 74.5\% (65.7 to 81.8) | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Serious ${ }^{5}$ | Very low |
| Patel 2013 |  |  | 295 | 93.9\% (90.7 to 96.1) |  |  |  |  |  |
| Generini 1997 |  |  | 73 | 86.3\% (76.6 to 92.4) |  |  |  |  |  |
| Falcini 2007 |  |  | 266 | 99.6\% (97.9 to 99.9) |  |  |  |  |  |
| Sanchez-Manubens |  |  | 399 | 84.2\% (80.3 to 87.5) |  |  |  |  |  |
| 2017 |  |  | 319 | 89.7\% (85.8 to 92.5) |  |  |  |  |  |
| Tacke 2014 |  |  |  | $\begin{aligned} & \text { Median } 88.0 \% \\ & \text { IQR } 84.7 \text { to } 92.9 \end{aligned}$ |  |  |  |  |  |
| Ebbeson 2005 | Outside Europe | $\begin{aligned} & <1 \\ & \text { year } \end{aligned}$ | 32 | 87.5\% (71.9 to 95.0) | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Li 2018 |  |  | 40 | 42.5\% (28.5 to |  |  |  |  |  |
| Kim 2009 |  |  | 22 | $57.8)^{6}$ |  |  |  |  |  |
| Yoon 2016 |  |  | 26 | 63.6\% (43.0 to |  |  |  |  |  |
| Teng 2012 |  |  | 109 | 80.3) ${ }^{7}$ |  |  |  |  |  |
| Liu 2012 |  |  | 65 | $\begin{aligned} & 61.5 \% ~(42.5 \text { to } \\ & 77.6)^{8} \end{aligned}$ |  |  |  |  |  |
|  |  |  |  | 78.9\% (70.3 to 85.5) |  |  |  |  |  |
|  |  |  |  | 90.8\% (81.3 to 95.7) |  |  |  |  |  |
|  |  |  |  | Median 86.3\% IQR 74.5 to 89.7 |  |  |  |  |  |
| Ebbeson 2005 | Outside Europe | $\begin{aligned} & \geq 1 \\ & \text { year } \end{aligned}$ | 92 | 93.5\% (86.5 to 97.0) | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Li 2018 |  |  | 160 | 82.5\% (75.9 to |  |  |  |  |  |
| Kim 2009 |  |  | 131 | 87.6) ${ }^{6}$ |  |  |  |  |  |
| Yoon 2016 |  |  | 213 | 66.4\% (58.0 to |  |  |  |  |  |
| Teng 2012 |  |  | 242 | $73.9)^{7}$ |  |  |  |  |  |
| Liu 2012 |  |  | 80 | $\begin{aligned} & 78.4 \%(72.4 \text { to } \\ & 83.4)^{8} \end{aligned}$ |  |  |  |  |  |


| Studies | Locatio n | Age | Sampl <br> e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | $\begin{aligned} & 81.0 \% \text { ( } 75.6 \text { to } 85.4 \text { ) } \\ & 91.3 \%(83.0 \text { to } 95.7) \\ & \text { Median } 81.8 \% \\ & \text { IQR } 79.1 \text { to } 89.1 \end{aligned}$ |  |  |  |  |  |
| Boudiaf 2016 <br> Saundankar 2014 <br> Zhang 2016 <br> Zhang 2012 <br> Zhu 2015 <br> Chen 2016 <br> Peng 2019 <br> Advani 2019 <br> Shamizadeh 2014 <br> Kubota 2008 <br> Piao 2010 <br> Nomura 2012 <br> Tajima 2015 <br> Garrido-Garcia 2017 <br> Gorrab 2016 <br> Kim 2017 <br> Jun 2017 <br> Jun 2015 <br> Kim 2018 <br> Hu 2019 <br> Sittiwangkul 2011 <br> LuAnn Minich 2007 <br> Wang 2009 <br> Bal 2014 <br> Sehgal 2015 <br> Huang $2006^{9}$ | Outside Europe | All | 133 353 518 577 231 2304 1420 542 104 136 735 207 100 399 146 14916 146 355 329 293 170 562 243 106 312 768 | 97.7\% (93.6 to 99.2) 96.0\% (93.5 to 97.6) 64.5\% (60.3 to 68.5) 75.9\% (72.3 to 79.2) 57.6\% (51.1 to 63.8) $73.7 \%$ (71.9 to 75.5) 82.7\% (80.7 to 84.6) 86.5\% (83.4 to 89.2) $76.0 \%$ (66.9 to 83.2) 96.3\% (91.7 to 98.4) 72.2\% (68.9 to 75.4) 95.2\% (91.3 to 97.4) 94.0\% (87.5 to 97.2) 85.0\% (81.1 to 88.1) 91.1\% (85.4 to 94.7) 83.1\% (82.5 to 83.7) $78.8 \%$ (71.4 to 84.6) 91.5\% (88.2 to 94.0) 85.1\% (80.5 to 88.6) 90.4\% (86.5 to 93.3) 89.4\% (83.9 to 93.2) 85.9\% (82.8 to 88.6) 96.3\% (96.1 to 98.0) 83.0\% (74.8 to 89.0) 93.9\% (90.7 to 96.1) 81.0\% (78.1 to 83.6) Median 85.5\% IQR 79.4 to 93.3 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |


| Studies | Locatio n | Age | Sampl e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1. $>33.3 \%$ of participants from studies at high risk of bias |  |  |  |  |  |  |  |  |  |
| 2. $>33.3 \%$ participants from studies that were indirect |  |  |  |  |  |  |  |  |  |
| 3. $<33.3 \%$ of studies had $>100$ participants |  |  |  |  |  |  |  |  |  |
| 4. Confidence intervals were non-overlapping |  |  |  |  |  |  |  |  |  |
| 5. $<33.3 \%$ of studies had $>300$ participants |  |  |  |  |  |  |  |  |  |
| 6. Li 2018 groups data for children <3 months old and for children over 3 months old |  |  |  |  |  |  |  |  |  |
| 7. Kim 2009 groups data for children $\leq 5$ months old and for children between 5 months to $<5$ years old |  |  |  |  |  |  |  |  |  |
| 8. Yoon 2016 groups data for children $\leq 6$ months old and for children over 6 months old |  |  |  |  |  |  |  |  |  |
| 9. This study is one of two studies that formed the basis of the previous 2013 recommendations |  |  |  |  |  |  |  |  |  |

At presentation: polymorphous rash

| Studies | Locatio <br> n | Age | Sample size | \% with Symptom $\text { ( } 95 \% \mathrm{CI} \text { ) }$ | Risk of bias | Indirectness | Inconsisten cy | Imprecisio $\mathrm{n}$ | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Moore 2014 (primary care) | UK | All | 74 | 63.5\% (52.1 to 73.6) <br> Median 63.5\% <br> IQR - | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Shiozawa 2013 (at time of presentation of principal symptoms other than fever) | Outside Europe | $\begin{aligned} & <1 \\ & \text { year } \end{aligned}$ | 51 | $\begin{aligned} & 78.4 \% ~(65.4 \text { to } \\ & 87.5)^{4} \\ & \text { Median } 78.4 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Shiozawa 2013 (at time of presentation of principal symptoms other than fever) | Outside Europe | $\begin{aligned} & \geq 1 \\ & \text { year } \end{aligned}$ | 49 | $\begin{aligned} & 24.5 \% ~(14.6 \text { to } \\ & 38.1)^{4} \\ & \text { Median } 24.5 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Shiozawa 2013 (day 5 of fever) | Outside Europe | $\begin{aligned} & <2 \\ & \text { year } \end{aligned}$ | 51 | $\begin{aligned} & 92.2 \% ~(81.5 \text { to } \\ & 96.9)^{4} \\ & \text { Median } 92.2 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Shiozawa 2013 (day 5 of fever) | Outside Europe | $\begin{aligned} & \geq 2 \\ & \text { year } \end{aligned}$ | 49 | $\begin{aligned} & 79.6 \%(66.4 \text { to } \\ & 88.5)^{4} \end{aligned}$ | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |


| Studies | Locatio n | Age | Sample size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | Median 79.6\% IQR - |  |  |  |  |  |

1. $>33.3 \%$ of participants from studies at high risk of bias
2. $>33.3 \%$ participants from studies that were partially direct
3. $<33.3 \%$ of studies had $>100$ participants
4. Shiozawa 2013 groups data for children $\leq 2$ years old and for children over 2 years old

During course of illness: cervical lymphadenopathy

| Studies | Locatio n | Age | Sampl <br> e size | $\begin{aligned} & \text { \% with Symptom } \\ & \text { (95\% CI) } \end{aligned}$ | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Maric 2015 <br> Giannouli 2013 <br> Perrin 2009 | Europe (not UK) | All | $\begin{aligned} & 78 \\ & 66 \\ & 39 \end{aligned}$ | 43.6\% (33.1 to 54.6) <br> 72.7\% (61.0 to 82.0) <br> 38.5\% (24.9 to 54.1) <br> Median 43.6\% <br> IQR 41.1 to 58.2 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Very serious ${ }^{3}$ | Very low |
| Manlhiot 2012 <br> Tang 2016 <br> Bai 2017 <br> Behmadi 2019 <br> Sonobe 2007 <br> Kil 2017 <br> Chang 2014 <br> Sittiwangkul 2013 <br> Tewelde 2014 <br> Ghelani 2012 <br> Yellen 2010 | Outside Europe | All | $\begin{aligned} & 738 \\ & 716 \\ & 298 \\ & 105 \\ & 13301 \\ & 387 \\ & 226 \\ & 147 \\ & 67 \\ & 127 \\ & 137 \end{aligned}$ | $74.7 \%$ (71.4 to 77.7) <br> $73.0 \%$ (69.7 to 76.2) <br> 24.8\% (20.3 to 30.0) <br> 63.8\% (54.3 to 72.4) <br> $75.3 \%$ (74.6 to 76.0) <br> 63.0\% (58.1 to 67.7) <br> 40.3\% (34.1 to 46.8) <br> 42.9\% (35.1 to 50.9) <br> 37.3\% (26.7 to 49.3) <br> 72.4\% (64.1 to 79.5) <br> 53.3\% (45.0 to 61.4) <br> Median 63.0\% <br> IQR 41.6 to 72.7 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |


| Studies | Locatio <br> n | Age | Sampl <br> e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Meric 2015 <br> Giannouli 2013 <br> Perrin 2009 | Europe (not UK) | All | $\begin{aligned} & 33 \\ & 22 \\ & 20 \end{aligned}$ | 27.3\% (15.1 to 44.2) <br> 59.1\% (38.7 to 76.7) <br> 30.0\% (14.6 to 51.9) <br> Median 30.0\% <br> IQR 28.7 to 44.6 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Very serious ${ }^{3}$ | Very low |
| Manlhoit 2012 <br> Tang 2016 <br> Bai 2017 <br> Behmadi 2019 <br> Sonobe 2007 <br> Lee 2016 <br> Kil 2017 <br> Sittiwangkul 2013 <br> Falcini 2012 <br> Tewelde 2014 <br> Ghelani 2012 <br> Yellen 2010 | Outside Europe | All | 62 300 85 71 2556 111 228 61 228 38 76 53 | 28.6\% (23.0 to 34.9) <br> 33.7\% (28.6 to 39.2) <br> 64.7\% (54.1 to 74.0) <br> 19.7\% (12.1 to 30.4) <br> 38.6\% (36.8 to 40.5) <br> 40.5\% (31.9 to 49.8) <br> 32.5\% (26.7 to 38.8) <br> 18.0\% (10.4 to 29.5) <br> 39.5\% (33.4 to 45.9) <br> 21.1\% (11.1 to 36.4) <br> 30.3\% (21.1 to 41.3) <br> 18.9\% (10.6 to 31.4) <br> Median 31.4\% <br> IQR 20.8 to 38.8 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Stemberger Maric 2018 <br> Patel 2013 <br> Generini 1997 <br> Falcini 2007 <br> Sanchez-Manubens 2016 <br> Tacke 2014 | Europe (not UK) | All | $\begin{aligned} & 110 \\ & 314 \\ & 73 \\ & 266 \\ & 399 \\ & 319 \end{aligned}$ | 30.9\% (23.0 to 40.1) <br> $75.5 \%$ (70.4 to 79.9) <br> 34.2\% (24.4 to 45.7) <br> $71.8 \%$ ( 66.1 to 76.9 ) <br> 28.8\% (24.6 to 33.5) <br> $71.8 \%$ (66.6 to 76.4) <br> Median 53.0\% <br> IQR 31.7 to 71.8 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Serious ${ }^{5}$ | Very low |
| Ebbeson 2004 <br> Ruan 2013 <br> Li 2018 | Outside Europe | $\begin{aligned} & <1 \\ & \text { year } \end{aligned}$ | $\begin{aligned} & 32 \\ & 49 \\ & 40 \end{aligned}$ | $\begin{aligned} & 3.1 \%(0.55 \text { to } 15.7) \\ & 14.3 \%(7.10 \text { to } \\ & 26.7)^{6} \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Very serious ${ }^{3}$ | Very low |


| Studies | Locatio n | Age | Sampl e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Kim 2009 <br> Yoon 2016 <br> Teng 2012 <br> Liu 2012 |  |  | $\begin{aligned} & 22 \\ & 26 \\ & 109 \\ & 65 \end{aligned}$ | ```30.0% (18.1 to 45.4)}\mp@subsup{}{}{7 31.8% (16.4 to 52.7)}\mp@subsup{}{}{8 15.4% (6.15 to 33.5)9 36.7% (28.3 to 46.1) 24.6% (15.8 to 36.3) Median 24.6% IQR 14.9 to 30.9``` |  |  |  |  |  |
| Ebbeson 2004 <br> Ruan 2013 <br> Li 2018 <br> Kim 2009 <br> Yoon 2016 <br> Teng 2012 <br> Liu 2012 | Outside Europe | $\geq 1$ <br> year | $\begin{aligned} & 92 \\ & 1160 \\ & 160 \\ & 131 \\ & 213 \\ & 242 \\ & 80 \end{aligned}$ | $\begin{aligned} & 29.3 \%(21.0 \text { to } 39.3) \\ & 31.4 \%(28.8 \text { to } \\ & 34.1)^{6} \\ & 66.9 \%(59.3 \text { to } \\ & 73.7)^{7} \\ & 35.9 \%(28.2 \text { to } \\ & 44.4)^{8} \\ & 57.3 \%(50.6 \text { to } \\ & 63.7)^{9} \\ & 62.4 \%(56.1 \text { to } 68.3) \\ & 63.8 \%(52.8 \text { to } 73.4) \\ & \text { Median } 57.3 \% \\ & \text { IQR } 33.7 \text { to } 63.1 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Boudiaf 2016 <br> Saundankar 2014 <br> Zhang 2016 <br> Zhang 2012 <br> Zhu 2015 <br> Chen 2016 <br> Sun 2018 <br> Peng 2019 <br> Advani 2019 <br> Shamsizadeh 2014 | Outside <br> Europe | All | $\begin{aligned} & 133 \\ & 353 \\ & 518 \\ & 577 \\ & 231 \\ & 2304 \\ & 1008 \\ & 1420 \\ & 542 \\ & 104 \end{aligned}$ | $\begin{aligned} & 28.6 \%(21.6 \text { to } 36.8) \\ & 59.5 \%(54.3 \text { to } 64.5) \\ & 62.0 \%(57.7 \text { to } 66.1) \\ & 69.3 \%(65.4 \text { to } 73.0) \\ & 66.7 \%(60.4 \text { to } 72.4) \\ & 54.6 \%(52.5 \text { to } 56.6) \\ & 76.7 \%(74.0 \text { to } 79.2) \\ & 36.8 \%(34.3 \text { to } 39.3) \\ & 39.5 \%(35.5 \text { to } 43.7) \\ & 42.3 \%(33.3 \text { to } 51.9) \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |


| Studies | Locatio <br> n | Age | $\begin{aligned} & \text { Sampl } \\ & \text { e size } \end{aligned}$ | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Kubota 2008 <br> Piao 2010 <br> Nomura 2012 <br> Tajima 2015 <br> Garrido-Garcia 2017 <br> Gorrab 2016 <br> Kim 2017 <br> Jun 2017 <br> Jun 2015 <br> Kim 2018 <br> Hu 2019 <br> Sittiwangkul 2011 <br> LuAnn Minich 2007 <br> Wang 2009 <br> Bal 2014 <br> Sehgal 2015 <br> Huang 2006 |  |  | 136 735 207 100 399 146 14916 146 355 329 293 170 562 243 106 312 768 | 66.9\% (58.6 to 74.3) 68.8\% (65.4 to 72.1) $72.0 \%$ (65.5 to 77.7) 89.0\% (81.4 to 93.8) 67.7\% (62.9 to 72.1) 30.8\% (23.9 to 38.7) 59.4\% (58.6 to 60.2) 46.6\% (38.7 to 54.7) 66.2\% (61.1 to 70.9) 21.6\% (17.5 to 26.3) 29.7\% (24.8 to 35.2) 33.5\% (26.9 to 40.9) 44.0\% (39.9 to 48.1) 26.7\% (21.6 to 32.6) $74.5 \%$ (65.5 to 81.9) 46.8\% (41.3 to 52.3) 69.3\% (65.9 to 72.4) Median 59.4\% IQR 38.2 to 68.3 |  |  |  |  |  |
| 1. $>33.3 \%$ of particip <br> 2. $>33.3 \%$ participan <br> 3. $<33.3 \%$ of studies <br> 4. Confidence interv <br> 5. $<33.3 \%$ of studies <br> 6. Ruan 2013 group <br> 7. Li 2018 groups da <br> 8. Kim 2009 groups <br> 9. Yoon 2016 group <br> 10. This study is one | from studi <br> m studies <br> $>100$ particip <br> re non-ov <br> >300 parti <br> for childr <br> children <br> or children <br> for childr <br> studies | $s$ at $h$ <br> hat w <br> ipant <br> rlapp <br> ipant <br> \ll6 <br> 3 mon <br> $\leq 5 \mathrm{~m}$ <br> $\leq 6$ <br> at form | risk of indirec <br> nths old s old and ths old nths old $d$ the ba | as <br> and for children betwe for children over 3 mo d for children between and for children over 6 is of the previous 2013 | 6 month hs old 5 months months old recomme | o 5 years old $<5$ years old ations |  |  |  |

At presentation: cervical lymphadenopathy

| Studies | Locatio $\mathrm{n}$ | Age | Sample size | $\begin{aligned} & \text { \% with Symptom } \\ & (95 \% \mathrm{CI}) \end{aligned}$ | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Moore 2014 (primary care) | UK | All | 74 | $35.1 \% \text { (25.2 to 46.5) }$ <br> Median 35.1\% IQR - | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Shiozawa 2013 (at time of presentation of principal symptoms other than fever) | Outside Europe | $\begin{aligned} & <2 \\ & \text { year } \end{aligned}$ | 51 | $\begin{aligned} & 35.3 \%(23.6 \text { to } \\ & 49.0)^{4} \\ & \text { Median } 35.3 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Shiozawa 2013 (at time of presentation of principal symptoms other than fever) | Outside Europe | $\begin{aligned} & \geq 2 \\ & \text { year } \end{aligned}$ | 49 | $\begin{aligned} & 75.5 \%(61.9 \text { to } \\ & 85.4)^{4} \\ & \text { Median } 75.5 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Shiozawa 2013 (day 5 of fever) | Outside Europe | $\begin{aligned} & <2 \\ & \text { year } \end{aligned}$ | 51 | $\begin{aligned} & 64.7 \% ~(51.0 \text { to } \\ & 76.4)^{4} \end{aligned}$ <br> Median 64.7\% IQR - | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Shiozawa 2013 (day 5 of fever) | Outside Europe | $\begin{aligned} & \geq 2 \\ & \text { year } \end{aligned}$ | 49 | $\begin{aligned} & 93.9 \%(83.9 \text { to } \\ & 97.9)^{4} \\ & \text { Median } 93.9 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| 1. $>33.3 \%$ of participants from studies at high risk of bias <br> 2. $>33.3 \%$ participants from studies that were partially direct <br> 3. $<33.3 \%$ of studies had $>100$ participants <br> 4. Shiozawa 2013 groups data for children $\leq 2$ years old and for children over 2 years old |  |  |  |  |  |  |  |  |  |


| Studies | Locatio n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 0 symptoms: <br> Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Moore $2014^{5}$ (at first presentation) | UK | all | 74 | 28.4\% (19.4 to 38.5) | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |


| Studies | Locatio n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1 symptom <br> Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Moore $2014{ }^{5}$ (at first presentation) | UK | all | 74 | $39.2 \%$ (28.9 to 50.6) | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| 2 symptoms: <br> Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Moore $2014^{5}$ (at first presentation) | UK | all | 74 | 9.5\% (4.7 to 18.3) | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Shiozawa $2013^{6}$ (at time of presentation of principal symptoms other than fever) | Outside Europe | $\begin{aligned} & <1 \\ & \text { year } \end{aligned}$ | 51 | $\begin{aligned} & 54.9 \%(41.4 \text { to } \\ & 67.7)^{4} \end{aligned}$ | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Shiozawa $2013^{6}$ (at time of presentation of principal symptoms other than fever) | Outside Europe | $\begin{aligned} & \geq 1 \\ & \text { year } \end{aligned}$ | 49 | $\begin{aligned} & 16.3 \%(8.51 \text { to } \\ & 29.0)^{4} \end{aligned}$ | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| 3 symptoms: <br> Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Moore $2014^{5}$ (at first presentation) | UK | all | 74 | $6.8 \%$ (2.9 to 14.9) | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| 4 symptoms: <br> Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Moore $2014^{5}$ (at first presentation) | UK | all | 74 | $5.4 \%$ (2.1 to 13.1) | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| 5 symptoms: <br> Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Moore $2014^{5}$ (at first presentation) | UK | all | 74 | $5.4 \%$ (2.1 to 13.1) | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very Iow |
| 6 symptoms: <br> Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Moore $2014{ }^{5}$ (at first presentation) | UK | all | 74 | 1.4\% (0.2 to 7.3) | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |


| Studies | Locatio n | Age | Sample size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 7 symptoms: <br> Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Moore $2014^{5}$ (at first presentation) | UK | all | 74 | 1.4\% (0.2 to 7.3) | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| 8 symptoms: <br> Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Moore $2014^{5}$ (at first presentation) |  | all | 74 | 2.7\% (0.7 to 9.3) | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| 1. $>33.3 \%$ of participants from studies at high risk of bias |  |  |  |  |  |  |  |  |  |
| 2. $>33.3 \%$ participants from studies that were partially direct |  |  |  |  |  |  |  |  |  |
| 3. $<33.3 \%$ of studies had $>100$ participants |  |  |  |  |  |  |  |  |  |
| 4. Shiozawa 2013 groups data for children $\leq 2$ years old and for children over 2 years old |  |  |  |  |  |  |  |  |  |
| 5. Symptoms included Rash, Lymphadenopathy, Conjunctivitis, Red, dry, or cracked lips, Strawberry tongue, Redness in mouth, Peeling skin, Red palms/soles, Oedema |  |  |  |  |  |  |  |  |  |
| 6. Symptoms included conjunctival injection, oral changes, polymorphous rash, changes in extremities, cervical lymphadenopathy |  |  |  |  |  |  |  |  |  |

Other signs and symptoms
During course of illness: anterior uveitis

| Studies | Locatio $\mathrm{n}$ | Age | $\begin{aligned} & \text { Sampl } \\ & \text { e size } \end{aligned}$ | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Lee 2016 | Outside Europe | All | 111 | $36.9 \%(28.5 \text { to } 46.2)$ <br> Median 36.9\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{3}$ | Very low |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Sanchez-Manubens 2016 | Europe (not UK) | All | 399 | $2.8 \%(1.55 \text { to } 4.87)$ <br> Median 2.8\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Not serious | Very low |


| Studies | Locatio <br> n | Age | $\begin{aligned} & \text { Sampl } \\ & \text { e size } \end{aligned}$ | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 1. $>33.3$ <br> 2. $>33.3$ <br> 3. $<33.3$ | from stud | s at h | risk of |  |  |  |  |  |  |


| Studies | Locatio n | Age | Sampl <br> e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Sanchez-Manubens $2016$ | Europe <br> (not UK) | All | 399 | $2.8 \% \text { (1.55 to 4.87) }$ <br> Median 2.8\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Not serous | Very low |
| Boudiaf 2016 | Outside Europe | All | 133 | 24.1 (17.6 to 32.0) <br> Median 24.1\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{3}$ | Very low |
| 1. $>33.3 \%$ of participants from studies at high risk of bias <br> 2. $>33.3 \%$ participants from studies that were indirect <br> 3. $<33.3 \%$ of studies had $>300$ participants |  |  |  |  |  |  |  |  |  |

During course of illness: arthritis

| Studies | Locatio n | Age | Sampl <br> e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Sanchez-Manubens $2016$ | Europe (not UK) | All | 399 | $13.8 \% \text { (10.8 to } 17.5)$ <br> Median 13.8\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Not serous | Very low |
| Boudiaf 2016 | Outside Europe | All | 133 | 7.5\% (4.14 to 13.3) | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{3}$ | Very low |


| Studies | Locatio <br> n | Age | Sampl e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | Median 7.5\% IQR - |  |  |  |  |  |
| 1. $>33.3 \%$ of participants from studies at high risk of bias <br> 2. $>33.3 \%$ participants from studies that were indirect <br> 3. $<33.3 \%$ of studies had $>300$ participants |  |  |  |  |  |  |  |  |  |

During course of illness: arthritis or arthralgia

| Studies | Locatio n | Age | Sample size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Manlhiot 2012 <br> Behmadi 2019 <br> Yun 2011 <br> Sittiwangkul 2013 | Outside Europe | All | $\begin{aligned} & 738 \\ & 105 \\ & 83 \\ & 147 \end{aligned}$ | $\begin{aligned} & 12.6 \%(10.4 \text { to } 15.2) \\ & 6.7 \% \text { (3.3 to } 13.1) \\ & 15.7 \%(9.39 \text { to } 25.0) \\ & 4.1 \%(1.9 \text { to } 8.2) \\ & \text { Median } 9.7 \% \\ & \text { IQR } 6.1 \text { to } 13.4 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Manlhiot 2012 <br> Behmadi 2019 <br> Yun 2011 <br> Sittiwangkul 2013 <br> Falcini 2012 | Outside Europe | All | $\begin{aligned} & 217 \\ & 71 \\ & 38 \\ & 61 \\ & 228 \end{aligned}$ | $\begin{aligned} & 12.4 \%(8.7 \text { to } 17.5) \\ & 8.5 \%(3.93 \text { to } 17.2) \\ & 5.3 \%(1.5 \text { to } 17.3) \\ & 0.0 \%(0.0 \text { to } 5.9) \\ & 4.4 \%(2.4 \text { to } 7.9) \end{aligned}$ <br> Median 5.3\% IQR 4.4 to 8.5 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Generini 1997 <br> Martins 2018 <br> Tacke 2014 | Europe (not UK) | All | $\begin{aligned} & 73 \\ & 63 \\ & 319 \end{aligned}$ | $\begin{aligned} & 1.4 \%(0.24 \text { to } 7.4) \\ & 12.7 \%(6.6 \text { to } 23.1) \\ & 10.3 \%(7.5 \text { to } 14.2) \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Very serious ${ }^{3}$ | Very low |


| Studies | Locatio <br> n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | Median 10.3\% IQR 5.9 to 11.5 |  |  |  |  |  |
| Saundankar 2014 <br> Peng 2019 <br> Shamsizedeh 2014 <br> Gorrab 2016 <br> Sehgal 2015 <br> Baker 2009 | Outside <br> Europe | All | $\begin{aligned} & 353 \\ & 1420 \\ & 104 \\ & 146 \\ & 312 \\ & 198 \end{aligned}$ | $\begin{aligned} & 21.0 \%(17.0 \text { to } 25.5) \\ & 10.6 \%(9.14 \text { to } 12.3) \\ & 14.4 \%(8.9 \text { to } 22.4) \\ & 30.8 \%(23.9 \text { to } 38.7) \\ & 8.3 \%(5.8 \text { to } 11.9) \\ & 14.6 \%(10.4 \text { to } 20.2) \\ & \text { Median } 14.5 \% \\ & \text { IQR } 11.6 \text { to } 19.4 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| 1. $>33.3 \%$ of participants from studies at high risk of bias |  |  |  |  |  |  |  |  |  |
| 2. $>33.3 \%$ participants from studies that were indirect |  |  |  |  |  |  |  |  |  |
| 3. $<33.3 \%$ of studies had $>100$ participants |  |  |  |  |  |  |  |  |  |
| 4. Confidence intervals were non-overlapping |  |  |  |  |  |  |  |  |  |
| 5. $<33.3 \%$ of studies had $>300$ participants |  |  |  |  |  |  |  |  |  |

During course of illness: BCG scar reaction

| Studies | Locatio <br> n | Age | $\begin{aligned} & \text { Sample } \\ & \text { size } \end{aligned}$ | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Maric 2015 | Europe (not UK) | All | 78 | $10.3 \% \text { (5.3 to 19.0) }$ <br> Median 10.3\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Tang 2016 <br> Bai 2017 <br> Chang 2014 <br> Sittiwangkul 2013 | Outside Europe | All | $\begin{aligned} & 716 \\ & 298 \\ & 226 \\ & 147 \end{aligned}$ | 14.0\% (11.6 to 16.7) 85.9\% (81.5 to 89.4) 43.8\% (37.5 to 50.3) 10.9\% (6.81 to 17.0) Median 28.9\% IQR 13.2 to 54.3 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |


| Studies | Locatio <br> n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Maric 2015 | Europe (not UK) | All | 33 | 6.1\% (1.7 to 19.6) Median 6.1\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Tang 2016 <br> Bai 2017 <br> Sittiwangkul 2013 | Outside Europe | All | $\begin{aligned} & 300 \\ & 85 \\ & 61 \end{aligned}$ | 12.3\% (9.1 to 16.5) 76.5\% (66.4 to 84.2) 4.9\% (1.7 to 13.5) Median 12.3\% IQR 8.6 to 44.4 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Ruan 2013 <br> Li 2018 <br> Loh 2019 <br> Kim 2009 <br> Teng 2012 | Outside Europe | $\begin{aligned} & <1 \\ & \text { year } \end{aligned}$ | $\begin{aligned} & 49 \\ & 40 \\ & 99 \\ & 22 \\ & 109 \end{aligned}$ | $\begin{aligned} & 18.4 \%(10.0 \text { to } \\ & 31.4)^{6} \\ & 10.0 \%(4.0 \text { to } 23.1)^{7} \\ & 69.7(60.1 \text { to } 77.9) \\ & 72.7 \%(51.9 \text { to } \\ & 86.9)^{8} \\ & 45.9 \%(36.8 \text { to } 55.2) \\ & \text { Median } 45.9 \% \\ & \text { IQR } 18.4 \text { to } 72.7 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Ruan 2013 <br> Li 2018 <br> Loh 2019 <br> Kim 2009 <br> Teng 2012 | Outside Europe | $\geq 1$ <br> year | $\begin{aligned} & 1160 \\ & 160 \\ & 180 \\ & 131 \\ & 242 \end{aligned}$ | $\begin{aligned} & 3.4 \%(2.5 \text { to } 4.6)^{6} \\ & 1.3 \%(0.34 \text { to } 4.4)^{7} \\ & 27.8 \%(21.8 \text { to } 34.7) \\ & 33.6 \%(26.1 \text { to } \\ & 42.0)^{8} \\ & 16.1 \%(12.0 \text { to } 21.2) \end{aligned}$ <br> Median 16.1\% IQR 3.4 to 27.8 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Boudiaf 2016 <br> Zhang 2012 <br> Piao 2010 <br> Garrido-Garcia 2017 <br> Kim 2018 <br> Hu 2019 | Outside Europe | All | $\begin{aligned} & 133 \\ & 577 \\ & 735 \\ & 399 \\ & 329 \\ & 293 \end{aligned}$ | 1.5\% (0.4 to 5.3) <br> $1.7 \%$ ( 0.9 to 3.2 ) <br> 1.1\% (0.6 to 2.1) <br> 24.3\% (20.4 to 28.8) <br> 25.5\% (21.1 to 30.5) <br> 36.9\% (31.5 to 42.5) | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very <br> low |


| Studies | Locatio n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Uehara 2010 |  |  | 15524 | 49.9\% (49.1 to 50.7) <br> Median 24.3\% <br> IQR 1.6 to 31.2 |  |  |  |  |  |

1. $>33.3 \%$ of participants from studies at high risk of bias
2. $>33.3 \%$ participants from studies that were indirect
3. $<33.3 \%$ of studies had $>100$ participants
4. Confidence intervals were non-overlapping
5. $<33.3 \%$ of studies had $>300$ participants
6. Ruan 2013 groups data for children $<6$ months old and for children between 6 months to 5 years old
7. Li 2018 groups data for children $<3$ months old and for children over 3 months old
8. Kim 2009 groups data for children $\leq 5$ months old and for children between 5 months to $<5$ years old

During course of illness: changes in the extremities: desquamation (and/or decrustation of at least 1 toe)

| Studies | Locatio n | Age | Sampl e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Bai 2017 <br> Tang 2016 <br> Chang 2014 | Outside Europe | All | $\begin{aligned} & 298 \\ & 716 \\ & 226 \end{aligned}$ | 97.7\% (95.2 to 98.9) <br> 68.3\% (64.8 to 71.6) <br> 95.6\% (92.1 to 97.6) <br> Median 95.6\% <br> IQR 82.0 to 96.7 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Bai 2017 <br> Tang 2016 | Outside Europe | All | $\begin{aligned} & 85 \\ & 300 \end{aligned}$ | $\begin{aligned} & 77.6 \%(67.7 \text { to } 85.2) \\ & 43.0 \%(37.5 \text { to } 48.7) \\ & \text { Median } 60.3 \% \\ & \text { IQR } 51.7 \text { to } 69.0 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Ruan 2013 | Outside Europe | $\begin{aligned} & <1 \\ & \text { year } \end{aligned}$ | 49 | 51.0\% (37.5 to $64.4)^{6}$ <br> Median 51.0\% | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |


| Studies | Locatio n | Age | Sampl e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | IQR - |  |  |  |  |  |
| Ruan 2013 | Outside Europe | $\begin{aligned} & \geq 1 \\ & \text { year } \end{aligned}$ | 1160 | $\begin{aligned} & 50.3 \% ~(47.5 \text { to } \\ & 53.2)^{6} \\ & \text { Median } 50.3 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| Piao 2010 <br> Sanchez-Manubens <br> 2016 <br> Chen 2016 <br> Sun 2018 <br> Kim 2018 <br> Huang 20067 | Outside Europe | All | 735 399 2304 1008 329 768 | 54.0\% (50.4 to 57.6) <br> $31.1 \%$ ( 26.7 to 35.8 ) <br> $32.2 \%$ (30.3 to 34.1) <br> 51.3\% (48.2 to 54.4) <br> 53.8\% (48.4 to 59.1) <br> 82.9\% (80.1 to 85.4) <br> Median 52.6\% <br> IQR 37.0 to 54.0) | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| 1. $>33.3 \%$ of partici <br> 2. $>33.3 \%$ participa <br> 3. $<33.3 \%$ of studie <br> 4. Confidence inter <br> 5. $<33.3 \%$ of studie <br> 6. Ruan 2013 group <br> 7. This study is one | from studi <br> m studies <br> $>100$ part <br> ere non-ov <br> >300 partic <br> for childr <br> studies th | at hi hat we ipants rlappin ipants $\mathrm{n}<6 \mathrm{~m}$ at form | risk of indirect <br> nths old d the ba | ias <br> and for children between is of the previous 2013 | 6 month comme | o 5 years old ations |  |  |  |

At presentation: changes in the extremities: desquamation

| Studies | Locatio n | Age | Sample size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Moore 2014 | UK | All | 74 | $18.9 \% \text { (11.6 to } 29.3 \text { ) }$ <br> Median 18.9\% IQR - | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| 1. $>33.3 \%$ of participants from studies at high risk of bias <br> 2. $>33.3 \%$ participants from studies that were partially direct |  |  |  |  |  |  |  |  |  |


| Studies | Locatio <br> n | Age | Sample size | \% with Symptom $(95 \% \mathrm{Cl})$ | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

3. $<33.3 \%$ of studies had $>100$ participants

During course of illness: changes in the extremities: oedema

| Studies | Locatio n | Age | Sampl <br> e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Tang 2016 | Outside Europe | All | 716 | 60.1\% (56.4 to 63.6) Median 60.1\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Tang 2016 | Outside Europe | All | 300 | $22.7 \%(18.3 \text { to } 27.7)$ <br> Median 22.7\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Sanchez-Manubens $2016$ | Europe (not UK) | All | 399 | $32.8 \% \text { (28.4 to } 37.6 \text { ) }$ <br> Median 32.8\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Ruan 2013 | Outside Europe | $\begin{aligned} & <1 \\ & \text { year } \end{aligned}$ | 49 | $\begin{aligned} & 59.2 \% ~(45.3 \text { to } \\ & 71.8)^{6} \\ & \text { Median } 59.2 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| Ruan 2013 | Outside Europe | $\begin{aligned} & \geq 1 \\ & \text { year } \end{aligned}$ | 1160 | $\begin{aligned} & 79.5 \%(77.1 \text { to } \\ & 81.7)^{6} \\ & \text { Median } 79.5 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| Sun 2018 <br> Bal 2014 <br> Huang 20067 | Outside Europe | All | $\begin{aligned} & 1008 \\ & 106 \\ & 768 \end{aligned}$ | 44.0\% (41.0 to 47.1) <br> 47.2\% (37.9 to 56.6) <br> 48.0\% (44.5 to 51.6) <br> Median 47.2\% <br> IQR 45.6 to 47.6 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |


| Studies | Locatio <br> n | Age | Sampl <br> e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

1. $>33.3 \%$ of participants from studies at high risk of bias
2. $>33.3 \%$ participants from studies that were indirect
3. $<33.3 \%$ of studies had $>100$ participants
4. Confidence intervals were non-overlapping
5. $<33.3 \%$ of studies had $>300$ participants
6. Ruan 2013 groups data for children $<6$ months old and for children between 6 months to 5 years old
7. This study is one of two studies that formed the basis of the previous 2013 recommendations

At presentation: changes in the extremities: oedema

| Studies | Locatio <br> n | Age | $\begin{aligned} & \text { Sample } \\ & \text { size } \end{aligned}$ | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Moore 2014 | UK | All | 74 | $13.5 \%$ ( 7.51 to 23.1) <br> Median 13.5\% <br> IQR - | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| 1. $>33.3 \%$ of participants from studies at high risk of bias <br> 2. $>33.3 \%$ participants from studies that were partially direct <br> 3. $<33.3 \%$ of studies had $>100$ participants |  |  |  |  |  |  |  |  |  |

During course of illness: changes in the extremities: red palms/soles

| Studies | Locatio n | Age | Sampl <br> e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Sanchez-Manubens 2016 | Europe (not UK) | All | 399 | $\begin{aligned} & 29.1 \%(24.8 \text { to } 33.7) \\ & \text { Median } 29.1 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Ruan 2013 | Outside Europe | $\begin{aligned} & <1 \\ & \text { year } \end{aligned}$ | 49 | 87.8\% (75.8 to 94.3) ${ }^{6}$ <br> Median 87.8\% | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |


| Studies | Locatio n | Age | Sampl <br> e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | IQR - |  |  |  |  |  |
| Ruan 2013 | Outside Europe | $\begin{aligned} & \geq 1 \\ & \text { year } \end{aligned}$ | 1160 | $\begin{aligned} & 83.3 \% ~(81.0 \text { to } \\ & 85.3)^{6} \\ & \text { Median } 83.3 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| Sun 2018 | Outside Europe | All | 1008 | $78.3 \% \text { (75.6 to 80.7) }$ <br> Median 78.3\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |

1. $>33.3 \%$ of participants from studies at high risk of bias
2. $>33.3 \%$ participants from studies that were indirect
3. $<33.3 \%$ of studies had $>100$ participants
4. Confidence intervals were non-overlapping
5. $<33.3 \%$ of studies had $>300$ participants
6. Ruan 2013 groups data for children $<6$ months old and for children between 6 months to 5 years old

At presentation: changes in the extremities: red palms/soles

| Studies | Locatio n | Age | Sample <br> size | \% with Symptom $\text { ( } 95 \% \mathrm{CI} \text { ) }$ | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Moore 2014 | UK | All | 74 | $17.6 \%$ (10.6 to 27.8) <br> Median 17.6\% IQR - | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| 1. $>33.3 \%$ of participants from studies at high risk of bias <br> 2. $>33.3 \%$ participants from studies that were partially direct <br> 3. $<33.3 \%$ of studies had $>100$ participants |  |  |  |  |  |  |  |  |  |

During course of illness: desquamation

| Studies | Locatio n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Teng 2012 | Outside Europe | $\begin{aligned} & <1 \\ & \text { year } \end{aligned}$ | 109 | 50.5\% (41.2 to 59.7) <br> Median 50.5\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| Teng 2012 | Outside <br> Europe | $\geq 1$ <br> year | 242 | $40.5 \% \text { (34.5 to 46.8) }$ <br> Median 40.5\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| Bal 2014 | Outside Europe | All | 106 | 24.5\% (17.3 to 33.5) <br> Median 24.5\% <br> IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| 1. $>33.3 \%$ of participants from studies at high risk of bias <br> 2. $>33.3 \%$ participants from studies that were indirect <br> 3. $<33.3 \%$ of studies had $>100$ participants <br> 4. Confidence intervals were non-overlapping <br> 5. $<33.3 \%$ of studies had $>300$ participants |  |  |  |  |  |  |  |  |  |

During course of illness: anal desquamation

| Studies | Locatio <br> n | Age | Sample size | $\begin{aligned} & \text { \% with Symptom } \\ & \text { (95\% CI) } \end{aligned}$ | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Tang 2016 <br> Bai 2017 <br> Chang 2014 | Outside Europe | All | $\begin{aligned} & 716 \\ & 298 \\ & 226 \end{aligned}$ | 43.7\% (40.1 to 47.4) <br> 46.0\% (40.4 to 51.7) <br> 24.3\% (19.2 to 30.3) <br> Median 43.7\% <br> IQR 34.0 to 44.9 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Tang 2016 Bai 2017 Falcini 2012 | Outside Europe | All | $\begin{aligned} & 300 \\ & 85 \\ & 24 \end{aligned}$ | $\begin{aligned} & 38.3 \%(33.0 \text { to } 44.0) \\ & 51.8 \%(41.3 \text { to } 62.1) \\ & 10.5 \%(7.2 \text { to } 15.2) \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

| Studies | Locatio n | Age | Sample size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | Median 38.3\% IQR 24.4 to 45.1 |  |  |  |  |  |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Ruan 2013 <br> Kim 2009 <br> Liu 2012 | Outside Europe | $\begin{aligned} & <1 \\ & \text { year } \end{aligned}$ | $\begin{aligned} & 49 \\ & 22 \\ & 65 \end{aligned}$ | $\begin{aligned} & 30.6 \%(19.5 \text { to } \\ & 44.5)^{6} \\ & 4.5 \%(0.8 \text { to } 21.8)^{7} \\ & 13.8 \%(7.5 \text { to } 24.3) \\ & \text { Median } 13.8 \% \\ & \text { IQR 9.2\% to 22.2\% } \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Ruan 2013 <br> Kim 2009 <br> Liu 2012 | Outside <br> Europe | $\geq 1$ <br> year | $\begin{aligned} & 1160 \\ & 131 \\ & 80 \end{aligned}$ | $\begin{aligned} & 40.5 \%(37.7 \text { to } \\ & 43.4)^{6} \\ & 11.5 \%(7.1 \text { to } 18.0)^{7} \\ & 20.0 \%(12.7 \text { to } 30.1) \\ & \text { Median } \mathbf{2 0 . 0} \% \\ & \text { IQR } 15.8 \text { to } \mathbf{3 0 . 3} \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Boudiaf 2016 <br> Saundankar 2014 <br> Zhang 2016 <br> Zhang 2012 <br> Sun 2018 <br> Shamsizadeh 2014 <br> Piao 2010 <br> Wang 2009 <br> Huang $2006^{8}$ | Outside Europe | All | $\begin{aligned} & 133 \\ & 353 \\ & 518 \\ & 577 \\ & 1008 \\ & 104 \\ & 735 \\ & 243 \\ & 768 \end{aligned}$ | 83.5\% (76.2 to 88.8) <br> 16.1\% (12.7 to 20.4) <br> 22.8\% (19.4 to 26.6) <br> 34.7\% (30.9 to 38.6) <br> 29.2\% (26.4 to 32.1) <br> 31.7\% (23.6 to 41.2) <br> 6.4\% (4.8 to 8.4) <br> 67.9\% (61.8 to 73.5) <br> 45.2\% (41.7 to 48.7) <br> Median 31.7\% <br> IQR 22.8 to 45.2 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| 1. $>33.3 \%$ of partic <br> 2. $>33.3 \%$ participa <br> 3. $<33.3 \%$ of studie <br> 4. Confidence inter <br> 5. $<33.3 \%$ of studie <br> 6. Ruan 2013 grou |  | ies at that icipan verlap icipan <6 | gh risk of re indirec <br> ng <br> months old | ias <br> and for children between | 6 months | 5 years old |  |  |  |


7. Kim 2009 groups data for children $\leq 5$ months old and for children between 5 months to $<5$ years old
8. This study is one of two studies that formed the basis of the previous 2013 recommendations

During course of illness: gastrointestinal symptoms

| Studies | Locatio n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| $\begin{aligned} & \text { Manlhiot } 2012 \\ & \text { Yun } 2011 \end{aligned}$ | Outside Europe | All | $\begin{aligned} & 738 \\ & 83 \end{aligned}$ | $\begin{aligned} & 31.6 \%(28.3 \text { to } 35.0) \\ & 32.5 \%(23.4 \text { to } 43.2) \\ & \text { Median } 32.1 \% \\ & \text { IQR } 31.8 \text { to } 32.3 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Manlhiot 2012 <br> Yun 2011 <br> Falcini 2012 | Outside Europe | All | $\begin{aligned} & 217 \\ & 38 \\ & 228 \end{aligned}$ | $\begin{aligned} & 33.2 \%(27.3 \text { to } 39.7) \\ & 36.8 \%(23.4 \text { to } 52.7) \\ & 4.4 \% \text { (2.4 to } 7.9) \\ & \text { Median } 33.2 \% \\ & \text { IQR } 18.8 \text { to } 35.0 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Tacke 2014 <br> Fabi 2018 | Europe <br> (not UK) | All | $\begin{aligned} & 319 \\ & 302 \end{aligned}$ | $\begin{aligned} & 26.3 \%(21.8 \text { to } 31.4) \\ & 35.1 \%(29.9 \text { to } 40.6) \\ & \text { Median } 30.7 \% \\ & \text { IQR } 28.5 \text { to } 32.9 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Very serious ${ }^{3}$ | Very low |
| Teng 2012 | Outside Europe | $\begin{aligned} & <1 \\ & \text { year } \end{aligned}$ | 109 | $18.3 \% \text { (12.2 to } 26.7 \text { ) }$ <br> Median 18.3\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Teng 2012 | Outside Europe | $\begin{aligned} & \geq 1 \\ & \text { year } \end{aligned}$ | 242 | $20.2 \% \text { (15.7 to } 25.8)$ <br> Median 20.2\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Sehgal 2015 <br> Saundankar 2014 | Outside Europe | All | $\begin{aligned} & 312 \\ & 353 \end{aligned}$ | $\begin{aligned} & 55.1 \%(49.6 \text { to } 60.6) \\ & 54.1 \%(48.9 \text { to } 59.2) \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

| Studies | Locatio <br> n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Chen 2016 |  |  | 2304 | 27.5\% (25.7 to 29.3) |  |  |  |  |  |
| Gamez-Gonzalez 2013 |  |  | 214 | 33.2\% (27.2 to 39.7) |  |  |  |  |  |
| Baker 2009 |  |  | 198 | 60.6\% (53.7 to 67.2) |  |  |  |  |  |
|  |  |  |  | Median 54.1\% |  |  |  |  |  |

4. $>33.3 \%$ of participants from studies at high risk of bias
5. $>33.3 \%$ participants from studies that were indirect
6. $<33.3 \%$ of studies had $>100$ participants
7. Confidence intervals were non-overlapping
8. $<33.3 \%$ of studies had $>300$ participants

During course of illness: gastrointestinal symptoms: abdominal pain

| Studies | Locatio <br> n | Age | Sample <br> size | $\begin{aligned} & \text { \% with Symptom } \\ & (95 \% \mathrm{CI}) \end{aligned}$ | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| $\begin{aligned} & \text { Bai } 2017 \\ & \text { Yun } 2011 \end{aligned}$ | Outside Europe | All | $\begin{aligned} & 298 \\ & 83 \end{aligned}$ | $\begin{aligned} & 75.2 \%(70.0 \text { to } 79.7) \\ & 9.6 \%(5.0 \text { to } 17.9) \\ & \text { Median } 42.4 \% \\ & \text { IQR } 26.0 \text { to } 58.8 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| $\begin{aligned} & \text { Bai } 2017 \\ & \text { Yun } 2011 \end{aligned}$ | Outside Europe | All | $\begin{aligned} & 85 \\ & 38 \end{aligned}$ | $\begin{aligned} & 64.7 \%(54.1 \text { to } 74.0) \\ & 18.4 \%(9.2 \text { to } 33.4) \\ & \text { Median } 41.6 \% \\ & \text { IQR } 23.0 \text { to } 53.1 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Boudiaf 2016 <br> Gamez-Gonzalez 2013 <br> Gorrab 2016 <br> Baker 2009 | Outside Europe | All | $\begin{aligned} & 133 \\ & 214 \\ & 146 \\ & 198 \end{aligned}$ | $\begin{aligned} & 22.6 \%(16.3 \text { to } 30.4) \\ & 15.9 \%(11.6 \text { to } 21.4) \\ & 22.6 \%(16.6 \text { to } 30.0) \\ & 17.7 \%(13.0 \text { to } 23.6) \end{aligned}$ Median 20.2\% | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |


| Studies | Locatio <br> n | Age | Sample size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | IQR 17.3 to 22.6 |  |  |  |  |  |
| 1. $>33.3 \%$ of participants from studies at high risk of bias |  |  |  |  |  |  |  |  |  |
| 2. $>33.3 \%$ participants from studies that were indirect |  |  |  |  |  |  |  |  |  |
| 3. $<33.3 \%$ of studies had $>100$ participants |  |  |  |  |  |  |  |  |  |
| 4. Confidence intervals were non-overlapping |  |  |  |  |  |  |  |  |  |
| 5. $<33.3 \%$ of studies had $>300$ participants |  |  |  |  |  |  |  |  |  |

During course of illness: gastrointestinal symptoms: decreased food/fluid intake

| Studies | Locatio <br> n | Age | Sample size | \% with Symptom $\text { ( } 95 \% \mathrm{CI} \text { ) }$ | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Shamsizadeh 2014 <br> Baker 2009 | Outside Europe | All | $\begin{aligned} & 104 \\ & 198 \end{aligned}$ | $\begin{aligned} & 34.6 \%(26.2 \text { to } 44.2) \\ & 36.9 \%(30.5 \text { to } 43.8) \\ & \text { Median } 35.8 \% \\ & \text { IQR } 35.2 \text { to } 36.3 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |

1. $>33.3 \%$ of participants from studies at high risk of bias
2. $>33.3 \%$ participants from studies that were indirect
3. $<33.3 \%$ of studies had $>100$ participants
4. Confidence intervals were non-overlapping
5. $<33.3 \%$ of studies had $>300$ participants

During course of illness: gastrointestinal symptoms: diarrhoea

| Studies | Locatio n | Age | Sample size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Behmadi 2019 <br> Yun 2011 <br> Chang 2014 <br> Sittiwangkul 2013 | Outside Europe | All | $\begin{aligned} & 105 \\ & 83 \\ & 226 \\ & 147 \end{aligned}$ | $\begin{aligned} & 8.6 \%(4.6 \text { to } 15.5) \\ & 14.5 \%(8.5 \text { to } 23.6) \\ & 45.1 \%(38.8 \text { to } 51.7) \\ & 48.3 \%(40.4 \text { to } 56.3) \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

| Studies | Locatio <br> n | Age | Sample size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio $\mathrm{n}$ | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | Median 29.8\% IQR 13.0 to 45.9 |  |  |  |  |  |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Behmadi 2019 <br> Yun 2011 <br> Sittiwangkul 2013 | Outside Europe | All | $\begin{aligned} & 71 \\ & 38 \\ & 61 \end{aligned}$ | $\begin{aligned} & 18.3 \% ~(11.0 \text { to } 28.9) \\ & 18.4 \%(9.2 \text { to } 33.4) \\ & 44.3 \%(32.5 \text { to } 56.7) \\ & \text { Median } 18.4 \% \\ & \text { IQR } 18.4 \text { to } 31.4 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Boudiaf 2016 <br> Zhang 2016 <br> Shamsizadeh 2014 <br> Gamez-Gonzalez 2013 <br> Gorrab 2016 <br> Baker 2009 | Outside Europe | All | $\begin{aligned} & 133 \\ & 518 \\ & 104 \\ & 214 \\ & 146 \\ & 198 \end{aligned}$ | $\begin{aligned} & 15.0 \%(10.0 \text { to } 22.1) \\ & 3.3 \%(2.1 \text { to } 5.2) \\ & 17.3 \%(11.2 \text { to } 25.7) \\ & 15.9 \%(11.6 \text { to } 21.4) \\ & 25.3 \%(19.0 \text { to } 33.0) \\ & 26.3 \%(20.6 \text { to } 32.8) \\ & \text { Median } 16.6 \% \\ & \text { IQR } 15.2 \text { to } 23.3 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| 1. $>33.3 \%$ of participan <br> 2. $>33.3 \%$ participants <br> 3. $<33.3 \%$ of studies h <br> 4. Confidence intervals <br> 5. $<33.3 \%$ of studies h | from stu rom studie >100 pa were non- d >300 pa | ies at that icipan verlap icipan | gh risk of re indirec <br> ng |  |  |  |  |  |  |

During course of illness: gastrointestinal symptoms: diarrhoea and/or abdominal pain

| Studies | Locatio <br> n | Age | Sample size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio $\mathrm{n}$ | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Generini 1997 | Europe (not UK) | All | 73 | 19.2\% (11.8 to 29.7) <br> Median 19.2\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |



1. $>33.3 \%$ of participants from studies at high risk of bias
2. $>33.3 \%$ participants from studies that were indirect
3. $<33.3 \%$ of studies had $>100$ participants
4. Confidence intervals were non-overlapping
5. $<33.3 \%$ of studies had $>300$ participants

During course of illness: gastrointestinal symptoms: diarrhoea and/or vomiting

| Studies | Locatio n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Stemberger Maric 2018 | Europe <br> (not UK) | All | 110 | 46.4\% (37.3 to 55.7) <br> Median 46.4\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Very serious ${ }^{3}$ | Very low |
| Li 2018 <br> Liu 2012 | Outside Europe | $\begin{aligned} & <1 \\ & \text { year } \end{aligned}$ | $\begin{aligned} & 40 \\ & 65 \end{aligned}$ | $\begin{aligned} & 42.5 \%(28.5 \text { to } \\ & 57.8)^{6} \\ & 40.0 \%(29.0 \text { to } 52.1) \\ & \text { Median } 41.3 \% \\ & \text { IQR } 40.6 \text { to } 41.9 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Li 2018 <br> Liu 2012 | Outside Europe | $\geq 1$ <br> year | $\begin{aligned} & 160 \\ & 80 \end{aligned}$ | $\begin{aligned} & 16.9 \% \text { (11.9 to } \\ & 23.4)^{6} \\ & 23.8 \% \text { (15.8 to } 34.1 \text { ) } \\ & \text { Median } 20.4 \% \\ & \text { IQR } 18.6 \text { to } \mathbf{2 2 . 1} \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| 1. $>33.3 \%$ of participants from studies at high risk of bias <br> 2. $>33.3 \%$ participants from studies that were indirect <br> 3. $<33.3 \%$ of studies had $>100$ participants <br> 4. Confidence intervals were non-overlapping <br> 5. $<33.3 \%$ of studies had $>300$ participants <br> 6. Li 2018 groups data for children $<3$ months old and for children over 3 months old <br> 7. Kim 2009 groups data for children $\leq 5$ months old and for children between 5 months to $<5$ years old |  |  |  |  |  |  |  |  |  |


| Studies | $\begin{aligned} & \text { Locatio } \\ & \mathrm{n} \\ & \hline \end{aligned}$ | Age | Sample size | $\begin{aligned} & \text { \% with Symptom } \\ & (95 \% \mathrm{Cl}) \end{aligned}$ | Risk of bias | Indirectness | Inconsisten cy | $\begin{aligned} & \text { Imprecisio } \\ & \mathrm{n} \end{aligned}$ | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

8. Yoon 2016 groups data for children $\leq 6$ months old and for children over 6 months old

During course of illness: gastrointestinal symptoms: hepatomegaly

| Studies | Locatio <br> n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Bai 2017 | Outside Europe | All | 298 | 82.9\% (78.2 to 86.7) <br> Median 82.9\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Bai 2017 | Outside Europe | All | 85 | 62.4\% (51.7 to 71.9) <br> Median 62.4\% <br> IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Shamsizadeh 2014 | Outside <br> Europe | All | 104 | 6.7\% (3.3 to 13.3) Median 6.7\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| 1. $>33.3 \%$ of participants from studies at high risk of bias |  |  |  |  |  |  |  |  |  |
| 2. $>33.3 \%$ participants from studies that were indirect |  |  |  |  |  |  |  |  |  |
| 3. $<33.3 \%$ of studies had $>100$ participants |  |  |  |  |  |  |  |  |  |
| 4. Confidence intervals were non-overlapping |  |  |  |  |  |  |  |  |  |
| 5. $<33.3 \%$ of studies had $>300$ participants |  |  |  |  |  |  |  |  |  |

During course of illness: gastrointestinal symptoms: jaundice

| Studies | Locatio <br> n | Age | Sampl e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Typical + Incomplete Kawasaki disease (AHA criteria or equivalent)

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| Studies | Locatio $\mathrm{n}$ | Age | Sampl e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Generini 1997 <br> Sanchez-Manubens 2016 <br> Tacke 2014 | Europe (not UK) | All | $\begin{aligned} & 73 \\ & 399 \\ & 319 \end{aligned}$ | $\begin{aligned} & \text { 2.7\% (0.75 to 9.45) } \\ & 5.3 \%(3.5 \text { to } 7.9) \\ & 1.9 \%(0.9 \text { to } 4.0) \\ & \text { Median } 2.7 \% \\ & \text { IQR } 2.3 \text { to } 4.0 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Very serious ${ }^{3}$ | Very low |

1. $>33.3 \%$ of participants from studies at high risk of bias
2. $>33.3 \%$ participants from studies that were indirect
3. $<33.3 \%$ of studies had $>100$ participants
4. Confidence intervals were non-overlapping
5. $<33.3 \%$ of studies had $>300$ participants

During course of illness: gastrointestinal symptoms: vomiting

| Studies | Locatio $\mathrm{n}$ | Age | Sampl <br> e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Behmadi 2019 <br> Yun 2011 <br> Sittiwangkul 2013 | Outside Europe | All | $\begin{aligned} & 105 \\ & 83 \\ & 147 \end{aligned}$ | 6.7\% (3.3 to 13.1) <br> 19.3\% (12.2 to 29.0) <br> 10.9\% (6.8 to 17.0) <br> Median 10.9\% <br> IQR 8.8 to 15.1 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Behmadi 2019 <br> Yun 2011 <br> Sittiwangkul 2013 | Outside Europe | All | $\begin{aligned} & 71 \\ & 38 \\ & 61 \end{aligned}$ | $\begin{aligned} & 18.3 \% \text { (11.0 to } 28.9) \\ & 15.8 \%(7.4 \text { to } 30.4) \\ & 8.2 \% \text { ( } 3.6 \text { to } 17.8 \text { ) } \\ & \text { Median } 15.8 \% \\ & \text { IQR } 12.0 \text { to } \mathbf{1 7 . 1} \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Sanchez-Manubens $2016$ | Europe (not UK) | All | 399 | 24.1\% (20.1 to 28.5) Median 24.1\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Very serious ${ }^{3}$ | Very low |


| Studies | Locatio n | Age | Sampl e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Boudiaf 2016 <br> Zhang 2016 <br> Shamsizadeh 2014 <br> Gamez-Gonzalez 2013 <br> Gorrab 2016 <br> Baker 2009 | Outside Europe | All | $\begin{aligned} & 133 \\ & 518 \\ & 104 \\ & 214 \\ & 146 \\ & 198 \end{aligned}$ | $\begin{aligned} & 21.1 \%(15.0 \text { to } 28.7) \\ & 2.3 \% ~(1.3 \text { to } 4.0) \\ & 23.1 \%(16.0 \text { to } 32.1) \\ & 20.1 \%(15.3 \text { to } 26.0) \\ & 27 \%(20.8 \text { to } 35.1) \\ & 44.4 \%(37.7 \text { to } 51.4) \\ & \text { Median } 22.1 \% \\ & \text { IQR } 20.4 \text { to } \mathbf{2 6 . 0} \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| 1. $>33.3 \%$ of participants from studies at high risk <br> 2. $>33.3 \%$ participants from studies that were indire <br> 3. $<33.3 \%$ of studies had $>100$ participants <br> 4. Confidence intervals were non-overlapping <br> 5. $<33.3 \%$ of studies had $>300$ participants |  |  |  |  |  |  |  |  |  |

During course of illness: irritability

| Studies | Locatio n | Age | Sampl e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Manlhiot 2012 <br> Bai 2017 <br> Sittiwangkul 2013 | Outside Europe | All | $\begin{aligned} & 738 \\ & 298 \\ & 147 \end{aligned}$ | 13.4\% (11.1 to 16.1) 89.3\% (85.2 to 92.3) 82.3\% (75.4 to 87.6) <br> Median 82.3\% <br> IQR 47.9 to 85.8 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Manlhiot 2012 <br> Bai 2017 <br> Sittiwangkul 2013 | Outside Europe | All | $\begin{aligned} & 217 \\ & 85 \\ & 61 \end{aligned}$ | $\begin{aligned} & 12.9 \%(9.1 \text { to } 18.0) \\ & 76.5 \%(66.4 \text { to } 84.2) \\ & 73.8 \%(61.6 \text { to } 83.2) \end{aligned}$ Median 73.8\% | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |


| Studies | Locatio n | Age | Sampl e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | IQR 43.4 to 75.2 |  |  |  |  |  |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Sanchez-Manubens $2016$ | Europe <br> (not UK) | All | 399 | $29.8 \% \text { (25.6 to } 34.5 \text { ) }$ <br> Median 29.8\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Very serious ${ }^{3}$ | Very low |
| Saundankar 2014 <br> Shamsizadeh 2014 <br> Baker 2009 | Outside Europe | All | $\begin{aligned} & 353 \\ & 104 \\ & 198 \end{aligned}$ | $\begin{aligned} & 85.0 \%(80.9 \text { to } 88.3) \\ & 26.0 \%(18.5 \text { to } 35.1) \\ & 49.5 \%(42.6 \text { to } 56.4) \end{aligned}$ <br> Median 49.5\% <br> IQR 37.8 to 67.3 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| 1. $>33.3 \%$ of participants from studies at high risk of bias <br> 2. $>33.3 \%$ participants from studies that were indirect <br> 3. $<33.3 \%$ of studies had $>100$ participants <br> 4. Confidence intervals were non-overlapping <br> 5. $<33.3 \%$ of studies had $>300$ participants |  |  |  |  |  |  |  |  |  |

During course of illness: lassitude

| Studies | Locatio n | Age | Sample size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Manlhiot 2012 <br> Bai 2017 | Outside Europe | All | $\begin{aligned} & 738 \\ & 298 \end{aligned}$ | $\begin{aligned} & 5.4 \% \text { ( } 4.0 \text { to } 7.3 \text { ) } \\ & 88.6 \%(84.5 \text { to } 91.7) \\ & \text { Median } 47.0 \% \\ & \text { IQR } 26.2 \text { to } 67.8 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Manlhiot 2012 <br> Bai 2017 | Outside Europe | All | $\begin{aligned} & 217 \\ & 85 \end{aligned}$ | $\begin{aligned} & 6.0 \% \text { ( } 3.5 \text { to } 10.0 \text { ) } \\ & 78.8 \%(69.0 \text { to } 86.2) \\ & \text { Median } 42.4 \% \\ & \text { IQR } 24.2 \text { to } 60.6 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

| Studies | Locatio <br> n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Baker 2009 | Outside <br> Europe | All | 198 | 18.7\% (13.9 to 24.7) <br> Median 18.7\% <br> IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| 1. $>33.3 \%$ of participants from studies at high risk of bias |  |  |  |  |  |  |  |  |  |
| 2. $>33.3 \%$ participants from studies that were indirect |  |  |  |  |  |  |  |  |  |
| 3. $<33.3 \%$ of studies had $>100$ participants |  |  |  |  |  |  |  |  |  |
| 4. Confidence intervals were non-overlapping |  |  |  |  |  |  |  |  |  |
| 5. $<33.3 \%$ of studies had $>300$ participants |  |  |  |  |  |  |  |  |  |

During course of illness: limb sclerosis

| Studies | Locatio <br> n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Bai 2017 | Outside Europe | All | 298 | 37.9\% (32.6 to 43.6) <br> Median 37.9\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Bai 2017 | Outside Europe | All | 85 | 63.5\% (52.9 to 73.0) <br> Median 63.5\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |

1. $>33.3 \%$ of participants from studies at high risk of bias
2. $>33.3 \%$ participants from studies that were indirect
3. $<33.3 \%$ of studies had $>100$ participants
4. Confidence intervals were non-overlapping
5. $<33.3 \%$ of studies had $>300$ participants

During course of illness: oral changes: red, dry or cracked lips

| Studies | Location | Age | Sampl e size | $\begin{aligned} & \text { \% with Symptom } \\ & \text { (95\% CI) } \end{aligned}$ | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Tang 2016 | Outside Europe | All | 716 | $90.9 \% \text { (88.6 to 92.8) }$ <br> Median 90.9\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Tang 2016 | Outside Europe | All | 300 | $65.0 \% \text { (59.4 to } 70.2 \text { ) }$ <br> Median 65.0\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Sanchez-Manubens 2016 | Europe <br> (not UK) | All | 399 | $65.7 \% \text { (60.9 to } 70.2 \text { ) }$ <br> Median 65.7\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Ruan 2013 | Outside Europe | $\begin{aligned} & <1 \\ & \text { year } \end{aligned}$ | 49 | $\begin{aligned} & 73.5 \%(59.7 \text { to } \\ & 83.8)^{6} \\ & \text { Median } 73.5 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| Ruan 2013 | Outside Europe | $\geq 1$ <br> year | 1160 | $\begin{aligned} & 89.9 \%(88.1 \text { to } \\ & 91.5)^{6} \end{aligned}$ <br> Median 89.9\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| Sun 2018 | Outside Europe | All | 1008 | $83.6 \% \text { (81.2 to 85.8) }$ <br> Median 83.6\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | $\mathrm{n} / \mathrm{a}$ | Serious ${ }^{5}$ | Very low |
| 1. $>33.3 \%$ of partic <br> 2. $>33.3 \%$ participa <br> 3. $<33.3 \%$ of studie <br> 4. Confidence inter <br> 5. $<33.3 \%$ of studie <br> 6. Ruan 2013 group | s from stud from studie > $>100$ par were non-o d >300 par ta for child | $s$ at hi hat we ipants erlappi ipants < $<6$ | risk of indirect | ias <br> and for children between | 6 month | to 5 years old |  |  |  |

At presentation: oral changes: red, dry or cracked lips

| Studies | Locatio <br> n | Age | Sample size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio $\mathrm{n}$ | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Moore 2014 | UK | All | 74 | $31.1 \%$ (21.7 to 42.3) | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| 1. $>33.3 \%$ of participants from studies at high risk of bias <br> 2. $>33.3 \%$ participants from studies that were partially direct <br> 3. $<33.3 \%$ of studies had $>100$ participants |  |  |  |  |  |  |  |  |  |

At presentation: oral changes: redness in mouth

| Studies | Locatio n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Moore 2014 | UK | All | 74 | $\begin{aligned} & \text { 23.0\% (14.9 to 33.8) } \\ & \text { Median } 23.0 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |

1. $>33.3 \%$ of participants from studies at high risk of bias
2. $>33.3 \%$ participants from studies that were partially direct
3. $<33.3 \%$ of studies had $>100$ participants

During course of illness: oral changes: strawberry tongue

| Studies | Locatio n | Age | Sampl e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Tang 2016 | Outside Europe | All | 716 | $76.8 \% \text { (73.6 to } 79.8 \text { ) }$ <br> Median 76.8\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Tang 2016 | Outside Europe | All | 300 | 45.3\% (39.8 to 51.0) <br> Median 45.3\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |


| Studies | Locatio $\mathrm{n}$ | Age | Sampl <br> e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Sanchez-Manubens $2016$ | Europe <br> (not UK) | All | 399 | 55.6\% (50.7 to 60.4) <br> Median 55.6\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| Ruan 2013 | Outside Europe | $\begin{aligned} & <1 \\ & \text { year } \end{aligned}$ | 49 | $\begin{aligned} & 34.7 \% ~(22.9 \text { to } \\ & 48.7)^{6} \\ & \text { Median } 34.7 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| Ruan 2013 | Outside Europe | $\begin{aligned} & \geq 1 \\ & \text { year } \end{aligned}$ | 1160 | $\begin{aligned} & 58.9 \%(56.0 \text { to } \\ & 61.7)^{6} \\ & \text { Median } 58.9 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| Sun 2018 | Outside Europe | All | 1008 | 71.6\% (68.8 to 74.3) <br> Median 71.6\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| 1. $>33.3 \%$ of participants from studies at high risk of |  |  |  |  |  |  |  |  |  |
| 2. $>33.3 \%$ participants from studies that were indirect |  |  |  |  |  |  |  |  |  |
| 3. $<33.3 \%$ of studies had $>100$ participants |  |  |  |  |  |  |  |  |  |
| 4. Confidence intervals were non-overlapping |  |  |  |  |  |  |  |  |  |
| 5. $<33.3 \%$ of studies had $>300$ participants |  |  |  |  |  |  |  |  |  |
| 6. Ruan 2013 groups data for children <6 months old and for children between 6 months to 5 years old |  |  |  |  |  |  |  |  |  |

At presentation: oral changes: strawberry tongue

| Studies | Locatio n | Age | Sample size | \% with Symptom $\text { ( } 95 \% \mathrm{CI} \text { ) }$ | Risk of bias | Indirectness | Inconsisten <br> cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Moore 2014 | UK | All | 74 | 24.3\% (16.0 to 35.2) <br> Median 24.3\% IQR - | Very serious ${ }^{1}$ | Serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very Iow |
| 1. $>33.3 \%$ of participants from studies at high risk of bias <br> 2. $>33.3 \%$ participants from studies that were partially direct |  |  |  |  |  |  |  |  |  |


| Studies | Locatio | Age | Sample size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

3. $<33.3 \%$ of studies had $>100$ participants

During course of illness: otitis media

| Studies | Locatio <br> n | Age | Sample <br> size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Manlhiot 2012 | Outside Europe | All | 738 | $8.0 \%(6.25 \text { to } 10.2)$ <br> Median 8.0\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Manlhiot 2012 | Outside Europe | All | 217 | $7.8 \%$ ( 5.0 to 12.2) <br> Median 7.8\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Serious ${ }^{5}$ | Very low |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Generini 1997 <br> Tacke 2014 | Europe <br> (not UK) | All | $\begin{aligned} & 73 \\ & 319 \end{aligned}$ | $\begin{aligned} & 1.4 \%(0.24 \text { to } 7.4) \\ & 11.9 \%(8.8 \text { to } 15.9) \\ & \text { Median } 6.7 \% \\ & \text { IQR } 4.0 \text { to } 9.3 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Very serious ${ }^{3}$ | Very low |

1. $>33.3 \%$ of participants from studies at high risk of bias
2. $>33.3 \%$ participants from studies that were indirect
3. $<33.3 \%$ of studies had $>100$ participants
4. Confidence intervals were non-overlapping
5. $<33.3 \%$ of studies had $>300$ participants

During course of illness: respiratory symptoms

| Studies | Locatio <br> n | Age | Sample size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Typical Kawasaki disease (AHA criteria or equivalent)

Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

| Studies | Locatio <br> n | Age | Sample <br> size | \% with Symptom $\text { ( } 95 \% \mathrm{CI} \text { ) }$ | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Manlhiot 2012 <br> Yun 2011 <br> Sittiwangkul 2013 | Outside Europe | All | $\begin{aligned} & 738 \\ & 83 \\ & 147 \end{aligned}$ | $\begin{aligned} & 32.8 \%(29.5 \text { to } 36.3) \\ & 55.4 \%(44.7 \text { to } 65.6) \\ & 38.8 \%(31.3 \text { to } 46.8) \\ & \text { Median } 38.8 \% \\ & \text { IQR } 35.8 \text { to } 47.1 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Manlhiot 2012 <br> Yun 2011 <br> Falcini 2012 <br> Sittiwangkul 2013 | Outside Europe | All | $\begin{aligned} & 217 \\ & 38 \\ & 228 \\ & 61 \end{aligned}$ | $\begin{aligned} & 35.5 \%(29.4 \text { to } 42.1) \\ & 52.6 \%(37.3 \text { to } 67.5) \\ & 6.1 \%(3.7 \text { to } 10.0) \\ & 39.3 \%(28.1 \text { to } 51.9) \\ & \text { Median } 37.4 \% \\ & \text { IQR } 28.2 \text { to } 42.6 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Stemberger Maric 2018 | Europe (not UK) | All | 110 | $54.5 \%$ (45.2 to 63.5) | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Very serious ${ }^{3}$ | Very low |
| Saundankar 2014 <br> Zhang 2016 <br> Chen 2016 | Outside Europe | All | $\begin{aligned} & 353 \\ & 518 \\ & 2304 \end{aligned}$ | 33.1\% (28.4 to 38.2) <br> 35.9\% (31.9 to 40.1) <br> 31.0\% (29.2 to 33.0) <br> Median 34.5\% <br> IQR 32.6 to 40.6 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |

1. $>33.3 \%$ of participants from studies at high risk of bias
2. $>33.3 \%$ participants from studies that were indirect
3. $<33.3 \%$ of studies had $>100$ participants
4. Confidence intervals were non-overlapping
5. $<33.3 \%$ of studies had $>300$ participants

During course of illness: respiratory symptoms: infection

| Studies | Locatio <br> n | Age | Sample size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

Typical + Incomplete Kawasaki disease (AHA criteria or equivalent)

| Studies | Locatio $\mathrm{n}$ | Age | Sample size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Generini 1997 | Europe (not UK) | All | 73 | $17.8 \% \text { (10.7 to } 28.1 \text { ) }$ <br> Median 17.8\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| $\begin{aligned} & \text { Li } 2018 \\ & \text { Liu } 2012 \end{aligned}$ | Outside Europe | $\begin{aligned} & <1 \\ & \text { year } \end{aligned}$ | $\begin{aligned} & 40 \\ & 65 \end{aligned}$ | $\begin{aligned} & 55.0 \% \text { ( } 39.8 \text { to } \\ & 69.3)^{6} \\ & 60.0 \% \text { ( } 47.9 \text { to } 71.0 \text { ) } \\ & \text { Median } 57.5 \% \\ & \text { IQR } 56.3 \text { to } 58.8 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Li 2018 Liu 2012 | Outside Europe | $\begin{aligned} & \geq 1 \\ & \text { year } \end{aligned}$ | $\begin{aligned} & 160 \\ & 80 \end{aligned}$ | ```21.3% (15.6 to 28.2) 43.8% (33.4 to 54.7) Median 32.6% IQR 26.9 to 38.2``` | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |

1. $>33.3 \%$ of participants from studies at high risk of bias
2. $>33.3 \%$ participants from studies that were indirect
3. $<33.3 \%$ of studies had $>100$ participants
4. Confidence intervals were non-overlapping
5. $<33.3 \%$ of studies had $>300$ participants
6. Li 2018 groups data for children $<3$ months old and for children over 3 months old

During course of illness: respiratory symptoms: sputum

| Studies | Locatio <br> n | Age | $\begin{aligned} & \text { Sample } \\ & \text { size } \end{aligned}$ | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Yun 2011 | Outside Europe | All | 83 | 10.8\% (5.8 to 19.3) <br> Median 10.8\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Yun 2011 | Outside Europe | All | 38 | 23.7\% (13.0 to 39.2) <br> Median 23.7\% | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |


| Studies | Locatio <br> n | Age | Sample size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  | IQR - |  |  |  |  |  |
| 1. $>33.3 \%$ of participants from studies at high risk of bias |  |  |  |  |  |  |  |  |  |
| 2. $>33.3 \%$ participants from studies that were indirect |  |  |  |  |  |  |  |  |  |
| 3. $<33.3 \%$ of studies had $>100$ participants |  |  |  |  |  |  |  |  |  |
| 4. Confidence intervals were non-overlapping |  |  |  |  |  |  |  |  |  |
| 5. $<33.3 \%$ of studies had $>300$ participants |  |  |  |  |  |  |  |  |  |

During course of illness: respiratory symptoms: cough

| Studies | Locatio <br> n | Age | Sample size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio $\mathrm{n}$ | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Yun 2011 Chang 2014 | Outside Europe | All | $\begin{aligned} & 83 \\ & 226 \end{aligned}$ | $\begin{aligned} & 47.0 \%(36.6 \text { to } 57.6) \\ & 69.5 \%(63.2 \text { to } 75.1) \\ & \text { Median } 58.3 \% \\ & \text { IQR 52.6 to } 63.9 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Tacke 2014 | Europe (not UK) | All | 319 | $\begin{aligned} & 16.9 \%(13.2 \text { to } 21.4) \\ & \text { Median } 16.9 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Not serious | Very serious ${ }^{3}$ | Very low |
| Yun 2011 | Outside Europe | All | 38 | $44.7 \% \text { (30.2 to 60.3) }$ <br> Median 44.7\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Shamsizadeh 2014 <br> Baker 2009 <br> Zhang 2016 | Outside Europe | All | $\begin{aligned} & 104 \\ & 198 \\ & 518 \end{aligned}$ | $\begin{aligned} & 12.5 \%(7.5 \text { to } 20.2) \\ & 27.8 \%(22.0 \text { to } 34.4) \\ & 12.2 \%(9.62 \text { to } 15.3) \end{aligned}$ <br> Median 12.5\% <br> IQR 12.4 to 20.2 | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |

1. $>33.3 \%$ of participants from studies at high risk of bias
2. $>33.3 \%$ participants from studies that were indirect

| Studies | Locatio n | Age | Sample size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

3. $<33.3 \%$ of studies had $>100$ participants
4. Confidence intervals were non-overlapping
5. $<33.3 \%$ of studies had $>300$ participants

During course of illness: respiratory symptoms: rhinorrhoea

| Studies | Locatio <br> n | Age | Sample size | \% with Symptom $(95 \% \mathrm{Cl})$ | Risk of bias | Indirectness | Inconsisten cy | Imprecisio $\mathrm{n}$ | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Yun 2011 Chang 2014 | Outside Europe | All | $\begin{aligned} & 83 \\ & 226 \end{aligned}$ | $\begin{aligned} & 30.1 \%(21.3 \text { to } 40.7) \\ & 57.5 \%(51.0 \text { to } 63.8) \\ & \text { Median } 43.8 \% \\ & \text { IQR } \mathbf{3 7 . 0} \text { to } 50.7 \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Yun 2011 | Outside Europe | All | 38 | $\begin{aligned} & 31.6 \%(19.1 \text { to } 47.5) \\ & \text { Median } 31.6 \% \\ & \text { IQR - } \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Baker 2009 <br> Shamsizadeh 2014 | Outside Europe | All | $\begin{aligned} & 198 \\ & 104 \end{aligned}$ | $\begin{aligned} & \text { 18.7\% (13.9 to } 24.7) \\ & 29.8 \%(21.9 \text { to } 39.2) \\ & \text { Median } 24.3 \% \\ & \text { IQR } 21.5 \text { to } \mathbf{2 7 . 0} \end{aligned}$ | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | Serious ${ }^{4}$ | Serious ${ }^{5}$ | Very low |
| 1. $>33.3 \%$ of participants from studies at high risk of bias |  |  |  |  |  |  |  |  |  |
| 2. $>33.3 \%$ participa 3. 4. 5. 5. | rom studie >100 par were non >300 pa | that ticipan verlap ticipan | re indirec <br> ng |  |  |  |  |  |  |

During course of illness: respiratory symptoms: pharyngitis

| Studies | Locatio <br> n | Age | Sampl <br> e size | \% with Symptom (95\% CI) | Risk of bias | Indirectness | Inconsisten cy | Imprecisio <br> n | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Typical + Incomplete Kawasaki disease (AHA criteria or equivalent) |  |  |  |  |  |  |  |  |  |
| Sanchez-Manubens $2016$ | Europe (not UK) | All | 399 | 49.6\% (44.8 to 54.5) <br> Median 49.6\% IQR - | Very serious ${ }^{1}$ | Very serious ${ }^{2}$ | n/a | Very serious ${ }^{3}$ | Very low |
| 1. $>33.3 \%$ of participants from studies at high risk of bias <br> 2. $>33.3 \%$ participants from studies that were indirect <br> 3. $<33.3 \%$ of studies had $>100$ participants <br> 4. Confidence intervals were non-overlapping <br> 5. $<33.3 \%$ of studies had $>300$ participants |  |  |  |  |  |  |  |  |  |

Case control

| No. of studies | Study design | Sample <br> size | Sensitivity (95\%CI) | Specificity (95\%CI) | $\begin{aligned} & \text { Effect size } \\ & (95 \% \mathrm{CI}) \end{aligned}$ | Risk of bias | Indirectness | Inconsistency | Imprecision | Quality |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| BCG scar activation <br> Typical + incomplete Kawasaki disease, Outside Europe, all ages ${ }^{3}$ |  |  |  |  |  |  |  |  |  |  |
| $\begin{aligned} & 1 \text { (Loh } \\ & 2019) \end{aligned}$ | Case control | 370 | $\begin{aligned} & 0.43 \text { ( } 0.37 \text {, } \\ & 0.49 \text { ) } \end{aligned}$ | $\begin{aligned} & 0.90(0.82, \\ & 0.95) \end{aligned}$ | $\begin{aligned} & \mathrm{LR}+4.31 \\ & (2.29,8.14) \end{aligned}$ | Very serious ${ }^{1}$ | Serious | N/A | Not serious | Very low |
|  |  |  |  |  | $\begin{aligned} & \text { LR- } 0.64 \\ & (0.56,0.72) \end{aligned}$ | Very serious ${ }^{1}$ | Serious | N/A | Not serious | Very low |
|  |  | 1. $>33.3 \%$ of studies were at high risk of bias <br> 2. $>33.3 \%$ of studies were partially applicable <br> 3. Study presented sensitivity/specificity for under/over 1 s separately, but data was not available to construct $2^{*} 2$ tables for these subgroups and partial data was internally inconsistent |  |  |  |  |  |  |  |  |

## Appendix G - Excluded studies

## Clinical studies

| Study | Code [Reason] |
| :--- | :--- |
| Abuhammour, W. and Yousef, N. (2008) Incomplete <br> Kawasaki disease: Experience wwith 14 patients with <br> cardiac complications. Journal of Pediatric Infectious <br> Diseases 3(2): $91-95$ | - Study conducted outside Europe and |
| had fewer than 100 patients |  |

FINAL

## Study

Garcia Rodriguez, F., Flores Pineda, A. D. J., Villarreal Trevino, A. V. et al. (2016) Kawasaki disease at a pediatric hospital in Mexico. Boletin Medico del Hospital Infantil de Mexico 73(3): 166-173
Grasa, C. D., Fernandez-Cooke, E., SanchezManubens, J. et al. (2019) Kawasaki disease in infants 3 months of age and younger: A multicentre Spanish study. Annals of the Rheumatic Diseases 78(2): 289-290
Guleria, Sandesh, Bhattarai, Dharmagat, Pilania, Rakesh Kumar et al. (2018) Koplik Spots: A Physical Finding That Should Never Be Missed in Children With Suspected Kawasaki Disease. Journal of clinical rheumatology: practical reports on rheumatic \& musculoskeletal diseases
Hao, Shiying, Jin, Bo, Tan, Zhou et al. (2016) A Classification Tool for Differentiation of Kawasaki Disease from Other Febrile Illnesses. The Journal of pediatrics 176: 114-120.e8

Heuclin, T., Dubos, F., Hue, V. et al. (2009) Increased Detection Rate of Kawasaki Disease Using New Diagnostic Algorithm, Including Early Use of Echocardiography. Journal of Pediatrics 155(5): 695
Hua, Wang, Ma, Feiyue, Wang, Ying et al. (2019) A new scoring system to predict Kawasaki disease with coronary artery lesions. Clinical rheumatology 38(4): 1099-1107
Huang, W. C., Huang, L. M., Chang, I. S. et al. (2009) Epidemiologic features of Kawasaki disease in Taiwan, 2003 2006. Pediatrics 123(3): e401-e405
Huang, Xijing, Huang, Ping, Zhang, Li et al. (2015) Influenza infection and Kawasaki disease. Revista da Sociedade Brasileira de Medicina Tropical 48(3): 243-8

Huang, Ying-Hsien, Lin, Kuan-Miao, Ho, Shu-Chen et al. (2019) Increased Incidence of Kawasaki Disease in Taiwan in Recent Years: A 15 Years Nationwide Population-Based Cohort Study. Frontiers in pediatrics 7: 121
Jakob, A., Whelan, J., Kordecki, M. et al. (2016) Kawasaki Disease in Germany: A Prospective, Population-based Study Adjusted for Underreporting. Pediatric Infectious Disease Journal 35(2): 129-134
Jakob, Andre, von Kries, Rudiger, Horstmann, Judith et al. (2018) Failure to Predict High-risk Kawasaki Disease Patients in a Population-based Study Cohort in Germany. The Pediatric infectious disease journal 37(9): 850-855
James, R. and Burgner, D. (2015) Orange-brown chromonychia in Kawasaki disease. Archives of Disease in Childhood 100(9): 872

## Code [Reason]

- This study is not written in English.
- Study happened within Europe but outside of the UK and had fewer than 50 patients with Kawasaki disease.
- This is a case study.
- This study assesses an algorithm that requires knowledge of laboratory investigations. It does not have separate data that assesses the signs and symptoms alone.
- Study happened within Europe but outside of the UK and had fewer than 50 patients with Kawasaki disease.
- No data on the signs and symptoms.
- This is an epidemiological study about incidence rates. Signs and symptoms were beyond the scope of the study.
- Study happened outside Europe and had fewer than 100 patients
[This study looks at a small subset (45) of the 1,053 Kawasaki disease cases mentioned in the abstract.]
- This is an epidemiological study about incidence rates. Signs and symptoms were beyond the scope of the study.
- No data on the signs and symptoms.
- No data on the signs and symptoms.
- This is a case study.


## Study

Juan, Chien-Chang, Hwang, Betau, Lee, Pi-Chang et al. (2007) The clinical manifestations and risk factors of a delayed diagnosis of Kawasaki disease. Journal of the Chinese Medical Association: JCMA 70(9): 374-9

Kang, Hye Jin; Kim, Gee Na; Kil, Hong Ryang (2013) Changes of clinical characteristics and outcomes in patients with Kawasaki disease over the past 7 years in a single center study. Korean journal of pediatrics 56(9): 389-95
Kayiran, Sinan Mahir; Dindar, Aygun; Gurakan, Berkan (2010) An evaluation of children with Kawasaki disease in Istanbul: a retrospective follow-up study. Clinics (Sao Paulo, Brazil) 65(12): 1261-5
Kim, Gi Beom, Han, Ji Whan, Park, Yong Won et al. (2014) Epidemiologic features of Kawasaki disease in South Korea: data from nationwide survey, 2009-2011. The Pediatric infectious disease journal 33(1): 24-7 Kim, Jae-Jung, Hong, Young Mi, Yun, Sin Weon et al. (2012) Assessment of risk factors for Korean children with Kawasaki disease. Pediatric cardiology 33(4): 51320
Kim, Taeyeun, Choi, Wooksun, Woo, Chan-Wook et al. (2007) Predictive risk factors for coronary artery abnormalities in Kawasaki disease. European journal of pediatrics 166(5): 421-5
Kyung Sim, B., Park, H., Kim, J. J. et al. (2019) Assessment of the clinical heterogeneity of Kawasaki disease using genetic variants of BLK and FCGR2A. Korean Circulation Journal 49(1): 99-108
Lin, Ming-Chih, Lai, Mei-Shu, Jan, Sheng-Ling et al. (2015) Epidemiologic features of Kawasaki disease in acute stages in Taiwan, 1997-2010: effect of different case definitions in claims data analysis. Journal of the Chinese Medical Association: JCMA 78(2): 121-6
Lin, R. Y. and Krata, L. M. (2010) Trends in kawasaki disease hospitalizations: New York state 1990-2009. Internet Journal of Asthma, Allergy and Immunology 8(1)

Ling, Xuefeng B., Kanegaye, John T., Ji, Jun et al.
(2013) Point-of-care differentiation of Kawasaki disease from other febrile illnesses. The Journal of pediatrics 162(1): 183-188.e3

Ling, Xuefeng B., Lau, Kenneth, Kanegaye, John T. et al. (2011) A diagnostic algorithm combining clinical and molecular data distinguishes Kawasaki disease from other febrile illnesses. BMC medicine 9: 130

## Code [Reason]

- Study happened outside Europe and had fewer than 100 patients
- No data on the signs and symptoms.
- Study happened outside Europe and had fewer than 100 patients
- This is an epidemiological study about incidence rates. Signs and symptoms were beyond the scope of the study.
- No data on the signs and symptoms.
- No data on the signs and symptoms.
- No data on the signs and symptoms.
- This is an epidemiological study about incidence rates. Signs and symptoms were beyond the scope of the study.
- This is an epidemiological study about incidence rates. Signs and symptoms were beyond the scope of the study. [Although some data was collected on conjunctivitis, the intention of the database was not to document medical records. The aim of the database was to measure the costs of care.]
- This study is about the development of a diagnostic algorithm for Kawasaki disease. Details of how it works is not provided. There is no useful data such as diagnostic yield.
- This study assesses an algorithm that requires knowledge of laboratory investigations. It does not have separate data that assesses the signs and symptoms alone.

FINAL

## Study

Lue, Hung-Chi, Chen, Lei-Ru, Lin, Ming-Tai et al. (2014) Epidemiological features of Kawasaki disease in Taiwan, 1976-2007: results of five nationwide questionnaire hospital surveys. Pediatrics and neonatology 55(2): 92-6
Manlhiot, C., O'Shea, S., Bernknopf, B. et al. (2018) Epidemiology of Kawasaki Disease in Canada 2004 to 2014: Comparison of Surveillance Using Administrative Data vs Periodic Medical Record Review. Canadian Journal of Cardiology 34(3): 330-332
Mao, Youying, Yin, Lei, Xia, Hui et al. (2016) Incidence and clinical features of paediatric vasculitis in Eastern China: 14-year retrospective study, 1999-2013. The Journal of international medical research 44(3): 710-7 Muta, Hiromi, Ishii, Masahiro, lemura, Motofumi et al. (2007) Effect of revision of Japanese diagnostic criterion for fever in Kawasaki disease on treatment and cardiovascular outcome. Circulation journal: official journal of the Japanese Circulation Society 71(11): 1791-3
Nakamura, Yosikazu, Yashiro, Mayumi, Sadakane, Atsuko et al. (2009) Six principal symptoms and coronary artery sequelae in Kawasaki disease. Pediatrics international: official journal of the Japan Pediatric Society 51(5): 705-8
Nakamura, Yosikazu, Yashiro, Mayumi, Uehara, Ritei et al. (2008) Epidemiologic features of Kawasaki disease in Japan: results from the nationwide survey in 2005-2006. Journal of epidemiology 18(4): 167-72
No, Sol Ji, Kim, Dong Ouk, Choi, Kyong Min et al.
(2013) Do predictors of incomplete Kawasaki disease exist for infants? Pediatric cardiology 34(2): 286-90
Ozeki, Y., Yamada, F., Saito, A. et al. (2018) Epidemiologic features of Kawasaki disease distinguished by seasonal variation: an age-specific analysis. Annals of Epidemiology 28(11): 796-800
Ozeki, Yukie, Yamada, Fumiya, Kishimoto, Tsuyoshi et al. (2017) Epidemiologic features of Kawasaki disease: Winter versus summer. Pediatrics international: official journal of the Japan Pediatric Society 59(7): 821-825
Ozen, S., Bakkaloglu, A., Dusunsel, R. et al. (2007) Childhood vasculitides in Turkey: A nationwide survey. Clinical Rheumatology 26(2): 196-200
Rezai, Mohammad Sadegh and Shahmohammadi, Soheila (2014) Erythema at BCG Inoculation Site in Kawasaki Disease Patients. Materia socio-medica 26(4): 256-60
Rhim, Jung-Woo, Youn, You-Sook, Han, Ji-Whan et al. (2014) Changes in Kawasaki disease during 2 decades at a single institution in Daejeon, Korea. The Pediatric infectious disease journal 33(4): 372-5

## Code [Reason]

- This is an epidemiological study about incidence rates. Signs and symptoms were beyond the scope of the study.
- This is an epidemiological study about incidence rates. Signs and symptoms were beyond the scope of the study.
- This is an epidemiological study about incidence rates. Signs and symptoms were beyond the scope of the study.
- No data on the signs and symptoms.
- No data on the signs and symptoms.
- This is an epidemiological study about incidence rates. Signs and symptoms were beyond the scope of the study.
- No data on the signs and symptoms.
- This is an epidemiological study about incidence rates. Signs and symptoms were beyond the scope of the study.
- This is an epidemiological study about incidence rates. Signs and symptoms were beyond the scope of the study.
- This is an epidemiological study about incidence rates. Signs and symptoms were beyond the scope of the study.
- This study is a review article. The reference list was checked to ensure that we have included studies that meet our inclusion and exclusion criteria.
- No data on the signs and symptoms.


## Study

Saguil, A., Fargo, M., Grogan, S. et al. (2015) Diagnosis and management of kawasaki disease. American Family Physician 91(6): 365-371
Salo, Eeva, Griffiths, Elizabeth P., Farstad, Teresa et al. (2012) Incidence of Kawasaki disease in northern European countries. Pediatrics international: official journal of the Japan Pediatric Society 54(6): 770-2
Shapiro, Cal, Maenz, Lynn, Hossain, Alomgir et al. (2007) Onset to first visit intervals in childhood rheumatic diseases. The Journal of rheumatology 34(9): 1913-7
Singh, S., Gupta, M. K., Bansal, A. et al. (2007) A comparison of the clinical profile of Kawasaki disease in children from Northern India above and below 5 years of age. Clinical and experimental rheumatology 25(4): 6547
Singh, Surjit, Agarwal, Sikha, Bhattad, Sagar et al.
(2016) Kawasaki disease in infants below 6 months: a clinical conundrum? International journal of rheumatic diseases 19(9): 924-8
Singh, Surjit, Gupta, Aman, Jindal, Ankur Kumar et al. (2018) Pulmonary presentation of Kawasaki disease-A diagnostic challenge. Pediatric pulmonology 53(1): 103107

Song, Dooll, Yeo, Yunku, Ha, KeeSoo et al. (2009) Risk factors for Kawasaki disease-associated coronary abnormalities differ depending on age. European journal of pediatrics 168(11): 1315-21
Sudo, Daisuke, Monobe, Yoshiro, Yashiro, Mayumi et al. (2012) Coronary artery lesions of incomplete Kawasaki disease: a nationwide survey in Japan. European journal of pediatrics 171(4): 651-6
Sun, L., Tang, Y., Wang, Y. et al. (2018) Changes in profiles of kawasaki disease noted over time in Suzhou, China. Cardiology (Switzerland) 141(1): 69-70
Sundberg, Melissa, Perron, Catherine O., Kimia, Amir et al. (2018) A method to identify pediatric high-risk diagnoses missed in the emergency department.
Diagnosis (Berlin, Germany) 5(2): 63-69
Taddio, Andrea, Rossi, Eleonora Dei, Monasta, Lorenzo et al. (2017) Describing Kawasaki shock syndrome: results from a retrospective study and literature review. Clinical rheumatology 36(1): 223-228
Takahashi, Takuto, Sakakibara, Hiroshi, Morikawa, Yoshihiko et al. (2015) Development of coronary artery lesions in indolent Kawasaki disease following initial spontaneous defervescence: a retrospective cohort study. Pediatric rheumatology online journal 13(1): 44
Thapa, R. and Pal, P. (2010) Transverse orange-brown chromonychia in Kawasaki disease. International Journal of Dermatology 49(2): 227-228

## Code [Reason]

- This is a narrative review. There is no study data.
- This is an epidemiological study about incidence rates. Signs and symptoms were beyond the scope of the study.
- No data on the signs and symptoms.
- Study happened outside Europe and had fewer than 100 patients
- Study happened outside Europe and had fewer than 100 patients
- Study happened outside Europe and had fewer than 100 patients
[This study took place in India and it focused on 11 children.]
- No data on the signs and symptoms.
- No data on the signs and symptoms.
- Duplicate reference
- This study is about computer software
- No data on the signs and symptoms.
- No data on the signs and symptoms.
- Study happened outside Europe and had fewer than 100 patients


## Study

Tomita, Yasuhiko, Shimaya, Maki, Yamaura, Yasuko et al. (2018) Kawasaki disease: Epidemiological differences between past and recent periods, and implications of distribution dynamism. Pediatrics international: official journal of the Japan Pediatric Society 60(4): 349-356
Tona, Risa, Shinohara, Shogo, Fujiwara, Keizo et al. (2014) Risk factors for retropharyngeal cellulitis in Kawasaki disease. Auris, nasus, larynx 41(5): 455-8
Topcu, S., Dogan, O. A., Oz, N. et al. (2014) Clinical evaluations of 49 cases with kawasaki disease: A retrospective cohort study. Cocuk Enfeksiyon Dergisi 8(2): 64-70
Tseng C-F, Fu Y-C, Fu L-S, et al. Clinical spectrum of Kawasaki disease in infants. Chinese Medical Journal 2001;64(3):168-73.
Tulloh, Robert M. R., Mayon-White, Richard, Harnden, Anthony et al. (2018) Kawasaki disease: a prospective population survey in the UK and Ireland from 2013 to 2015. Archives of disease in childhood

Ulloa-Gutierrez, R., Salgado, A. P., Garrido-Garcia, L. M. et al. (2016) Kawasaki disease (KD) in infants <6 months of age among 20 latin american (LA) countries: a prospective multinational multicenter study of the rekamlatina network. European journal of pediatrics. Conference: 6th congress of the european academy of paediatric societies. Switzerland. Conference start: 20161021. Conference end: 20161025 175(11): 1778

Wilder, Matthew S., Palinkas, Lawrence A., Kao, Annie S. et al. (2007) Delayed diagnosis by physicians contributes to the development of coronary artery aneurysms in children with Kawasaki syndrome. The Pediatric infectious disease journal 26(3): 256-60
Yamashita, Maho, Ae, Ryusuke, Yashiro, Mayumi et al. (2017) Difference in Risk Factors for Subtypes of Acute Cardiac Lesions Resulting from Kawasaki Disease. Pediatric cardiology 38(2): 375-380
Yamazaki-Nakashimada, Marco Antonio, Deguchi, Kuntaro, Gamez-Gonzalez, Berenise et al. (2019) Orange-brown chromonychia: A valid sign in Kawasaki disease in children of different ethnicities. International journal of rheumatic diseases
Yeo, Yunku, Kim, TaeYeon, Ha, KeeSoo et al. (2009) Incomplete Kawasaki disease in patients younger than 1 year of age: a possible inherent risk factor. European journal of pediatrics 168(2): 157-62

## Code [Reason]

- This is an epidemiological study about incidence rates. Signs and symptoms were beyond the scope of the study.
- This is a study about imaging rather than signs and symptoms
- Study happened outside Europe and had fewer than 100 patients
- Study happened outside Europe and had fewer than 100 patients
- No data on the signs and symptoms.
- Conference abstract.
- No data on the signs and symptoms.
- No data on the signs and symptoms.
- This is a case study.
- No data on the signs and symptoms.


## Appendix J - Research recommendations

|  | Which signs and symptoms (or combinations of signs and <br> symptoms) predict a diagnosis of Kawasaki disease in children under <br> 5 presenting with fever lasting 5 days or more? |
| :--- | :--- |
| Question | Children aged 5 years or under presenting with fever lasting 5 days or <br> longer. <br> Subgroup: Children aged under 1 |
| Population | Signs and symptoms of Kawasaki disease including: <br> - bilateral conjunctival injection without exudate |
| - erythema and cracking of lips; strawberry tongue; or erythema of oral and |  |
| pharyngeal mucosa |  |

# Appendix K - Reference list of included studies 

Advani, Najib; Santoso, Lucyana Alim; Sastroasmoro, Sudigdo (2019) Profile of Kawasaki Disease in Adolescents: Is It Different? Acta medica Indonesiana 51(1): 42-46

Bai, L., Feng, T., Yang, L. et al. (2017) Retrospective analysis of risk factors associated with Kawasaki disease in China. Oncotarget 8(33): 54357-54363

Baker, Annette L., Lu, Minmin, Minich, L. LuAnn et al. (2009) Associated symptoms in the ten days before diagnosis of Kawasaki disease. The Journal of pediatrics 154(4): 592-595.e2

Bal, Aswine K., Prasad, Deepa, Umali Pamintuan, Maria Angela et al. (2014) Timing of intravenous immunoglobulin treatment and risk of coronary artery abnormalities in children with Kawasaki disease. Pediatrics and neonatology 55(5): 387-92

Behmadi, Maryam; Alizadeh, Behzad; Malek, Abdolreza (2019) Comparison of Clinical Symptoms and Cardiac Lesions in Children with Typical and Atypical Kawasaki Disease. Medical sciences (Basel, Switzerland) 7(4)

Boudiaf, Houda and Achir, Moussa (2016) The Clinical Profile of Kawasaki Disease in Algerian Children: A Single Institution Experience. Journal of tropical pediatrics 62(2): 139-43

Chang, Luan-Yin, Lu, Chun-Yi, Shao, Pei-Lan et al. (2014) Viral infections associated with Kawasaki disease. Journal of the Formosan Medical Association = Taiwan yi zhi 113(3): 148-54

Chen, J. J., Ma, X. J., Liu, F. et al. (2016) Epidemiologic features of Kawasaki disease in Shanghai from 2008 Through 2012. Pediatric Infectious Disease Journal 35(1): 7-12

Ebbeson, Regan L., Riley, Mark R., Potts, Jim E. et al. (2004) Kawasaki disease at British Columbia's Children's Hospital. Paediatrics \& child health 9(7): 466-70

Fabi, Marianna, Corinaldesi, Elena, Pierantoni, Luca et al. (2018) Gastrointestinal presentation of Kawasaki disease: A red flag for severe disease? PloS one 13(9): e0202658

Falcini, F., Calabri, G. B., Ricci, L. et al. (2007) Update on Kawasaki disease: The 25-year experience at the "A. Mayer" Children's Hospital, Florence. Italian Journal of Pediatrics 33(1): 32-40

Falcini, Fernanda, Ozen, Seza, Magni-Manzoni, Silvia et al. (2012) Discrimination between incomplete and atypical Kawasaki syndrome versus other febrile diseases in childhood: results from an international registry-based study. Clinical and experimental rheumatology 30(5): 799-804

Gamez-Gonzalez, Luisa Berenise, Murata, Chiharu, Munoz-Ramirez, Mireya et al. (2013) Clinical manifestations associated with Kawasaki disease shock syndrome in Mexican children. European journal of pediatrics 172(3): 337-42

Generini, S., Ermini, M., Taccetti, G. et al. (1997) Clinical and laboratory features and disease outcome of kawasaki disease: the analysis of our experience and literature review. Journal of clinical rheumatology: practical reports on rheumatic \& musculoskeletal diseases 3(5): 241-7

Garrido-Garcia, Luis Martin, Castillo-Moguel, Ariel, Vazquez-Rivera, Mirella et al. (2017) Reaction of the BCG Scar in the Acute Phase of Kawasaki Disease in Mexican Children. The Pediatric infectious disease journal 36(10): e237-e241

Ghelani, Sunil J., Sable, Craig, Wiedermann, Bernhard L. et al. (2012) Increased incidence of incomplete Kawasaki disease at a pediatric hospital after publication of the 2004 American Heart Association guidelines. Pediatric cardiology 33(7): 1097-103

Giannouli, Georgia, Tzoumaka-Bakoula, Chryssa, Kopsidas, loannis et al. (2013) Epidemiology and risk factors for coronary artery abnormalities in children with complete and incomplete Kawasaki disease during a 10-year period. Pediatric cardiology 34(6): 1476-81

Gorrab, Arbia Abir, Fournier, Anne, Bouaziz, Asma Abed et al. (2016) Incidence Rate and Epidemiological and Clinical Aspects of Kawasaki Disease in Children of Maghrebi Origin in the Province of Quebec, Canada, Compared to the Country of Origin. Global pediatric health 3: 2333794x16630670

Hu, Ya-Chiao, Liu, Hsin-Min, Lin, Ming-Tai et al. (2019) Outcomes of Kawasaki Disease Children with Spontaneous Defervescence Within 10 Days. Frontiers in pediatrics 7: 158

Huang GY, Ma XJ, Huang M, et al. Epidemiologic pictures of Kawasaki disease in Shanghai from 1998 through 2002. Journal of Epidemiology 2006;16(1):9-14.

Jaggi, Preeti, Grcic, Michelle, Kovalchin, John et al. (2018) Using the Electronic Medical Record to Correlate Kawasaki Disease Phenotypes with Clinical Outcomes. Journal of the Pediatric Infectious Diseases Society 7(2): 119-123

Jun, Hyun Ok, Yu, Jeong Jin, Kang, So Yeon et al. (2015) Diagnostic characteristics of supplemental laboratory criteria for incomplete Kawasaki disease in children with complete Kawasaki disease. Korean journal of pediatrics 58(10): 369-73

Jun, Woo Young, Ann, Yu Kyung, Kim, Ja Yeong et al. (2017) Kawasaki Disease with Fever and Cervical Lymphadenopathy as the Sole Initial Presentation. Korean circulation journal 47(1): 107114

Kil, Hong-Ryang, Yu, Jae-Won, Lee, Sung-Churl et al. (2017) Changes in clinical and laboratory features of Kawasaki disease noted over time in Daejeon, Korea. Pediatric rheumatology online journal 15(1): 60

Kim, Gi Beom, Park, Sohee, Eun, Lucy Youngmin et al. (2017) Epidemiology and Clinical Features of Kawasaki Disease in South Korea, 2012-2014. The Pediatric infectious disease journal 36(5): 482-485

Kim, S. H.; Lee, H. J.; Lee, J. S. (2018) Clinical aspects of periungual desquamation in Kawasaki disease. Iranian Journal of Pediatrics 28(3): e59262

Kim, Seong Hyun; Kim, Ki Hwan; Kim, Dong Soo (2009) Clinical characteristics of Kawasaki disease according to age at diagnosis. Indian pediatrics 46(7): 585-90

Kubota, Masaru, Usami, Ikuya, Yamakawa, Masaru et al. (2008) Kawasaki disease with lymphadenopathy and fever as sole initial manifestations. Journal of paediatrics and child health 44(6): 359-62

Li, Wei, Zhang, Li, Huang, Ping et al. (2019) Clinical features and mid-term follow-up in infants younger than 3 months with Kawasaki disease in a Chinese population. Journal of paediatrics and child health 55(5): 523-527

Liu, Hao-Chuan, Lo, Chiao-Wei, Hwang, Betau et al. (2012) Clinical manifestations vary with different age spectrums in infants with Kawasaki disease. TheScientificWorldJournal 2012: 210382

Manlhiot, Cedric, Christie, Erin, McCrindle, Brian W. et al. (2012) Complete and incomplete Kawasaki disease: two sides of the same coin. European journal of pediatrics 171(4): 657-62

Maric, Lorna Stemberger, Knezovic, Ivica, Papic, Neven et al. (2015) Risk factors for coronary artery abnormalities in children with Kawasaki disease: a 10-year experience. Rheumatology international 35(6): 1053-8

Martins, Andreia, Conde, Marta, Brito, Maria et al. (2018) Arthritis in Kawasaki disease: A poorly recognised manifestation. Journal of paediatrics and child health 54(12): 1371-1374

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[^0]:    Fever under 5 s : evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

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[^13]:    Fever under 5 s : evidence reviews for signs and symptoms predicting Kawasaki disease
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[^14]:    Joanna Briggs critical appraisal checklist for case series
    Were there clear criteria for inclusion in the case series?
    Yes

[^15]:    Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

[^16]:    Fever under 5s: evidence reviews for signs and symptoms predicting Kawasaki disease FINAL (November 2019)

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