Systematic Review of Evidence-Based Guidelines on Medication Therapy for Upper Respiratory Tract Infection in Children with AGREE Instrument

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Abstract

Objectives: To summarize recommendations of existing guidelines on the treatment of upper respiratory tract infections (URTIs) in children, and to assess the methodological quality of these guidelines.

Methods: We searched seven databases and web sites of relevant academic agencies. Evidence-based guidelines on pediatric URTIs were included. AGREE II was used to assess the quality of these guidelines. Two researchers selected guidelines independently and extracted information on publication years, institutions, target populations, recommendations, quality of evidence, and strength of recommendations. We compared the similarities and differences of recommendations and their strength. We also analyzed the reasons for variation.

Results: Thirteen guidelines meeting our inclusion criteria were included. Huge differences existed among these 13 guidelines concerning the categorization of evidence and recommendations. Nearly all of these guidelines lacked the sufficient involvement of stake holders. Further, the applicability of these guidelines still needs to be improved. In terms of recommendations, penicillin and amoxicillin were suggested for group A streptococcal pharyngitis. Amoxicillin and amoxicillin-clavulanate were recommended for acute bacterial rhinosinusitis (ABRS). An observation of 2–3 days prior to antibiotic therapy initiation for mild acute otitis media (AOM) was recommended with amoxicillin as the suggested first choice agent. Direct evidence to support strong recommendations on the therapy for influenza is still lacking. In addition, the antimicrobial durations for pharyngitis and ABRs were still controversial. No consensus was reached for the onset of antibiotics for ABRs in children.

Conclusions: Future guidelines should use a consistent grading system for the quality of evidence and strength of recommendations. More effort needs to be paid to seek the preference of stake holders and to improve the applicability of guidelines. Further, there are still areas in pediatric URTIs that need more research.

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Introduction

Acute respiratory infections (ARIs) are classified as upper respiratory tract infections (URTIs) or lower respiratory tract infections (LRTIs) [1]. URTIs include the common cold, laryngitis, pharyngitis/tonsillitis, acute rhinitis, acute rhinosinusitis and acute otitis media (AOM) [2]. URTIs in children are a frequent illness accounting for a high proportion of doctor office visits [3,4]. A national survey report from the UK showed the consultation rates of URTIs were 3,103 and 1,002 per 10,000 person years at risk in children aged 0–4 and 5–15 years, respectively [5].

A proliferation of clinical guidelines published in peer-reviewed journals has been seen due to the high morbidity of URTIs. It is important that these guidelines provide appropriate guidance for the treatment of URTIs. Nevertheless, the growing number of guidelines has been accompanied with a growing concern about variance and conflicts among guideline recommendations and the quality of guidelines [18]. To date, there have been no systematic attempts to compare recommendations from available guidelines for the treatment of children with URTIs.

The aim of this study is to assess the quality of evidence-based guidelines for drug therapy of URTIs in children and to compare the recommendations of the existing evidence-based guidelines. Special attention was devoted to areas of disagreement and discussion with an ultimate aim to improve the clinical practice in treatment of URTIs for children. Such an assessment is important as it may explain some of the variability in guideline recommen-
dations and may assist health care providers in choosing among available guidelines.

**Methods**

**Data Sources**
We searched Pubmed, Guidelines International Network (GIN), U.S. National Guideline Clearinghouse (NGC) and four Chinese databases: Chinese Biomedical Literature Database (CBM), China Knowledge Resource Integrated Database (CNKI), VIP Database and Wanfang Database for evidence-based guidelines (until March 2013) using the following items: respiratory tract infections, common cold, laryngitis, pharyngitis, tonsillitis, rhinitis, rhinosinusitis, otitis media, middle ear inflammation, influenza, grippe as Medical Subject Headings (MeSH) or keywords. The searches were limited to guidelines published in English or Chinese. We also searched guidelines at web sites of academic agencies, such as American Academy of Pediatrics (AAP) and Infectious Diseases Society of America (IDSA). Retrieved references were considered if they met our inclusion criteria.

**Guideline Selection**

**Inclusion criteria.** Types: evidence-based guideline with systematic literature review and grading system for quality of evidence and/or strength of recommendation [6].
| Guidelines by Medical Condition | Country | Institution | Target Population | Conflicts of Interest | Method to Formulate Recommendations | Quality of Evidence | Strength of Recommendations | Reference |
|---------------------------------|---------|-------------|-------------------|----------------------|--------------------------------------|-------------------|-----------------------------|-----------|
| Pharyngitis                      | America | IDSA        | Adult and child   | SCI                  | Consensus development based on evidence | GRADE             | GRADE                       | [8]       |
| Rhinosinusitis                   | Finland | FSP         | Adult and child   | FPO                  | Consensus development based on evidence | Grading system from Evidence-based Medicine Working Group (A–D) | NA             | [9]       |
| Esposito 2008                    | Italy   | SIP         | child             | EI                   | Consensus development based on evidence | Self designed grading system in accordance with the Italian National Guidelines Plan (I–VI) | Self designed grading system in accordance with the Italian National Guidelines Plan (A–E) | [10]      |
| Chow 2012                        | America | IDSA        | Adult and child   | SCI                  | GRADE                               | GRADE             | GRADE                       | [11]      |
| Wald 2013                        | America | AAP         | child             | SCI                  | BRIDGE-Wiz²                         | Self designed grading system in accordance with the AAP policy statement (A–D,X) | Self designed grading system in accordance with the AAP policy statement “Classifying Recommendations for Clinical Practice Guidelines” (Strong recommendation, recommendation, option) | [12]      |
| Influenza                        | –       | WHO         | Adult and child   | SCI                  | GRADE                               | GRADE             | GRADE                       | [13]      |
| Bautista 2009                    | –       | WHO         | Adult and child   | SCI                  | Consensus development based on evidence | GRADE             | GRADE                       | [14]      |
| Harper 2009                      | America | IDSA        | Adult and child   | SCI                  | Consensus development based on evidence | CTFPHE            | CTFPHE                      | [15]      |
| Morciano 2009                    | Italy   | SNLG        | Adult and child   | FPO                  | Based on systematic review of available evidence | Self designed grading system in accordance with the Italian National Guidelines Plan (I–VI) | Self designed grading system in accordance with the Italian National Guidelines Plan (A–E) | [16]      |
| Otitis media                     | England | SIGN        | Child             | EI                   | Based on systematic review of available evidence | SIGN              | SIGN                        | [17]      |
| Takahashi 2012                   | Japan   | JOS         | Child             | NA                   | Based on available data             | Grading system from Japan Stroke Society (I–IV) | Grading system from the US Preventive Services Task Force report (A–E) | [18]      |
| Lieberthal 2013                  | America | AAP         | Child             | SCI                  | Consensus development based on evidence | Self designed grading system in accordance with the AAP policy statement (A–D,X) | Self designed grading system in accordance with the AAP policy statement “Classifying Recommendations for Clinical Practice Guidelines” (Strong recommendation, recommendation, option) | [19]      |
Diseases: URTIs including rhinosinusitis, pharyngitis, laryngitis, rhinitis, otitis media, tonsillitis, common cold and influenza.

Patients: children ages 0–18 years old

Interventions: drug therapy

Exclusion criteria. Types: guideline of hospital level; old version of guideline

Diseases: non-infectious upper respiratory diseases

Interventions: vaccines

Guideline Quality Assessment

Appraisal of guidelines with the AGREE instrument. Quality of evidence-based guidelines was assessed by using AGREE II from the following domains: scope and purpose, stakeholder involvement, rigor of development, clarity of presentation, applicability, editorial independence and overall guideline assessment. Each of the AGREE II items and the two global rating items were rated on a 7-point scale (1-strongly disagree to 7-strongly agree). A score was assigned depending on the completeness and quality of reporting. Domain scores were calculated by summing up all the scores of the individual items in a domain and by scaling the total as a percentage of the maximum possible score for that domain. The scaled domain score was calculated as: (obtained score-minimum possible score)/(maximum possible score-minimum possible score) [7].

Appraisal of agreement between reviewers. We used the intraclass correlation coefficient (ICC) as a measure of agreement between reviewers. The ICC was applied to each guideline. Calculations were carried out by using SPSS 13.0.

Data Extraction

Two researchers selected guidelines independently and extracted the following information: publication years, institutions, target populations, recommendations, quality of evidence, and strength of recommendations. We compared the similarities and differences of recommendations and their strength and analyzed the reasons for variation.

Results

Guideline Search and Review Process

A total of 1,785 citations and abstracts were identified in the initial searches. Finally, 13 guidelines meeting our inclusion criteria were included, covering a period from 2005 to 2013 (Figure 1). These 13 guidelines focused on drug therapy for pharyngitis, rhinosinusitis, otitis media and influenza. For the remaining URTIs (laryngitis, tonsillitis, rhinitis), no guidelines for children were found.

Characteristics and Quality of Guidelines

Half of the 13 guidelines are from America developed by the American Academy of Pediatrics (AAP) and the Infectious Diseases Society of America (IDSA). Five guidelines are developed specially for children, while the rest are for both adults and children. All guidelines, except one, announced the conflict of interests and none were founded by industrial partners. All guidelines stated the formulation of recommendations was based on evidence. However, there is a huge variation in the grading systems of evidence quality and recommendation strength used (Table 1).

Comparison of the Categorization of Evidence and Recommendations of the 13 Guidelines (Table 2)

There were huge differences among 13 guidelines concerning the categorization of evidence and recommendations. Eight
| Levels | CTFPHE | SIGN | GRADE | Self designed grading system in transition to GRADE² | Grading system from Evidence-based Medicine Working Group | Grading system from AAP | Grading system from Japan Stroke Society | Grading system from Italian National Guidelines System |
|--------|--------|------|-------|-----------------------------------------------|-------------------------------------------------|-----------------|-----------------------------|-------------------------------|
| Quality of evidence | 1 | I: Evidence from ≥1 properly RCT¹ | 1**: High quality meta-analyses, SR of RCTs, RCTs with a very low risk of bias. | High Quality Evidence: Further research is very unlikely to change our confidence in the estimate of effect. | High Quality Evidence: Further research is very unlikely to change our confidence in the estimate of effect. | A: Several relevant, high-quality scientific studies with homogeneous results | A: Well-designed RCTs or DS on relevant population | I: multiple RCTs and/or SR of randomized studies |
| | | | | | | | | |
| | | | | | | | Ib: At least one RCT |
| | 1: Well conducted meta-analyses, SR, RCTs with a low risk of bias. | 1**: Well conducted meta-analyses, SR, RCTs with a high risk of bias. |
| | | | | | | | |
| | | | | | | | II: Evidence from ≥1 well-designed clinical trial, without randomization; from cohort or case-controlled analytic studies; from multiple time-series; from dramatic results from uncontrolled experiments |
| | | | | | | | |
| | | | | | | | 2**: High quality SR of case control or cohort studies; High quality case control or cohort studies with a very low risk of confounding or bias and a high probability that the relationship is causal. |
| | | | | | | | |
| | | | | | | | Moderate Quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate. |
| | | | | | | | |
| | | | | | | | Moderate Quality Evidence: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate. |
| | | | | | | | |
| | | | | | | | B: At least 1 relevant, high-quality study or several adequate studies |
| | | | | | | | |
| | | | | | | | IIb: At least one well-designed, quasi-experimental study. |
| | | | | | | | |
| | | | | | | | II: one single adequate randomized trial |
| | | | | | | | |
| | | | | | | | 2: Well conducted case control or cohort studies with a low risk of confounding or bias and a moderate probability that the relationship is causal. |
| | | | | | | | |
| | | | | | | | 2: Case control or cohort studies with a high risk of confounding or bias and a significant risk that the relationship is not causal. |
| | | | | | | | |
| | | | | | | | 2: Non-analytic studies, eg case reports, case series. |
| | | | | | | | |
| | | | | | | | Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate. |
| | | | | | | | |
| | | | | | | | Low Quality Evidence: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate. |
| | | | | | | | |
| | | | | | | | C: At least 1 adequate scientific study |
| | | | | | | | |
| | | | | | | | C: OS |
| | | | | | | | |
| | | | | | | | III: At least one well-designed, non-experimental descriptive study |
| | | | | | | | |
| | | | | | | | III: non-randomized cohort, concurrent or historical studies, or their meta-analysis |
| | | | | | | | |
| | | | | | | | III: Evidence from opinions of respected authorities, based on clinical experience, descriptive studies, or reports of expert committees |
| | | | | | | | |
| | | | | | | | 3: Non-analytic studies, eg case reports, case series. |
| Levels | CTFPHE | SIGN | GRADE | Grading system from Evidence-based Medicine Working Group | Grading system from AAP | Grading system from Japan Stroke Society | Grading system from Italian National Guidelines System |
|--------|--------|------|-------|----------------------------------------------------------|--------------------------|------------------------------------------|-----------------------------------------------------|
| 4      | 4: Expert opinion | Very low quality: Any estimate of effect is very uncertain. | D: Expert opinion, case reports, or reasoning from first principles | IV: Expert committee reports, opinions and/or experience of respected authorities | IV: retrospective case-control studies |
| 5      |               |                                   |                                                                 | X: Exceptional situations in which validating studies cannot be performed and there is a clear preponderance of benefit or harm. | V: non-controlled case-series studies |
| 6      |               |                                   |                                                                 |                                  | V1: Expert opinion or opinions from panels as indicated in guidelines or consensus conferences |

### Strength of recommendation

| Levels | A: Good evidence to support a recommendation for or against use | B: Moderate evidence to support a recommendation for or against use | C: Poor evidence to support a recommendation |
|--------|---------------------------------------------------------------|---------------------------------------------------------------|----------------------------------------------|
| 1      | A: rated as 1++ or 1+, and directly applicable to the target population | B: rated as 2++, and directly applicable to the target population; or Extrapolated evidence from studies rated as 1++ or 1+ | C: Evidence level 3 or 4; or Extrapolated evidence from studies rated as 2++ |
| 2      | strong recommendation: the desirable effects of an intervention clearly outweigh the undesirable effects, or clearly do not | weak recommendation: the trade-offs are less certain--either because of low quality evidence or because evidence suggests that desirable and undesirable effects are closely balanced | |
| 3      | Strong Recommendation: quality of evidence is excellent; benefits strongly outweigh the harms | Recommendation: quality of evidence is not as strong; benefits exceed the harms | Option: suspect evidence or well-done studies but little clear advantage to one approach vs another |
| 4      | Strongly recommended; strong evidence is available, benefits substantially outweigh harms | Recommended: fair evidence is available, benefits outweigh harms. | No recommendation made: fair evidence is available, but the balance of benefits and harms is close |
| 5      | A: the specified strongly recommended and based on good quality scientific evidence | B: There are doubts as to whether the particular procedure should be always be recommended, but should be carefully considered. | C: There is substantial uncertainty concerning the procedure or intervention. |
different grading systems were used, two of which failed to give strength of recommendations. Four guidelines used GRADE [21–25]. One guideline used SIGN [26] and one used CTFPHE [27]. Seven guidelines used other grading systems. The variation in terms of grading system may decrease the comparability of guidelines and confuse readers.

Evaluation of the AGREE Domains of Guidelines Analyzed (Table 3)

**Scope and purpose.** This domain evaluates the overall objectives, expected benefit or outcomes, and target population of guidelines. The medium score for this domain was 90.28% (76.39%–97.22%), indicating that most guidelines satisfied criteria of this domain.

**Stake holder involvement.** This domain evaluates the degree of relevant professional group involvement and whether the views and preferences of the target population have been considered and the definition of target users has been clearly presented. The overall score in this domain was low with a medium of 61.11% (33.33%–83.33%). Most of the guidelines involved relevant professionals in the development process and declared the target population. However, the guideline developers did not seek the preference of target populations sufficiently resulting in a decrease in score of this domain.

**Rigour of development.** This domain addresses the method of evidence search, grading, summarizing and the formulation of recommendations. The medium score for this domain was 76.04% (32.81%–91.15%), with 1 guideline scoring 50%. This guideline failed to demonstrate the link between evidence and recommendations. It was not reviewed externally before its publication either.

**Clarity of presentation.** This domain evaluates presentation and format of guidelines. The medium score was 95.83% (90.28%–98.61%), indicating that all guidelines satisfied criteria of this domain.

**Applicability.** This domain evaluates the consideration of facilitators or barriers to its implementation, as well as monitoring criteria. The medium score of this domain was 56.25% (10.42%–83.22%), the lowest of all domains. Four of 13 guidelines scored ≤ 50%. Most guidelines failed to consider the applicability sufficiently in guideline development.

**Editorial independence.** This domain addresses founding issues and competing interests of guideline development members. The medium score was 81.25% (8.33%–97.92%), with 3 guidelines scoring ≤ 50%.

**Agreement among reviewers.** Table 3 summarizes the degree of agreement for 13 guidelines by ICC. The ICC values indicate overall agreement between reviewers was excellent (80%) for 8 of 13 guidelines and substantial (70%) for the other 5 guidelines.

**Recommendations**

**Recommendations towards drug therapy of Group A streptococcal pharyngitis for children (Table 4).** Recommendations on antibiotics from IDSA and ISCI are consensus. The first choice was penicillin or amoxicillin. For penicillin-allergic patients, cephalosporins, clindamycin, or macrolides were recommended. Only IDSA, however, gave a recommendation on the duration of antibiotics (10 days). In terms of adjunctive therapy, IDSA suggested nonsteroidal anti-inflammatory drugs (NSAIDs) as adjunct to an appropriate antibiotic for treatment of moderate to severe symptoms or control fever. Aspirin should be avoided in children due to the risk of Reye’s syndrome [28].

**Recommendations towards drug therapy of acute bacterial rhinosinusitis (ABRS)/sinusitis for children**
### Table 3. Quality Assessment by AGREE II of 13 Evidence-based Guidelines.

| Guidelines by Medical Condition, y | Domain 1 | Domain 2 | Domain 3 | Domain 4 | Domain 5 | Domain 6 | Agreement among reviewers for AGREE instrument items | Overall Assessment¹ |
|------------------------------------|----------|----------|----------|----------|----------|----------|---------------------------------------------------|---------------------|
|                                    | Scope and Purpose | Stakeholder Involvement | Rigour of Development | Clarity of Presentation | Applicability | Editorial Independence |                                                                  |                     |
| **Pharyngitis**                    |           |          |          |          |          |          |                                                                  |                     |
| Shulman 2012                       | 94.44     | 63.89    | 91.15    | 95.83    | 39.58    | 97.92    | 0.89                                                              | Y                   |
| **Rhinosinusitis**                 |           |          |          |          |          |          |                                                                  |                     |
| Blomgren 2005                      | 86.11     | 55.56    | 32.81    | 94.44    | 10.42    | 16.67    | 0.91                                                              | YM                  |
| Esposito 2008                      | 90.28     | 33.33    | 72.92    | 98.61    | 34.38    | 83.33    | 0.89                                                              | YM                  |
| Chow 2012                          | 90.28     | 63.89    | 72.40    | 98.61    | 83.33    | 87.50    | 0.91                                                              | YM                  |
| Wald 2013                          | 94.44     | 79.17    | 81.77    | 94.44    | 56.25    | 89.58    | 0.76                                                              | Y                   |
| **Influenza**                      |           |          |          |          |          |          |                                                                  |                     |
| Bellamy 2006                       | 86.11     | 56.94    | 84.38    | 95.83    | 60.42    | 81.25    | 0.78                                                              | YM(3Y,1YM)          |
| Bautista 2009                      | 93.06     | 61.11    | 72.40    | 95.83    | 55.21    | 77.08    | 0.83                                                              | YM(3Y,1YM)          |
| Harper 2009                        | 95.83     | 61.11    | 78.65    | 94.44    | 58.33    | 89.58    | 0.92                                                              | Y                   |
| Morciano 2009                      | 76.39     | 45.83    | 70.31    | 95.83    | 54.17    | 58.33    | 0.84                                                              | Y(3Y,1YM)           |
| **Otitis media**                   |           |          |          |          |          |          |                                                                  |                     |
| Takahashi 2012                     | 86.11     | 77.78    | 61.98    | 90.28    | 25.00    | 8.33     | 0.83                                                              | YM                  |
| Lieberthal 2013                    | 93.06     | 61.11    | 90.10    | 94.44    | 70.83    | 72.92    | 0.84                                                              | Y(3Y,1YM)           |
| Bain 2003                          | 84.72     | 62.50    | 76.04    | 95.83    | 66.67    | 62.50    | 0.76                                                              | Y(3Y,1YM)           |
| **URTI²**                          |           |          |          |          |          |          |                                                                  |                     |
| Snellman 2013                      | 97.22     | 83.33    | 83.33    | 94.44    | 72.92    | 95.83    | 0.77                                                              | Y(3Y,1YM)           |
| **Medium (range)**                 | 90.28 (76.39–97.22) | 61.11 (33.33–83.33) | 76.04 (32.81–91.15) | 95.83 (90.28–98.61) | 56.25 (10.42–83.33) | 81.25 (8.33–97.92) | /                                                                  | /                   |

Notes:
1. Y: Yes; YM: Yes, with modifications; N: No.
2. URTIs refer to guidelines which include multiple diseases in URTIs.

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Both WHO and IDSA placed a high value on the prevention of influenza rather than clinical trials on treatment of H5N1 patients. However, IDSA recommended zanamivir rather than oseltamivir, while WHO recommended oseltamivir for children (Table 5). All of the 5 guidelines supported the use of antibiotics in pediatric ABRS. However, AAP emphasized the onset of antibiotics should be in cases of severe onset or worsening course, while the other four recommended antibiotics for all clinical diagnosed ABRS. Guidelines from FSP and SIP recommended amoxicillin as first-line choice due to the low risk of treatment failure [10]. IDSA, however, suggested amoxicillin-clavulanic rather than amoxicillin as empiric antimicrobial therapy, considering the increasing prevalence of H. influenza among URTI of children and the high prevalence of β-lactamase-producing respiratory pathogens in ABRS [29,30]. For children with risk factors, amoxicillin-clavulanic was recommended. For non-type 1 hypertension, both ISCI and IDSA recommended doxycycline as an alternative for older children. Nevertheless, a variance appeared in terms of antibiotics for type 1 hypertension patients. ISCI and IDSA suggested levofloxacin, while AAP recommended cephalosporins based on recent studies which indicated the risk of a serious allergic reaction to cephalosporins in patients with penicillin or amoxicillin allergy appeared to be nil [31–33]. The duration of antibiotic therapy is still controversial [36]. SIGN recommended a 5 day course according to British National Formulary, while AAP recommended a 5 to 10 day course based on the age of children [35,37]. Further, only SIGN evaluated the efficacy and safety of adjunctive therapies [39].

**Recommendations towards drug therapy of influenza**

CTFPHE was first published in 1979 by Canadian Ministry of Health. The quality of evidence is based on the study design and the strength of recommendation depends on sufficiency of evidence. CTFPHE was the first grading system developed and is the foundation of many other grading systems. However, the CTFPHE still has some drawbacks [27]. For instance, a lack of strong relevance between quality of evidence and strength of recommendation and lack of consideration of results exist consistently among studies [39]. Consequently, many other

| Target of population | Shulman 2012, IDSA | Snellman 2013, ISCI |
|----------------------|-------------------|-------------------|
| **Antibiotics** | | |
| **Onset of antibiotics** | | |
| Diagnosis of pharyngitis | Culture positive cases of group A streptococcal pharyngitis |
| **Type of antibiotics** | | |
| **First-line** | Penicillin, amoxicillin (strong, high) | Penicillin, amoxicillin |
| **Second-line (penicillin allergy)** | A first-generation cephalosporin1, clindamycin, clarithromycin, azithromycin (strong, moderate) | Cephalosporins1, macrolides, clindamycin, amoxicillin-clavulaneate (2 low quality studies; 1 high quality study) |
| **Duration** | 10 days2 | – |
| **Adjuvant drugs** | | |
| **NSAIDs3** | | |
| For treatment of moderate to severe symptoms or control fever (high, strong) | – |
| **Corticosteroids** | | |
| NR* (moderate, weak) | – |

Notes:
1. The first-generation cephalosporins can be used for patients who are not anaphylactically sensitive.
2. Clarithromycin should be given for 5 days.
3. NSAIDs: nonsteroidal anti-inflammatory drug.
4. NR: not recommended.
5. doi:10.1371/journal.pone.0087711.t004

(Tables 5). All of the 5 guidelines supported the use of antibiotics in pediatric ABRS. However, AAP emphasized the onset of antibiotics should be in cases of severe onset or worsening course, while the other four recommended antibiotics for all clinical diagnosed ABRS. Guidelines from FSP and SIP recommended amoxicillin as first-line choice due to the low risk of treatment failure [10]. IDSA, however, suggested amoxicillin-clavulanic rather than amoxicillin as empiric antimicrobial therapy, considering the increasing prevalence of H. influenza among URTI of children and the high prevalence of β-lactamase-producing respiratory pathogens in ABRS [29,30]. For children with risk factors, amoxicillin-clavulanic was recommended. For non-type 1 hypertension, both ISCI and IDSA recommended doxycycline as an alternative for older children. Nevertheless, a variance appeared in terms of antibiotics for type 1 hypertension patients. ISCI and IDSA suggested levofloxacin, while AAP recommended cephalosporins based on recent studies which indicated the risk of a serious allergic reaction to cephalosporins in patients with penicillin or amoxicillin allergy appeared to be nil [31–33]. The duration of antibiotic therapy is still controversial [36]. SIGN recommended a 5 day course according to British National Formulary, while AAP recommended a 5 to 10 day course based on the age of children [35,37]. Further, only SIGN evaluated the efficacy and safety of adjunctive therapies [39].

**Discussions**

Variation of Evidence and Recommendation Grading System

Citicorps was first published in 1979 by Canadian Ministry of Health. The quality of evidence is based on the study design and the strength of recommendation depends on sufficiency of evidence. CTFPHE was the first grading system developed and is the foundation of many other grading systems. However, the CTFPHE still has some drawbacks [27]. For instance, a lack of strong relevance between quality of evidence and strength of recommendation and lack of consideration of results exist consistently among studies [39]. Consequently, many other
organizations developed their own grading systems [40,41]. SIGN grading system developed by the Scottish Intercollegiate Guidelines Network is one of the most-widely used systems [26]. However, it still failed to consider the consistency and indirectness of study results. In 2000, GRADE working group was founded based on organizations from 19 countries including WHO. The aim of this group is to develop a consolidated grading system for quality of evidence and strength of recommendation. In 2004, the first edition of this grading system was published and was recognized by more than 30 organizations including WHO and Cochrane Collaboration. Although the 13 guidelines included in our study were published after 2005, only four of them use the GRADE system. We suggest further guidelines use a comparable uniform grading system to evaluate the quality of evidence and strength of recommendations.

Quality of Guidelines

The potential benefits of guidelines are only as good as the quality of the guidelines themselves. Appropriate methodologies and rigorous strategies in the guideline development process are important for the successful implementation of the resulting recommendations [7]. For these 13 guidelines, two domains are the main problems which decrease the quality and reliability of guidelines. The first is a failure to seek patients’ views and preferences or fail to report this information. Many methods can be used to consider patients’ values such as: formal consultations with patients, participation of patients on guideline development group or external review group, or a literature review of patients’ values. However, these processes were seldom performed or described in guideline development or the final reports of guidelines. This problem is also found in other disease guidelines [42,43,44]. The second problem is a lack of consideration of applicability of guidelines. How to facilitate the application of guidelines is as important as how to develop a high quality guideline.

Table 5. Main Therapeutic Options on Acute Bacterial Rhinosinusitis (ABRS)/Sinusitis for Children According to Guidelines.

| Therapy recommended | Blomgren 2005, FSO | Esposito 2008, SIP | Chow 2012, IDSA | Wald 2013, AAP | Snellman 2013, ISCI |
|---------------------|--------------------|--------------------|-----------------|-----------------|-------------------|
| Target of population | Children >1 year | Children >1 year | Children | Children aged 1–18 years | Children |
| Antibiotics | | | | | |
| Onset of antibiotics | Clinical diagnosis of ABRS (B) | Clinical diagnosis of ABRS (I,A) | Clinical diagnosis of ABRS (strong, moderate) | Severe onset or worsening course ABRS (B, strong recommendation) | Clinical diagnosis of ABRS (high quality) |
| Types of antibiotics | | | | | |
| First-line | Amoxicillin | Mild: Amoxicillin (IV,B) | Amoxicillin-clavulanate (Strong, moderate) | Amoxicillin or amoxicillin-clavulanate (B, Recommendation). | Amoxicillin-clavulanate (high dose may consider in children <2 years) (Guideline). |
| Second-line | | | | | |
| With risk factors1 | – | Mild: Amoxicillin-clavulanic or cefalor IV, B | Amoxicillin-clavulanate (weak, moderate) or third-generation oral cephalosporin+clindamycin (weak, moderate). | Amoxicillin-clavulanate – |
| Hypersensitivity | | | | | |
| Non-type I | – | – | Third-generation oral cephalosporin+clindamycin (weak, moderate), doxycycline (>8 years) | Cefdinir, cefuroxime, cefpodoxime | Doxycycline (for older children), Levofloxacin |
| Type I | – | – | Levofloxacin (weak, low). | Cefdinir, cefuroxime, cefpodoxime, or cefixime +clindamycin |
| Durations | 7 days | Mild: 10–14 days (IV,B) | 10–14 days (weak, low-moderate), | 10–28 days | 3–14 days (Low quality evidence) |
| Adjunctive drugs | | | | | |
| Corticosteroids | Recommended to allergic patients | NR (II,A) | Recommended to allergic patients (weak, moderate) – | Recommended for recurrent or allergic patients (high quality evidence) |
| Decongestants | NR2 | NR (II,A) | NR(strong, low-moderate) – | NR |
| Antihistamines | NR | NR (II,A) | NR(Strong, low-moderate) – | NR |

Notes:
1Risk factors include: previous receive of antibiotic therapy; attendance at school, local or systematic diseases that favor infections due to antibiotic-resistant pathogens; from geographic regions with high endemic rates of invasive penicillin-nonsusceptible (PNS) S. pneumonia; severe infection; age <2; recent hospitalization; immunocompromised.
2NR: not recommended.

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The facilitators and barriers that may impact the application of guidelines should be considered when developing the guideline. Also, there is a need to consider how to disseminate and implement the guideline effectively using additional materials such as a quick reference guide, educational tools and patient leaflets. These factors are important but often ignored by guideline developers. Studies on the effectiveness of clinical guideline implementation strategies showed that successful guideline implementation strategies should be multifaceted, and actively engage clinicians throughout the process [45,46]. Thus, future guidelines should pay more attention to the implementation process of guidelines.

Factors Contributing to Inconsistencies of Guidelines

Although an important level of consensus appears throughout the various guidelines, there are still some conflicts in recommendations for drug choice and durations of therapy. There are three main reasons contributing to the variances. First, the geographic difference leads to the variance of pathogens and its drug resistance. Second, the recommendations of guidelines were based on different evidence. Recent studies may overturn the results of previous studies. Thus, the timely updated guidelines are more reliable. Third, the expectation and preference of guideline developers and patients may influence the final recommendation. Therefore, a local guideline is more useful for health professionals if there is a conflict among guidelines.

Suggestions for Future Research

The durations of antimicrobial therapy for rhinosinusitis and acute otitis media are still controversial. More studies are needed to compare the different durations of antibiotics in children. In addition, the antivirals for influenza also lack direct evidence. Many recommendations are based on indirect evidence. Thus, more clinical trials or prospective observational studies are needed.

Conclusions

Future guidelines should use a consistent grading system for quality of evidence and strength of recommendations and seek the preference of stake holders to improve the applicability of guidelines. Further, there are still some areas in pediatric URTI that need more research.

Supporting Information

Checklist S1. (DOC)
Table 7. Main Therapeutic Options on Acute Otitis Media (AOM) for Children According to Guidelines.

| Therapy recommended | Bain 2003, SIGN | Takahashi 2012, JOS | Lieberthal 2013, AAP |
|---------------------|----------------|-------------------|---------------------|
| Target population   | Children       | Children <15 yrs  | Children aged 6 mos-12 yrs |
| Antibiotics         |                |                   |                     |
| Onset of antibiotics| Mild: observation for 3 days without use of antimicrobial agents (A) | Mild: observation for 3 days without use of antimicrobial agents (A) | Mild AOM in children (>2 yrs) antibiotic therapy or observation for 2-3 days (B, Recommendation). Mild bilateral AOM in children (6-23 months): antibiotic therapy (B, Recommendation). Mild unilateral AOM in children (6-23 months): antibiotic therapy or observation for 2-3 days (B, Recommendation). Severe: antibiotic therapy (B, Strong Recommendation). |
| Types of antibiotics| Amoxicillin, amoxicillin-clavulanic, cefadroxil, cotrimoxazole, trimethoprim, erythromycin (B) | Amoxicillin, amoxicillin-clavulanic, ampicillin, cefditoren, cefixime (A) | First-line antibiotic (B, Recommendation). Second-line: an antibiotic with additional β-lactamase (C, Recommendation). |
| Duration            | 5 days (A), >2 years (B) | –                  | <2 years: 10 days; 2-5 years: 7 days; >6 years: 5-7 days |
| Adjunctive          | Decongestants | –                  | –                  |
|                     | Antihistamines| –                  | –                  |
|                     | Paracetamol   | Recommended for analgesia (A) | –                  |

Notes:
1. *The* child does not receive amoxicillin in the past 30 days or does not have concurrent purulent conjunctivitis or the child is not allergic to penicillin.
2. *The* child has received amoxicillin in the last 30 days or has concurrent purulent conjunctivitis, or has a history of recurrent AOM unresponsive to amoxicillin.
3. NR: Not Recommended.

*Parents should give paracetamol for analgesia but should be advised of the potential danger of overuse.

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Conceived and designed the experiments: L. Zhang. Performed the experiments: L. Zeng ZY YL MC. Analyzed the data: L. Zeng ZH. Wrote the paper: L. Zeng. Revised the language of the paper: EAE.

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