

LRI Children's Hospital

Community Acquired Pneumonia in Children

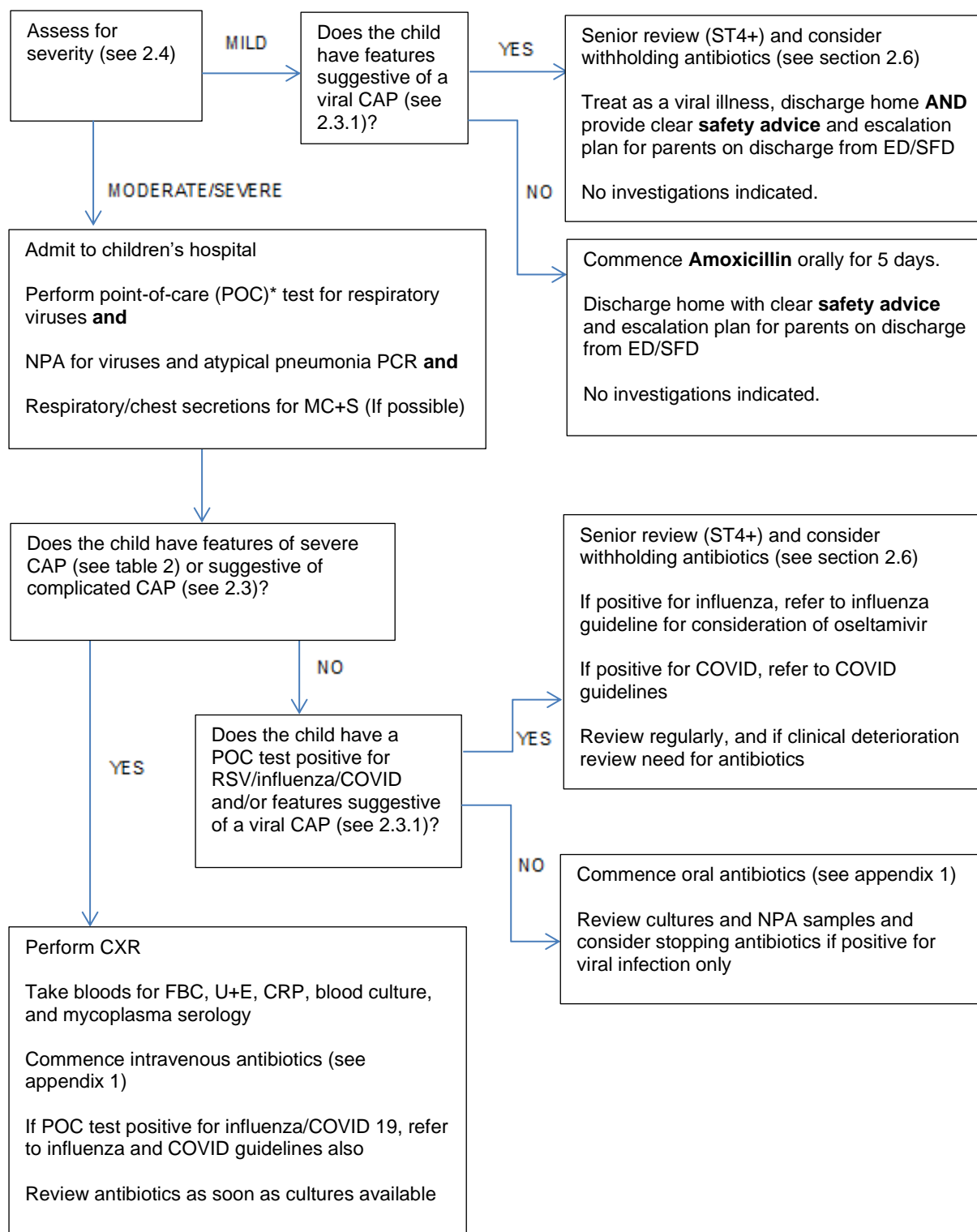
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2.5 INVESTIGATIONS	6
2.6 MANAGEMENT	7
2. 7 DISCHARGE CRITERIA	8
2.8 FOLLOW UP	9
3. Education and Training	9
4. Monitoring Compliance	9
5. Supporting References	9
6. Key Words	10
Contact and review details	11
APPENDIX 1: EMPIRICAL ANTIBIOTIC OPTIONS	12

Related guidelines:

- [Bronchiolitis UHL Childrens Hospital Guideline \(D11/2020\)](#)
- [Treatment of Children and Young People with Influenza like illness \(D10/2020\)](#)
- [Respiratory Viral Illness \(Including Flu\) Infection Prevention UHL Childrens Guideline \(D10/2019\)](#)
- [Upper Respiratory Tract Infection UHL Childrens Hospital Guideline \(D6/2020\)](#)
- [Cystic Fibrosis - Inpatient Chest Exacerbation UHL Childrens Medical Guideline \(C36/2016\)](#)
- [Cystic Fibrosis Emergencies UHL Childrens Medical Guidelines \(C64/2015\)](#)
- [Tuberculosis UHL Childrens Hospital Guideline \(C8/2019\)](#)
- [Empyema UHL Childrens Medical Guideline \(C127/2016\)](#)

MANAGEMENT ALGORITHM FOR CHILDREN WITH FEATURES SUGGESTIVE OF CAP (see 2.3)



2. Guideline Standards and Procedures

2.1 KEY PRACTICE POINTS

1. Bacterial pneumonia should be considered in children presenting with chest recession, raised respiratory rate, and pyrexia $>38.5^{\circ}\text{C}$
2. Blood tests are NOT recommended for routine use in the diagnosis and management of CAP, unless the child has symptoms and signs of severe pneumonia
3. Microbiological investigations should be performed in all patients admitted with CAP before commencing antibiotics as per NHSE recommendations
4. Chest X-Rays (CXR) should not be considered a routine investigation in children with mild to moderate CAP
5. Oral amoxicillin is the recommended first line antibiotic in children with mild CAP and those admitted with uncomplicated moderate bacterial CAP (unless not tolerated)
6. Follow up is not routinely indicated in children with mild to moderate

2.3.1 Viral versus bacterial CAP

Although viruses account for a significant proportion of childhood CAP (Jain 2015, Michelow 2004), mixed bacterial-viral infections are found in 23% to 33% of cases (Nascimento-Carvalho 2016, Michelow 2004, Cevey-Macherel 2009). Previous studies have consistently shown that it is not possible to distinguish viral and bacterial pneumonia by presenting symptoms, signs, or radiological changes with absolute certainty. However, clinical features which make either viral or bacterial CAP more likely have been identified in a number of studies (Nguyen 2019, Nascimento-Carvalho 2019, Bhuiyan 2019, Michelow 2004)

Table 1 Clinical Features Suggestive of Bacterial versus Viral CAP	
Bacterial	

Table 2 Severity Assessment (adapted from BTS 2011)			
	Mild CAP	Moderate CAP	Severe CAP
Infants	RR < 50/min CRT < 2 sec Mild recessions Taking full feeds	RR 50-70/min CRT ~2 sec Moderate recessions Reduced feeds (by 50%)	RR > 70/min CRT > 2 sec Nasal flaring Intermittent apnoea Grunting Unable to feed
Older Children	RR < 35/min CRT < 2 sec Mild breathlessness Taking full feeds	RR 35-50/min CRT ~2 sec Moderate recessions 	

- If present, pleural fluid should be sent for microscopy, culture, pneumococcal antigen, and atypical pneumonia PCR testing.
- Consider urinary pneumococcal/legionella antigen testing if ≥ 10 years old.

2.6 MANAGEMENT

All children with suspected **bacterial** CAP should receive antibiotics.

In children < 2 years admitted with moderate features of CAP, wheeze and temperature $< 38.5^{\circ}\text{C}$ a viral aetiology is more likely. Therefore consideration (by a registrar or Consultant) can be given to withholding antibiotics but the child should be monitored closely for signs of respiratory or cardiovascular deterioration. A history of conjugate pneum

- If unable to tolerate enteral fluids, commence intravenous (IV) fluids at two-thirds maintenance
 - Check Na, K, Urea, Creatinine at baseline, and then daily whilst on IV fluids to check for SIADH
- Analgesia to help keep the child comfortable and to allow deep breaths
- Chest physiotherapy has not been shown to reduce respiratory rate, severity score, or duration of hospitalisation and should not be routinely recommended

Failure to Respond

If the child is not improving despite being given antibiotics at an appropriate dose for 48 hours, consider:

- Complicated pneumonia (

2.8 FOLLOW UP

Children with complicated pneumonia (i.e. empyema) should be followed up after discharge until they have recovered completely and their chest x-ray has returned to normal – see empyema UHL Childrens Medical guideline.

Outpatient follow up is not otherwise routinely needed. However should be considered in children with:

- Severe pneumonia requiring a prolonged (more than 5 days) hospital stay
- Recurrent admissions/GP attendances with CAP

Follow up CXRs (in 6-8 weeks) are only otherwise indicated in the following circumstances:

- Lobar collapse
- Round pneumonia
- A child with persistent or recurrent symptoms

If unsure, discuss with the consultant responsible for the patient or with the respiratory team.

Good Practice Point – discuss prevention advice with carers i.e. vaccinations and smoking cessation in household.

3. Education and Training

None

4. Monitoring Compliance

What will be measured to monitor compliance	How will compliance be monitored	Monitoring Lead	Frequency	Reporting arrangements
Number of children prescribed empirical antibiotics as per guideline recommendations	Audit	DL	Annual	Departmental audit meeting
Number of children who received intravenous antibiotics who could have been managed with oral antibiotics as per severity assessment	Audit	DL	Annual	Departmental audit meeting

5. Supporting References

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6. Key Words

Pneumonia, Children

The Trust recognises the diversity of the local community it serves. Our aim therefore is to provide a safe environment free from discrimination and treat all individuals fairly with dignity and appropriately according to their needs. As part of its development, this policy and its impact on equality have been reviewed and no detriment was identified.

Contact and review details			
Guideline Lead (Name and Title) David Lo – Consultant in Paediatric Respiratory Medicine		Executive Lead Chief Medical Officer	
Details of Changes made during review:			
Date	Issue Number	Reviewed By	Description Of Changes (If Any)
July 2019	2	D Lo	See archived version July 2019 via paggs@uhl-tr.nhs.uk
September 2022	3	D Lo Approved by UHL Children’s Hospital Clinical Guidelines Group & UHL Antimicrobial Working Party	Removed admission as a specification for the scope of this guidance Updated management algorithm adding ref to COVID Reduced feeds parameters defined for assessment purposes Removed reference to flu season Added - atypical bacterial PCR, respiratory/chest secretion/ET aspirate for MC&S and consider urinary pneumococcal/legionella antigen - to testing recommendations Timing of when to start treatment recommendations and when to review treatment added Indications for respiratory review updated Added treatment advice for suspected aspiration pneumonia

APPENDIX 1: EMPIRICAL ANTIBIOTIC OPTIONS

First Line Antibiotics for Non-Severe CAP		
	Oral	Intravenous
No Penicillin Allergy	Amoxicillin	Amoxicillin
Allergy to Penicillin	Clarithromycin	Clarithromycin
First Line Antibiotics for Severe or Complicated CAP – START WITH INTRAVENOUS ANTIBIOTICS		
	Oral	Intravenous
No Penicillin Allergy	Co-Amoxiclav and Clarithromycin	Co-Amoxiclav and Clarithromycin**
Allergy to Penicillin (Non Severe*)	Discuss with microbiology	Cefuroxime and Clarithromycin**
Aspiration Pneumonia Suspected		
	Oral	Intravenous
No Penicillin Allergy	Co-Amoxiclav	Co-Amoxiclav
Allergy to Penicillin (Non Severe*)	Clarithromycin and Metronidazole	Cefuroxime and Metronidazole

***Non severe allergy = mild rash only, with no anaphylaxis. In patients with a severe history of penicillin allergy/reaction (i.e. anaphylaxis or Stevens Johnsons), discuss with microbiologists.**

****Oral Clarithromycin has good bioavailability and can be given via the oral route alongside IV Co-Amoxiclav or IV Cefuroxime provided the child is able to tolerate medications safely orally**

If there is no response to first line empirical antibiotic treatment consider adding a macrolide (clarithromycin) to cover for atypical pathogens, review available cultures and sensitivities, and/or discuss with microbiology for further advice.

For dosages refer to latest British National Formulary for Children (BNFC)