

































<b>Recommendations</b>	-
<b>Research recommendations</b>	<p><b>RR10. Is a minor injury unit, urgent care or walk-in centre clinically and cost effective i) as a standalone unit and ii) when located on the same site as an emergency department?</b></p>
	<p>emergency service (for example, ophthalmology or dental) with designated accommodation for the reception of patients.</p> <ul style="list-style-type: none"> <li>• <b>Type 3 A&amp;E department/Type 4 A&amp;E department/Urgent Care Centre:</b> Other type of A&amp;E/minor injury units (MIUs)/Walk-in Centres (WiCs)/Urgent Care Centre primarily designed for the receiving of accident and emergency patients.</li> </ul> <p>The committee considered that greater access to GPs (evening and Saturday GP appointments) would potentially reduce the need for walk-in centres in particular. It was noted that there is a tendency among rural populations to make use of GPs to a greater extent than urban populations, which may impact on ED demand. Future research on the impact of MIUs/UCCs/WiCs on ED demand should include measurement of case mix. Potential changes in case mix as a result of service reconfiguration could have significant economic implications. The shift of patients with minor conditions from EDs to these units, although reducing the pressure on the EDs, could be associated with an artefactual increase in time in ED (mean and variance) or admission rate due to the higher acuity of the residual ED case-mix.</p> <p>It was agreed that the absence of a consistent terminology for structures and processes and also the level and type of staffing relating to MIUs, UCCs and WiCs makes their evaluation challenging. Future studies should take into account several contextual factors including location (inner-city, urban and rural), opening times (24 hour versus restricted times) staffing composition and expertise, available resources, processes and overall service configuration in their analyses. Proximity to these units could be used as an instrumental variable to evaluate outcomes given the impossibility of randomising populations.</p> <p>The majority of the evidence came from studies with relatively short follow up periods, which the committee considered to be a significant limitation as it may not reflect long term effects. Future studies evaluating effects over longer time frames would offer the opportunity to account for secular trends and detect population effects.</p> <p>As well as ED demand, other outcomes should be patient-focussed and rooted in health economics evaluation. The potential impact on other services such as the ambulance service (particularly within rural areas) should be evaluated. Staff exposure to specific health problems within the ED may be reduced as a result of streaming particular groups of patients to specialist centres. Therefore, it would also be useful to assess the impact on staff training and potential staff de-skilling. In summary, the current level of evidence is insufficient to permit a recommendation on the internal or external configuration of such units. Opportunities should be taken to evaluate MIUs, UCCs and WICs using existing services or if local health economies choose to implement such services.</p>



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## Appendices

### Appendix A: Review protocol

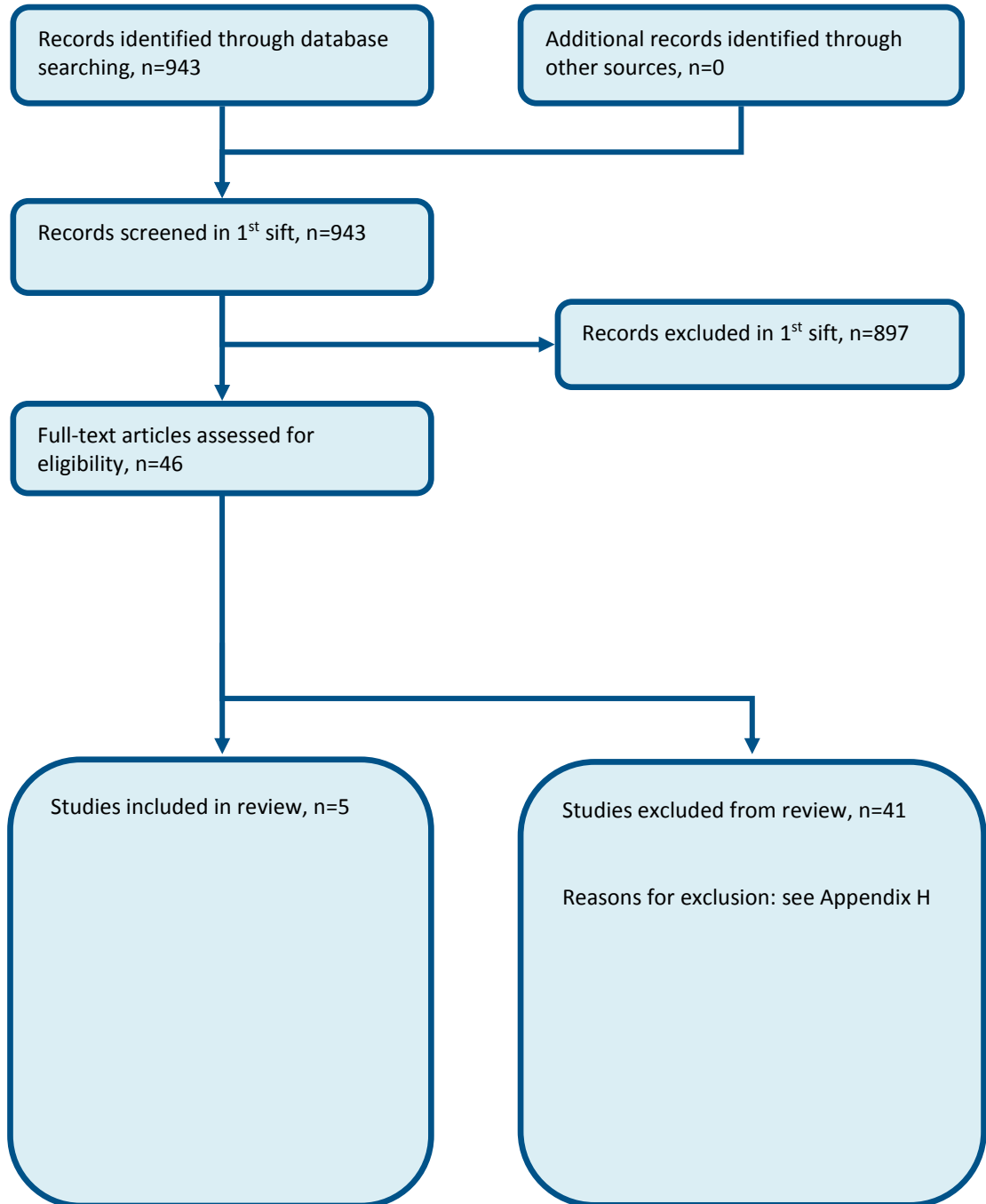
**Table 6: Review protocol: Minor injury unit, urgent care centre or walk-in centre**

Review question	Minor injury unit, urgent care centre or walk-in centre
Guideline condition and its definition	Acute medical emergencies.
Objectives	Is a minor injury unit, urgent care centre or walk-in centre clinically and cost effective: 1. as a stand-alone unit 2. when located on the same site as a full emergency department?
Review population	Adults and young people (16 years and over) presenting with a suspected or confirmed AME.
	Adults and young people (16 years and over).
	Line of therapy not an inclusion criterion.
Interventions and comparators: generic/class; specific/drug  (All interventions will be compared with each other, unless otherwise stated)	Presence of minor injury units, urgent care centres or walk in centres; as stand-alone units. Presence of minor injury units, urgent care centres or walk in centres; within a full emergency department. Absence of minor injury units, urgent care centres or walk in centres; absence.
Outcomes	<ul style="list-style-type: none"> <li>- Avoidable adverse events (Dichotomous) CRITICAL</li> <li>- Quality of life (Continuous) CRITICAL</li> <li>- Patient and/or carer satisfaction (Continuous) CRITICAL</li> <li>- Waiting time in ED (including A&amp;E 4 hour waiting target breach) (Continuous) CRITICAL</li> <li>- Mortality (Dichotomous) CRITICAL</li> <li>- ED Avoidance (Dichotomous) IMPORTANT</li> </ul>
Study design	<ul style="list-style-type: none"> <li>Systematic Review</li> <li>RCT</li> <li>Quasi-RCT</li> <li>Non randomised study</li> <li>Prospective cohort study</li> <li>Retrospective cohort study</li> <li>Controlled before and after study</li> <li>Before and after study</li> </ul>
Unit of randomisation	<ul style="list-style-type: none"> <li>Patient.</li> <li>Hospital.</li> <li>Ward.</li> </ul>
Crossover study	Not permitted.
Minimum duration of study	Not defined.
Subgroup analyses if there is heterogeneity	<ul style="list-style-type: none"> <li>- Case mix (frail elderly; not frail elderly); effects may be different in this subgroup.</li> <li>- Skill mix (doctor present; nurse led); effects may be different in this subgroup</li> <li>- Facilities (access to radiology; access to pathology); effects may be different in</li> </ul>

Review question	Minor injury unit, urgent care centre or walk-in centre
	<p>this subgroup.</p> <ul style="list-style-type: none"> <li>- Opening hours (24 hours a day; less than 24 hours a day); effects may be different in this subgroup.</li> <li>- Location (rural; urban); effects may be different in this subgroup.</li> </ul>
Exclusions	UK only.
Search criteria	<p>Databases: Medline, Embase, the Cochrane Library.            Date limits for search: 1995.            Language: English only.</p>

## Appendix B: Clinical article selection

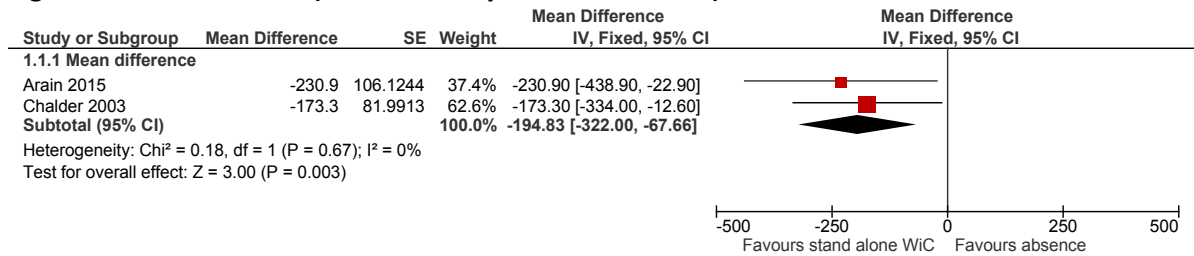
Figure 1: Flow chart of clinical article selection for the review of minor injury unit, urgent care centre or walk-in centre



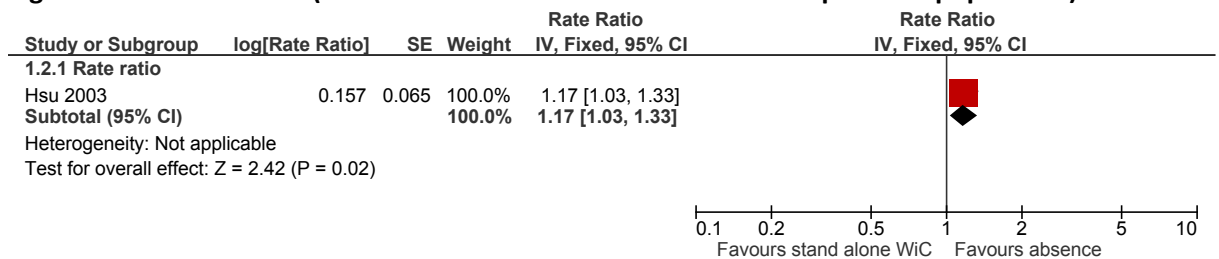
# Appendix C: Forest plots

## C.1 Stand-alone units versus absence

**Figure 2: ED avoidance (mean monthly attendance rates)**

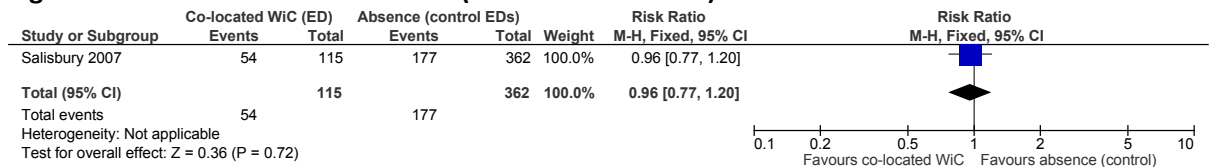


**Figure 3: ED avoidance (annual non-ambulance attendance rates per 1000 population)**



## C.2 Co-located on the same site as the ED versus absence

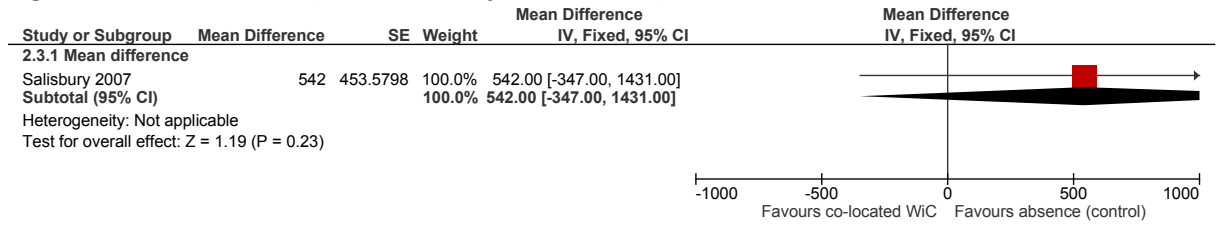
**Figure 4: Avoidable adverse events (ED re-consultations)**



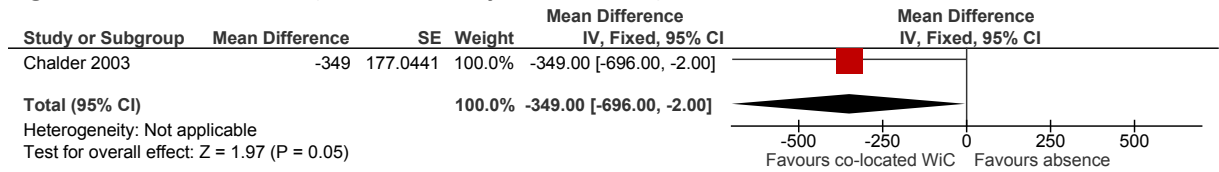
**Figure 5: Avoidable adverse events (ED + WiC re-consultations)**



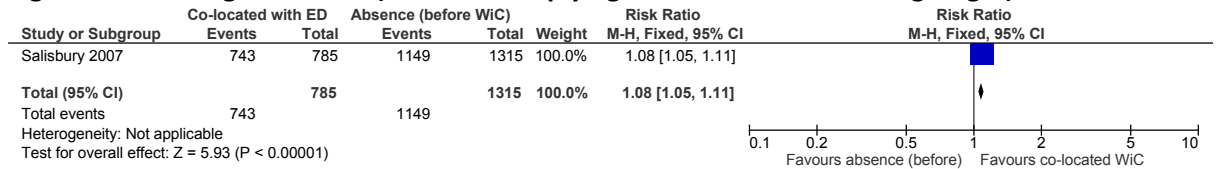
**Figure 6: ED avoidance (mean monthly attendances)**



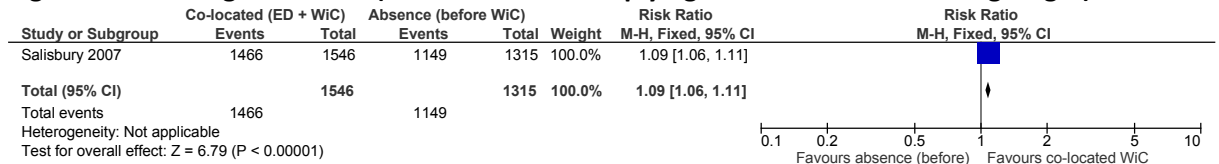
**Figure 7: ED avoidance (mean monthly attendances)**



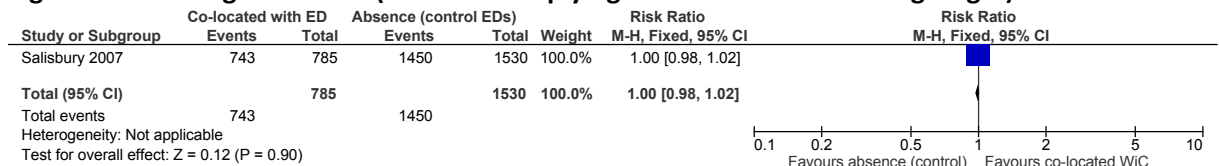
**Figure 8: Waiting time in ED (ED cases complying with A&E 4 hour waiting target)**



**Figure 9: Waiting time in ED (ED + WiC cases complying with A&E 4 hour waiting target)**

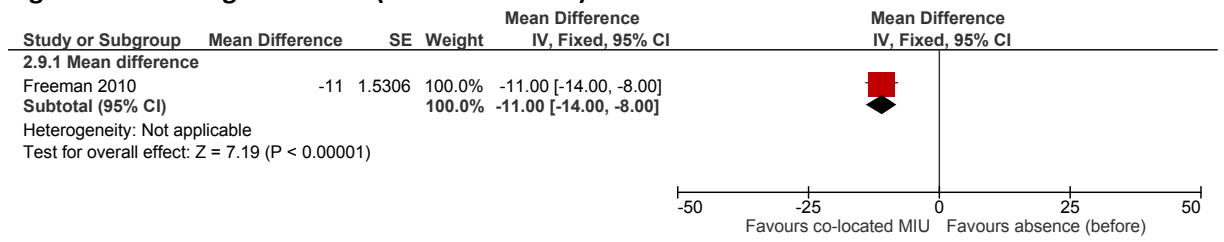


**Figure 10: Waiting time in ED (ED cases complying with A&E 4 hour waiting target)**

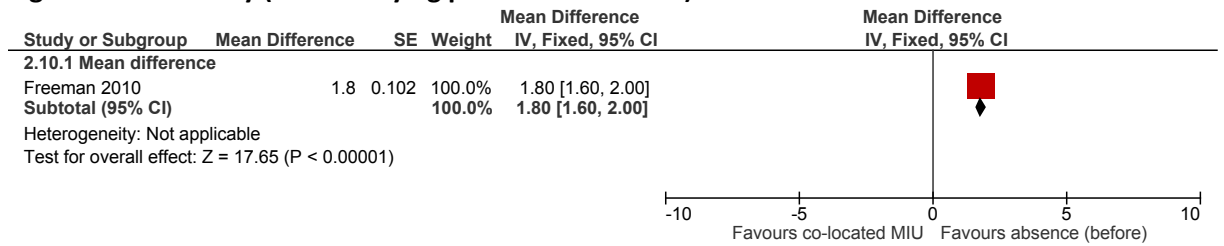




**Figure 11: Waiting time in ED (time to clinician)**



**Figure 12: Mortality (number dying per 1000 attendees)**



## Appendix D: Clinical evidence tables

Study	Arain 2015 trial: Arain 2015 <sup>1</sup>
Study type	Before and after study.
Number of studies (number of participants)	Not reported.
Countries and setting	Conducted in United Kingdom; setting: emergency department and minor injuries unit.
Line of therapy	Not applicable.
Duration of study	Other: 1 year before and 1 year after the opening of a GP led WiC.
Method of assessment of guideline condition	Partially adequate method of assessment/diagnosis: patients attending the ED.
Stratum	Overall.
Subgroup analysis within study	Not applicable.
Inclusion criteria	Patients attending the ED.
Exclusion criteria	None stated.
Recruitment/selection of patients	Consecutive patients.
Age, gender and ethnicity	Age: not reported. Gender (M:F): not reported. Ethnicity: not reported.
Further population details	1. Case mix: Not applicable/Not stated/Unclear.
Indirectness of population	No indirectness.
Interventions	<p>Intervention 1: Presence of minor injury unit, urgent care centre or walk in centre - as stand-alone units. GP-led walk-in centre. Duration: 1 year after the opening of a GP-led walk-in centre. Concurrent medication/care: not applicable. Further details: 1. Facilities: Not applicable/Not stated/Unclear 2. Location: urban 3. Opening hours: less than 24 hours a day 4. Skill mix: doctor present.</p> <p>Intervention 2: Absence of minor injury unit, urgent care centre or walk in centre - absence. Emergency department with absence of stand-alone walk-in centre. Duration: 1 year before the opening of a GP-led walk-in centre. Concurrent medication/care: not applicable. Further details: 1. Facilities: Not applicable/Not stated/Unclear 2. Location: urban. 3. Opening hours: 24 hours a day. 4. Skill mix: doctor present.</p>
Funding	Funding not stated.

Study	Arain 2015 trial: Arain 2015 <sup>1</sup>
RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: AS STAND ALONE UNITS versus ABSENCE.	
Protocol outcome 1: ED avoidance. - Actual outcome: Mean monthly ED attendances at 1 year before and 1 year after the opening of a GP-led walk-in centre; Mean: -230.9 (95% CI -438.9 to -21.9); Risk of bias: All domain - High, Selection - Very high, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness	
Protocol outcomes not reported by the study	Avoidable adverse events; Quality of life; Patient and/or carer satisfaction; Waiting time in ED (including A&E 4 hour waiting target breach); Mortality.

Study	Chalder 2003 trial: Chalder 2003 <sup>10</sup>
Study type	Controlled before and after study.
Number of studies (number of participants)	Not reported.
Countries and setting	Conducted in United Kingdom; setting: 20 A&E departments.
Line of therapy	Not applicable.
Duration of study	Other: 1 year before and 1 year after the opening of a local walk-in centre, or the opening of a walk-in centre in the matched site for control sites.
Method of assessment of guideline condition	Adequate method of assessment/diagnosis.
Stratum	Overall.
Subgroup analysis within study	Not applicable.
Inclusion criteria	Not reported.
Exclusion criteria	Not reported.
Recruitment/selection of patients	A&E department closest to the walk-in centre or town centre for control sites.
Age, gender and ethnicity	Age - Other: Age not reported. Gender (M:F): Not reported. Ethnicity: Not reported.
Further population details	1. Case mix: Not applicable/Not stated/Unclear.
Indirectness of population	No indirectness.
Interventions	(n=10) Intervention 1: Presence of minor injury unit, urgent care centre or walk in centre - as stand-alone units. EDs after the opening of local walk-in centres. Duration: 1 year after walk-in centres opened. Concurrent medication/care: Not applicable.

<b>Study</b>	<b>Chalder 2003 trial: Chalder 2003<sup>10</sup></b>
	Further details: 1. Facilities: Not applicable/Not stated/Unclear. 2. Location: Not applicable/Not stated/Unclear. 3. Opening hours: Not applicable/Not stated/Unclear. 4. Skill mix: Not applicable/Not stated/Unclear.  (n=10) Intervention 2: Absence of minor injury unit, urgent care centre or walk in centre - absence. EDs before the opening of local walk-in centres. Duration: 1 year before walk-in centres opened. Concurrent medication/care: not applicable. Further details: 1. Facilities: Not applicable/Not stated/Unclear. 2. Location: Not applicable/Not stated/Unclear. 3. Opening Hours: Not applicable/Not stated/Unclear. 4. Skill mix: Not applicable/Not stated/Unclear.
Funding	Academic or government funding (Department of Health).
<b>RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: AS STAND ALONE UNITS versus ABSENCE.</b>	
Protocol outcome 1: ED avoidance. - Actual outcome: Mean number of monthly consultations at 1 year (all walk-in centres); Mean -173 (95%CI -334 to -12); Risk of bias: All domain - Low, Selection - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness ; Baseline details: control sites were of a similar size and in the same region Actual outcome: Mean number of monthly consultations at 1 year (co-located walk-in centres only): Mean -349 (95%CI -696 to -2); Risk of bias: All domain - High, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness ; Baseline details: control sites were of a similar size and in the same region	
Protocol outcomes not reported by the study	Avoidable adverse events; Quality of life; Patient and/or carer satisfaction; Waiting time in ED (including A&E 4 hour waiting target breach); Mortality.

<b>Study</b>	<b>Freeman 2010 trial: Freeman 2010<sup>15</sup></b>
Study type	Before and after study.
Number of studies (number of participants)	(n=584,321).
Countries and setting	Conducted in United Kingdom; setting: A&E department.
Line of therapy	Not applicable.
Duration of study	Other: 2 years before and 4 years after the opening of a co-located minor injuries unit.
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: quality checks on hospital database accuracy undertaken on a weekly basis.
Stratum	Overall.
Subgroup analysis within study	Not applicable.

Study	Freeman 2010 trial: Freeman 2010 <sup>15</sup>
Inclusion criteria	Patients attending the ED.
Exclusion criteria	Patients attending for planned follow-up appointments; patients who were dead on arrival; aged <14 years.
Recruitment/selection of patients	Consecutive patients.
Age, gender and ethnicity	Age - Median: 40. Gender (M:F): 54/46. Ethnicity: not reported.
Further population details	1. Case mix: Not applicable/Not stated/Unclear.
Indirectness of population	No indirectness.
Interventions	<p>(n=333,832) Intervention 1: Presence of minor injury unit, urgent care centre or walk in centre - within a full emergency department. Co-located minor injuries unit. Duration: 4 years after the opening of a co-located MIU. Concurrent medication/care: not applicable. Further details: 1. Facilities: Not applicable/Not stated/Unclear. 2. Location: Urban 3. Opening Hours: Not applicable/Not stated/Unclear. 4. Skill mix: Not applicable/Not stated/Unclear.</p> <p>(n=250,489) Intervention 2: Absence of minor injury unit, urgent care centre or walk in centre - absence. ED with no co-located minor injuries unit. Duration: 2 years before the opening of a co-located MIU. Concurrent medication/care: not applicable. Further details: 1. Facilities: Not applicable/Not stated/Unclear. 2. Location: urban. 3. Opening hours: Not applicable/Not stated/Unclear. 4. Skill mix: Not applicable/Not stated/Unclear.</p>
Funding	Funding not stated.
RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: WITHIN A FULL EMERGENCY CARE DEPARTMENT versus ABSENCE.	
<p>Protocol outcome 1: Waiting time in ED (including A&amp;E 4 hour waiting target breach). - Actual outcome: Time (minutes) to clinician at visit; Other: -11 (95%CI -14 to -8); Risk of bias: All domain - Low, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness</p> <p>Protocol outcome 2: Mortality. - Actual outcome: Number dying per 1000 attendees at visit; Other: 1.8 (95%CI 1.6 to 1.9); Risk of bias: All domain - Low, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness</p>	
Protocol outcomes not reported by the study	Avoidable adverse events; Quality of life; Patient and/or carer satisfaction; ED Avoidance.
Study	Hsu 2003 trial: Hsu 2003 <sup>21</sup>

Study	Hsu 2003 trial: Hsu 2003 <sup>21</sup>
Study type	Controlled before and after study.
Number of studies (number of participants)	Not reported.
Countries and setting	Conducted in United Kingdom; setting: EDs with and without stand-alone walk-in centres in the same town.
Line of therapy	Not applicable.
Duration of study	Other: 6 months before and 1 year after the opening of a stand -alone walk-in centre.
Method of assessment of guideline condition	Method of assessment/diagnosis not stated.
Stratum	Overall.
Subgroup analysis within study	Not applicable.
Inclusion criteria	Not reported.
Exclusion criteria	Not reported.
Recruitment/selection of patients	Consecutive patients.
Age, gender and ethnicity	Age - Other: not reported. Gender (M:F): not reported. Ethnicity: not reported.
Further population details	1. Case mix: Not applicable/Not stated/Unclear.
Indirectness of population	No indirectness.
Interventions	<p>(n=3) Intervention 1: Presence of minor injury units, urgent care centre or walk in centre - as stand-alone units. EDs after the opening of local stand-alone walk-in centres. Duration: 6 months after the opening of walk-in centres. Concurrent medication/care: not applicable. Further details: 1. Facilities: Not applicable/Not stated/Unclear. 2. Location: Not applicable/Not stated/Unclear. 3. Opening hours: Not applicable/Not stated/Unclear. 4. Skill mix: Not applicable/Not stated/Unclear.</p> <p>(n=3) Intervention 2: Absence of minor injury unit, urgent care centre or walk in centre - absence. EDs before the opening of local stand-alone walk-in centres. Duration: 6 months before the opening of walk-in centres. Concurrent medication/care: not applicable. Further details: 1. Facilities: Not applicable/Not stated/Unclear. 2. Location: Not applicable/Not stated/Unclear. 3. Opening hours: Not applicable/Not stated/Unclear. 4. Skill mix: Not applicable/Not stated/Unclear.</p>
Funding	Academic or government funding (Department of Health).
RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: AS STAND ALONE UNITS versus ABSENCE.	
Protocol outcome 1: ED avoidance.	

<b>Study</b>	<b>Hsu 2003 trial: Hsu 2003<sup>21</sup></b>
	- Actual outcome: Annual rate of non-ambulance attendances (per 1000 population) at 6 months; Other: 1.17 (95%CI 1.03 to 1.33); Comments: Adjusted rate ratio Risk of bias: All domain - Low, Selection - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness
Protocol outcomes not reported by the study	Avoidable adverse events; Quality of life; Patient and/or carer satisfaction; Waiting time in ED (including A&E 4 hour waiting target breach); Mortality.

<b>Study</b>	<b>Salisbury 2007 trial: Salisbury 2007<sup>35</sup></b>
Study type	Controlled before and after study.
Number of studies (number of participants)	Not reported.
Countries and setting	Conducted in United Kingdom; setting: EDs with or without co-located walk-in centres.
Line of therapy	Not applicable.
Duration of study	Other: 9 months before and 6 months after the opening of co-located walk-in centres.
Method of assessment of guideline condition	Adequate method of assessment/diagnosis: patient records.
Stratum	Overall.
Subgroup analysis within study	Not applicable.
Inclusion criteria	Patients consulting at EDs with and without co-located walk-in centres.
Exclusion criteria	Patients under 16 years of age.
Recruitment/selection of patients	Randomly selected from those consulting in a 2-week period at least 3 months after the walk-in centres opened and the same period 1 year earlier.
Age, gender and ethnicity	Age - Other: age not reported. Gender (M:F): not reported. Ethnicity: not reported.
Further population details	1. Case mix: Not applicable/Not stated/Unclear.
Indirectness of population	No indirectness.
Interventions	(n=785) Intervention 1: Presence of minor injury units, urgent care centre or walk in centre - within a full emergency department. EDs after the opening of co-located walk-in centres. Duration: 6 months after the opening of co-located walk-in centres. Concurrent medication/care: not applicable. Further details: 1. Facilities: Not applicable/Not stated/Unclear. 2. Location: Not applicable/Not stated/Unclear. 3. Opening hours: Not applicable/Not stated/Unclear. 4. Skill mix: doctor present.

Study	Salisbury 2007 trial: Salisbury 2007 <sup>35</sup>
	<p>(n=1,315) Intervention 2: Absence of minor injury unit, urgent care centre or walk in centre - absence. EDs before the opening of co-located walk-in centres. Duration: 9 months before the opening of co-located walk in centres. Concurrent medication/care: not applicable. Further details: 1. Facilities: Not applicable/Not stated/Unclear. 2. Location: Not applicable/Not stated/Unclear. 3. Opening hours: Not applicable/Not stated/Unclear. 4. Skill mix: doctor present.</p> <p>(n=1,546) Intervention 3: Presence of minor injury units, urgent care centre or walk in centre - within a full emergency department. EDs and walk-in centres after the opening of co-located walk-in centres. Duration: 6 months after the opening of co-located walk-in centres. Concurrent medication/care: not applicable. Further details: 1. Facilities: Not applicable/Not stated/Unclear. 2. Location: Not applicable/Not stated/Unclear. 3. Opening hours: Not applicable/Not stated/Unclear. 4. Skill mix: doctor present.</p> <p>(n=1,530) Intervention 4: Absence of minor injury unit, urgent care centre or walk in centre - absence. Control EDs after the opening of co-located walk-in centres at intervention EDs. Duration: 6 months after the opening of co-located walk-in centres at intervention EDs. Concurrent medication/care: not applicable. Further details: 1. Facilities: Not applicable/Not stated/Unclear. 2. Location: Not applicable/Not stated/Unclear. 3. Opening hours: Not applicable/Not stated/Unclear. 4. Skill mix: Not applicable/Not stated/Unclear.</p>
Funding	Academic or government funding (Department of Health).
<p><b>RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: WITHIN A FULL EMERGENCY CARE DEPARTMENT versus ABSENCE.</b></p> <p>Protocol outcome 1: Waiting time in ED (including A&amp;E 4 hour waiting target breach). - Actual outcome: Mean visit duration at visit; Risk of bias: All domain - Low, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness - Actual outcome: Cases complying with A&amp;E 4 hour waiting target at visit; Group 1: 743/785, Group 2: 1149/1315; Risk of bias: All domain - Low, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness</p> <p><b>RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: WITHIN A FULL EMERGENCY CARE DEPARTMENT versus ABSENCE (CONTROL EDS).</b></p> <p>Protocol outcome 1: Avoidable adverse events. - Actual outcome: Re-consultations about the same problem at 4 weeks after visit; Group 1: 54/115, Group 2: 177/362; Risk of bias: All domain - High, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness</p> <p>Protocol outcome 2: Waiting time in ED (including A&amp;E 4 hour waiting target breach). - Actual outcome: Mean visit duration at visit; Risk of bias: Low; Indirectness of outcome: No indirectness. - Actual outcome: Cases complying with A&amp;E 4 hour waiting target at visit; Group 1: 743/785, Group 2: 1450/1530; Risk of bias: Low; Indirectness of outcome: No indirectness.</p>	



Study	Salisbury 2007 trial: Salisbury 2007 <sup>35</sup>
<p>RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: WITHIN A FULL EMERGENCY CARE DEPARTMENT (ED+WIC) versus ABSENCE.</p> <p>Protocol outcome 1: Waiting time in ED (including A&amp;E 4 hour waiting target breach).            - Actual outcome: Mean visit duration at visit; Risk of bias: All domain - Low, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness - Actual outcome: Cases complying with A&amp;E 4 hour waiting target at visit; Group 1: 1466/1546, Group 2: 1149/1315; Risk of bias: All domain - Low, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - Low, Crossover - Low; Indirectness of outcome: No indirectness</p> <p>RESULTS (NUMBERS ANALYSED) AND RISK OF BIAS FOR COMPARISON: WITHIN A FULL EMERGENCY CARE DEPARTMENT (ED+WIC) versus ABSENCE (CONTROL EDS).</p> <p>Protocol outcome 1: Avoidable adverse events.            - Actual outcome: Re-consultations about the same problem at 4 weeks after visit; Group 1: 149/330, Group 2: 177/362; Risk of bias: All domain - High, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness</p> <p>Protocol outcome 2: ED Avoidance.            - Actual outcome: Attendances at per month; Other: 542 (95%CI -347 to 1431); Risk of bias: All domain - High, Selection - High, Blinding - High, Incomplete outcome data - Low, Outcome reporting - Low, Measurement - High, Crossover - Low; Indirectness of outcome: No indirectness</p>	
Protocol outcomes not reported by the study	Quality of life; Patient and/or carer satisfaction; Mortality.

## Appendix E: Economic evidence tables

Study	Salisbury 2007 <sup>35</sup>			
Study details	Population & interventions	Costs	Health outcomes	Cost effectiveness
<p><b>Economic analysis:</b> CC</p> <p><b>Study design:</b> controlled before and after study</p> <p><b>Approach to analysis:</b> Regression analysis was used to compare the resource use, costs and outcomes data, adjusting for clustering. The data were obtained from anonymised patient records and by means of a patient survey. Sites were matched based on performance on the 4-hour target, proportion admitted and size of ED.</p> <p><b>Perspective:</b> UK NHS</p> <p><b>Follow-up:</b> 6 months ( 3 months before and 3 months after)</p> <p><b>Treatment effect duration<sup>(a)</sup>:</b> same as follow-up</p> <p><b>Discounting:</b> Costs: NA; Outcomes: NA</p>	<p><b>Population:</b> Patients consulting at EDs/co-located walk-in centres</p> <p><b>Cohort settings:</b> Start age: NR Male: NR</p> <p><b>Intervention 1:</b> EDs in matched control sites with no walk-in centre before and after the opening of walk-in centres in intervention sites</p> <p><b>Intervention 2:</b> EDs before and after the opening of standalone walk-in centres in the same town</p>	<p><b>Total costs, before (mean per patient):</b> Intervention 1: £113.39 Intervention 2: £110.96 Incremental (2-1): -£2.94 (95% CI: NR; p=NR)</p> <p><b>Total costs, after (mean per patient):</b> Intervention 1: £121.67 Intervention 2: £117.18 Incremental (2-1): -£4.49 (95% CI: NR; p=NR)</p> <p><b>Change in total cost (after-before) (mean per patient):</b> Intervention 1: £8.28 Intervention 2: £6.22 Incremental (2-1): -£3.06 (95% CI: -£16.50 to £10.39; p=NR)</p> <p><b>Currency &amp; cost year:</b> 2004-2005 UK pounds</p> <p><b>Cost components incorporated:</b> Variable costs relating to the following resources: Clinical staff time, administrative and clerical staff time, investigations, treatments, medications, admissions (included only in sensitivity analysis, onward referrals, and re-consultations.</p>	N/A	<p><b>ICER (Intervention 2 versus Intervention 1):</b> N/A</p> <p><b>Analysis of uncertainty:</b> Sensitivity analysis including admission costs was reported and showed no significant difference in the change in cost per patient (-£20.97; 95% CI: -£64.98 to £23.04)</p>

<b>Data sources</b>
<b>Health outcomes:</b> before and after data collected via survey and patient records on ED visit duration, re-consultation rates and patient satisfaction. Regression analysis adjusting for clustering was used. <b>Quality-of-life weights:</b> N/A. <b>Cost sources:</b> hospital records.
<b>Comments</b>
<b>Source of funding:</b> Department of Health. <b>Applicability and limitations:</b> Some uncertainty regarding the applicability of resource use and costs from 2004-2005 to current NHS context. The study is a comparative cost analysis; hence, QALYs were not used as an outcome measure. The study has a short follow-up period (3 months before and 3 months after), so follow-up may not have been long enough to capture all relevant differences in costs and outcomes. Sources of unit costs are not reported and may not be reflective of national unit costs.
<b>Overall applicability<sup>(b)</sup>:</b> partially applicable <b>Overall quality<sup>(c)</sup>:</b> potentially serious limitations

*Abbreviations: CC: comparative cost analysis; 95% CI: 95% confidence interval; CUA: cost–utility analysis; ICER: incremental cost-effectiveness ratio; NR: not reported; QALYs: quality-adjusted life years.*

- (a) *For studies where the time horizon is longer than the treatment duration, an assumption needs to be made about the continuation of the study effect. For example, does a difference in utility between groups during treatment continue beyond the end of treatment and if so for how long?*
- (b) *Directly applicable/Partially applicable/Not applicable.*
- (c) *Minor limitations/Potentially serious limitations/Very serious limitations.*

## Appendix F: GRADE tables

**Table 7: Clinical evidence profile: Stand-alone units versus absence**

Quality assessment							No of patients		Effect		Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Stand-alone units	Absence	Relative (95% CI)	Absolute		
<b>ED avoidance (follow-up mean 1 years; measured with: Mean monthly attendance rates; Better indicated by lower values)</b>												
2	observational studies	no serious risk of bias <sup>2</sup>	no serious inconsistency	no serious indirectness	serious <sup>1</sup>	None	not applicable	not applicable	not calculable	MD 194.83 lower (322 to 67.66 lower)	⊕○○○ VERY LOW	IMPORTANT
<b>ED avoidance (follow-up mean 1 years; assessed with: Annual attendance rates (per 1000 population))</b>												
1	observational studies	no serious risk of bias <sup>2</sup>	no serious inconsistency	no serious indirectness	serious <sup>1</sup>	None	not applicable	not applicable	1.17 (1.03 to 1.33)	not calculable	⊕○○○ VERY LOW	IMPORTANT

<sup>1</sup> Downgraded by 1 increment if the confidence interval crossed 1 MID, and downgraded by 2 increments if the confidence interval crossed both MIDs.

<sup>2</sup> All non-randomised studies automatically downgraded due to selection bias. Studies may be further downgraded by 1 increment if other factors suggest additional high risk of bias, or 2 increments if other factors suggest additional very high risk of bias.

**Table 8: Clinical evidence profile: Co-located on the same site as the ED versus absence**

Quality assessment							No of patients		Effect		Quality	Importance
No of studies	Design	Risk of bias	Inconsistency	Indirectness	Imprecision	Other considerations	Co-located with ED	Absence	Relative (95% CI)	Absolute		
<b>Avoidable adverse events (re-consultations) (follow-up 4 weeks; assessed with: Number of re-consultations - ED patients only)</b>												
1	observational studies	serious <sup>1</sup>	no serious inconsistency	no serious indirectness	no serious imprecision	none	54/115 (47%)	48.9%	RR 0.96 (0.77 to 1.2)	20 fewer per 1000 (from 112 fewer to	⊕○○○ LOW	CRITICAL

										98 more)		
<b>Avoidable adverse events (re-consultations) (follow-up 4 weeks; assessed with: Number of re-consultations (ED + WiC patients combined))</b>												
1	observational studies	serious <sup>1</sup>	no serious inconsistency	no serious indirectness	no serious imprecision	none	149/330 (45.2%)	48.9%	RR 0.92 (0.79 to 1.08)	39 fewer per 1000 (from 103 fewer to 39 more)	⊕○○○ LOW	CRITICAL
<b>ED avoidance (follow-up 6 months; measured with: Patient throughput (mean monthly attendances); Better indicated by lower values)</b>												
1	observational studies	no serious risk of bias <sup>3</sup>	no serious inconsistency	no serious indirectness	very serious <sup>2</sup>	none	not applicable	not applicable	not calculable	MD 542 higher (347 lower to 1431 higher)	⊕○○○ VERY LOW	IMPORTANT
<b>ED avoidance (follow-up 1 year; measured with: Mean monthly attendances); Better indicated by lower values)</b>												
1	observational studies	no serious risk of bias <sup>3</sup>	no serious inconsistency	no serious indirectness	serious <sup>1</sup>	none	not applicable	not applicable	not calculable	MD 349 lower (696 to 2 lower)	⊕○○○ VERY LOW	IMPORTANT
<b>Waiting time in ED (follow-up 3 months; assessed with: Cases complying with A&amp;E 4 hour waiting target)</b>												
1	observational studies	no serious risk of bias <sup>3</sup>	no serious inconsistency	no serious indirectness	no serious imprecision	none	743/785 (94.6%)	87.4%	RR 1.08 (1.05 to 1.11)	70 more per 1000 (from 44 more to 96 more)	⊕⊕○○ LOW	CRITICAL
<b>Waiting time in ED (follow-up 3 months; assessed with: Cases complying with A&amp;E 4 hour waiting target)</b>												
1	observational studies	no serious risk of bias <sup>3</sup>	no serious inconsistency	no serious indirectness	no serious imprecision	none	1466/1546 (94.8%)	87.4%	RR 1.09 (1.06 to 1.11)	79 more per 1000 (from 52 more to 96 more)	⊕⊕○○ LOW	CRITICAL
<b>Waiting time in ED (follow-up 3 months; assessed with: Cases complying with A&amp;E 4 hour waiting target)</b>												
1	observational studies	no serious risk of bias <sup>3</sup>	no serious inconsistency	no serious indirectness	no serious imprecision	none	743/785 (94.6%)	94.8%	RR 1 (0.98 to 1.02)	0 fewer per 1000 (from 19 fewer to 19 more)	⊕⊕○○ LOW	CRITICAL
<b>Waiting time in ED (follow-up 3 years; measured with: Average change in time to clinician (mins) per year; Better indicated by lower values)</b>												
1	observational studies	no serious risk of bias <sup>3</sup>	no serious inconsistency	no serious indirectness	no serious imprecision	none	not applicable	not applicable	not calculable	MD 11 lower (14 to 8 lower)	⊕⊕○○ LOW	CRITICAL
<b>Mortality (follow-up 3 years; measured with: Number dying per 1000 attendees; Better indicated by lower values)</b>												

1	observational studies	no serious risk of bias <sup>3</sup>	no serious inconsistency	no serious indirectness	unable to assess imprecision without MID values	none	not applicable	not applicable	not calculable	MD 1.8 higher (1.6 to 2 higher)	-	IMPORTANT
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<sup>1</sup> Downgraded by 1 increment if the majority of the evidence was at high risk of bias, and downgraded by 2 increments if the majority of the evidence was at very high risk of bias.

<sup>2</sup> Downgraded by 1 increment if the confidence interval crossed 1 MID or by 2 increments if the confidence interval crossed both MIDs.

<sup>3</sup> All non-randomised studies automatically downgraded due to selection bias. Studies may be further downgraded by 1 increment if other factors suggest additional high risk of bias, or 2 increments if other factors suggest additional very high risk of bias.

## Appendix G: Excluded clinical studies

**Table 9: Studies excluded from the clinical review**

Study	Exclusion reason
Banerjee 2012 <sup>2</sup>	Inappropriate comparison (audit – no comparator)
Beales 1995 <sup>3</sup>	Inappropriate comparison (descriptive article – no comparator)
Bickerton 2005 <sup>5</sup>	Inappropriate comparison. Incorrect interventions (retrospective case note analysis – no intervention or comparator)
Bickerton 2012 <sup>4</sup>	No relevant outcomes reported
Byrne 2000 <sup>6</sup>	Not review population (exclusively minor injuries patients)
CASPERS2016 <sup>7</sup>	Non-UK study (USA). Full text next available
Castledine 2008 <sup>8</sup>	Inappropriate comparison. Incorrect interventions (descriptive article – no intervention or comparator)
Chalder 2007 <sup>9</sup>	No extractable data
Chapman 2004 <sup>11</sup>	Systematic review is not relevant to review question or unclear PICO
Dale 1996 <sup>12</sup>	Inappropriate comparison (no comparator)
Davis 2005 <sup>13</sup>	Inappropriate comparison (descriptive article – no comparator)
Desborough 2012 <sup>14</sup>	Systematic review is not relevant to review question or unclear PICO
Freij 1996 <sup>16</sup>	No relevant outcomes reported
Gnani 2013 <sup>17</sup>	Inappropriate comparison (descriptive article – no comparator)
Grant 2002 <sup>18</sup>	Incorrect comparison (walk-in centre versus GP practice)
Gray 2003 <sup>19</sup>	Inappropriate comparison (descriptive study - no comparator)
Heaney 1997 <sup>20</sup>	No extractable outcome data
Ismail 2013 <sup>22</sup>	Systematic review is not relevant to review question or unclear PICO
Jackson 2005 <sup>23</sup>	Inappropriate comparison. No relevant outcomes (descriptive qualitative article – no comparator)
Land 2013 <sup>24</sup>	Inappropriate comparison. No relevant outcomes (patient survey – no comparator)
Marshall 1998 <sup>25</sup>	Inappropriate comparison (descriptive article – no comparator)
Mcintosh 1996 <sup>26</sup>	Inappropriate comparison (descriptive non-UK article – no comparator)
Paxton 1997 <sup>28</sup>	Incorrect interventions. Inappropriate comparison (patient survey – no comparator)
Roberts 1998 <sup>29</sup>	Systematic review is not relevant to review question or unclear PICO
Rourke 2009 <sup>30</sup>	Not review population (exclusively ENT conditions)
Rudge 2013 <sup>31</sup>	Inappropriate comparison (distance to MIU)
Sakr 2003 <sup>32</sup>	Not review population (exclusively minor injuries patients)
Salisbury 2002 <sup>36</sup>	Incorrect comparison (walk-in centre versus GP practice)
Salisbury 2002 <sup>34</sup>	Inappropriate comparison (descriptive article – no comparator)
Salisbury 2003 <sup>33</sup>	No extractable outcome data
Salisbury 2003 <sup>37</sup>	Systematic review is not relevant to review question or unclear PICO
Simpson 2001 <sup>38</sup>	Unclear intervention (centralisation of 3 A&E departments involving several changes including a new MIU and EAU)
Snooks 2004 <sup>39</sup>	Not review population (exclusively minor injuries patients)
Stark 2004 <sup>40</sup>	Inappropriate comparison (descriptive article – no comparator)

Study	Exclusion reason
Taylor 2008 <sup>41</sup>	Inappropriate comparison (descriptive article – no comparator)
Vaughan 2013 <sup>42</sup>	Incorrect interventions (nurse-led versus doctor-led hospital service)
VILLASENOR2016 <sup>43</sup>	Systematic review. No relevant references
Ward 2001 <sup>44</sup>	Inappropriate comparison (no comparator)
Weatherburn 2009 <sup>45</sup>	Not review intervention (telemetric cardiology service)
Welch 2009 <sup>46</sup>	Inappropriate comparison (descriptive article – no comparator)
Zimmerman 2013 <sup>47</sup>	Systematic review is not relevant to review question or unclear PICO



## **Appendix H: Excluded health economic studies**

No studies were excluded.