Influenza burden, prevention and treatment in asthma – a scoping review
by the EAACI Influenza in Asthma Task Force

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Abstract
To address uncertainties in the prevention and management of influenza in people with asthma, we performed a scoping review of the published literature on influenza burden; current vaccine recommendations; vaccination coverage; immunogenicity, efficacy, effectiveness and safety of influenza vaccines; and the benefits of antiviral drugs in people with asthma. We found significant variation in the reported rates of influenza detection in individuals with acute asthma exacerbations making it unclear to what degree influenza causes exacerbations of underlying asthma. The strongest evidence of an association was seen in studies of children. Countries in the European Union currently recommend influenza vaccination of adults with asthma; however, coverage varied between regions. Coverage was lower among children with asthma. Limited data suggest that good seroprotection and seroconversion can be achieved in both children and adults with asthma and that vaccination confers a degree of protection against influenza illness and asthma related morbidity to children with asthma. There were insufficient data to determine efficacy in adults. Overall, influenza vaccines appeared to be safe for people with asthma. We identify knowledge gaps and make recommendations on future research needs in relation to influenza in patients with asthma.

Introduction
Asthma is a heterogeneous chronic inflammatory lung disease characterized by intermittent airflow obstruction and increased reactivity to bronchoconstrictive triggers. Presently, an estimated 300 million people worldwide suffer from asthma (1). In Europe, more than 30 million people are affected, with six million suffering severe symptoms and 1.5 million life-threatening attacks (2). Asthma onset and acute asthma exacerbations (AAE) have been strongly associated with respiratory viral and (to a lesser extent) bacterial infections (3-11).

Influenza viruses cause respiratory infections, typically during winter months. Most significant disease is caused by influenza A and B, which may escape immune pressure by undergoing antigenic drift and recombination; such genetic changes lead to new subtypes, which can cause epidemic or pandemic outbreaks (12, 13). Each year, influenza virus infection affects 5 to 20% of the global population (14), while annual vaccination greatly contributes to reduce related morbidity and mortality (15). Upon vaccination, coordinated activity of innate and adaptive immune responses leads to generation of short- and long-lived antibody producing plasma cells and memory B cells. The importance of influenza as a cause of serious illness and AAE is increasingly appreciated (16, 17). Despite uncertainties about their effectiveness, influenza vaccines are widely recommended for those with asthma (18).

To address uncertainties over the prevention and management of influenza in patients with asthma, we performed a scoping review and discuss the current knowledge of the burden imposed by influenza on patients with asthma, current vaccine recommendations and vaccination coverage in various regions, the evidence regarding immunogenicity, efficacy, effectiveness and safety of influenza vaccines, and the possible benefits of antiviral drugs in patients with asthma. We conclude by identifying knowledge gaps and future research needs in relation to influenza in asthma patients.

1. Methods
We systematically searched relevant published, unpublished and in-progress original research and systematic reviews that were pertinent to the subject matter of this EAACI
Task Force. This involved searching databases of peer-reviewed published literature (Cochrane Library, EMBASE and CINAHL, MEDLINE and Google Scholar) and research in progress (i.e. [http://www.controlled-trials.com/](http://www.controlled-trials.com/) and [http://www.clinicaltrials.gov/](http://www.clinicaltrials.gov/)) for the period 2000–2015. In addition, we wrote to a panel of international experts in an attempt to identify additional unpublished or in-progress work. There were no restrictions on language of publication.

We were particularly interested in randomized controlled trials (RCTs) on questions relating to the immunogenicity, efficacy and/or effectiveness of influenza vaccine among asthmatics, and epidemiological studies relating to the influenza incidence/prevalence and public health burden and safety of influenza vaccine. Systematic reviews with or without meta-analysis were of interest if they focused on any of the questions of interest.

The searches (see Appendix 1) were undertaken and allocated to relevant sub-group leads depending on the focus of the inquiry. Where possible, studies reported in languages other than English were translated. Within subgroups, two reviewers independently reviewed the titles and if necessary abstracts of potentially eligible studies to decide on eligibility; disagreements on study eligibility were resolved through discussion among the reviewers. Studies were then – in the specific context of asthma/allergy – categorized by topic area, namely: i) Global burden of influenza; ii) Influenza vaccination recommendations and coverage; iii) Immunogenicity of influenza vaccines; iv) Efficacy of influenza vaccines; v) Effectiveness of influenza vaccines; vi). Safety of influenza vaccines; and vii). Antiviral treatments against influenza. Reports were then scrutinized and interpreted by experts on the subject matter. Any disagreements on issues to do with interpretation were resolved through discussion.

The searches were first conducted in 2014 and then updated in October 2015. The key characteristics and findings of all eligible studies were summarized in data tables, and the findings narratively synthesized. These are presented by topic in Table 1.

2. Global burden of influenza in patients with asthma

2a. Influenza-attributable health care utilization in asthma patients

Asthma is the most common underlying disease in patients with influenza admitted to healthcare facilities, both in adults (19-33) (7.6–46% (34, 35)) and children (26, 36-47) (8.3–42% (48, 49)). Children with asthma accounted for a significantly larger proportion of those hospitalized with pandemic H1N1 than with seasonal influenza (24, 50-53). Asthma was also the most common comorbidity among pregnant women hospitalized with H1N1 infection, with a prevalence ranging from 8-33% (54-57).

Asthma patients seemed to have a higher risk of being admitted to the intensive care unit (ICU) or to have a more severe disease course when hospitalized for influenza (26, 33, 43, 45, 50, 66, 68-70), with an OR of 4.92 (95% CI 1.38–17.33) in Argentinian children (69) and a RR of 15.20 (95% CI 1.16-199.63) in China >15 years of age (71). Severe disease was especially common in those developing pneumonia (33), but did not seem to be related to the severity of underlying asthma (51). In a multivariate
analysis of H1N1 pediatric patients, asthma was associated with increased mortality in ICU-admitted children and adolescents (OR 1.34, p=0.05) (68). However, a retrospective analysis performed in the UK on 1,520 patients showed that patients with asthma hospitalized because of influenza A infection were less likely to require ICU or die compared with those without asthma, with an OR of 0.51 (95%CI 0.36–0.72) for severe outcome (28). Two other studies confirmed this finding (25, 29), which may be even more pronounced in those with pre-admission inhaled (OR 0.34; 95% CI 0.18–0.66) or systemic corticosteroid treatment (OR 0.36 95% CI 0.18–0.72) for asthma and an earlier hospital admission (within 4 days from symptom onset - adjusted OR 0.60 95%CI 0.38–0.94) (28).

2b. Influenza-induced acute asthma exacerbations

Influenza, like other respiratory viral infections, can lead to AAE and was found to account for 32.0% (72) and 9.8% (73) of AAEs in two studies of adults with asthma outpatients. However, the method used to identify the presence of influenza in AAE episodes largely determines the outcome. A direct comparison of molecular (PCR), serological, and immunofluorescent (IFA) detection in the same cohort of adults with asthma found influenza rates of 24.0%, 18.3%, and 2.0% respectively in AAEs (74). Thus, the validity of studies that applied conventional influenza identification techniques such as cytopathic effect in cell culture followed by indirect immunofluorescence to confirm viral presence (75) is questionable. Among hospitalized adults with asthma patients, rates of influenza-induced AAE were reported at 10.0% (76), 19.0% (77), 20.7% (78), and as high as 37.9% (73).

Available data on influenza-associated AAE among children vary considerably. For outpatient visits, three studies did not detect any influenza-positive samples (75, 79, 80), while several groups identified influenza infection as the cause of AAE, with most reporting rates of influenza-positive samples between 2.4% and 15.5% (63, 81-84). A recent study showed an accentuated propensity for loss of asthma control when children were infected with pH1N1 (38%) compared with other common respiratory virus infections (85). Among children with asthma hospitalized for severe AAE, three studies reported an influenza-associated burden of 2.6% (82), 3.4% (86) and 3.7% (63) of AAEs. In summary, there were significant inter- and intra-group variations in the reported influenza-induced AAE burden among children and adults with asthma, with fewer studies on adults.

Various factors compromise the interpretation of the available data. First, the use of different diagnostic methods for influenza detection may significantly influence the study outcome. The advent of molecular techniques and their universal use may solve this issue. Second, the absence of harmonized criteria for appropriate inclusion of patients and the lack of a standardized definition and diagnosis of influenza infection disallow direct comparison of results and complicate the identification of relevant studies. Third, the lack of large cohorts of patients with asthma studied for influenza induced AAE, raises questions about the validity of the currently available findings. Furthermore, seasonal rather than perennial influenza testing and selective testing only in hospitalized AAE cases could lead to inaccurate estimation of the actual contribution of influenza to AAEs.

3. Current influenza vaccine recommendations and coverage for people with asthma

In common with most economically-developed nations (Fig. 1), all 28 countries of the European Union currently recommend influenza vaccination of people with chronic respiratory disease, including asthma (87-89). Trivalent (TIV) and quadrivalent
inactivated split-virus influenza vaccines are available, with TIV the most widely used throughout the EU. An intradermal vaccine is also available and more recently a quadrivalent live attenuated intranasal vaccine (LAIV) was approved for use in the UK in children aged 2–18 years (88, 90).

However, vaccine uptake for clinical risk groups varied from 31% in Portugal to 82% in Northern Ireland (88, 91), as did influenza vaccination coverage of adults with asthma, with 30–50% in Spain (92-95) and 40% in the UK (96). Self-reported uptake of seasonal influenza vaccine amongst adults with asthma in the US varied from 54% in the community to 71% in secondary care (97). Children with asthma had significantly lower coverage rates of approximately 18-20% in Spain (93, 98), 15% in France (99) and 2.5–20% in Italy, across multiple seasons (91). In comparison, in the USA vaccination coverage reached around 50% in adults and children (100).

To address the question of such historically low vaccination rates amongst those with asthma, many studies employed cross-sectional surveys to analyse the socio-demographic variables that predict vaccine uptake. The strongest factors associated with higher vaccine coverage, were older age, recent contact with healthcare providers, non-smoking status, presence of co-morbidities, more severe asthma, and (surprisingly) lower household income (92-95, 100-106), although regarding there are conflicting reports for the latter (98). Variables that were associated with lower vaccination uptake included younger age groups (96, 100, 107, 108), belonging to certain ethnic groups (102, 105, 108) and lack of awareness about the vaccine (91, 99, 100). A more detailed exploration assessing health attitudes suggested that the misbelief that the vaccination was not necessary or could actually cause influenza or side effects was predictive of non-adherence (97, 109-112) and contributed to lower vaccination coverage (96, 107, 108). Reports of low effectiveness probably also contributed to poor uptake (18, 113, 114).

Healthcare worker endorsement of vaccination was highly predictive of adherence (109, 110, 112). A survey of 1225 US allergists suggested that 85% of these offered influenza vaccinations to adults and children. Younger allergists (<45 years) were more likely to offer vaccination, especially to children (115). Pediatricians were more likely than family physicians to routinely recommend influenza vaccination to children with asthma (adjusted OR 3.49, 95% CI 1.68-7.22), with decisions also being made depending on asthma phenotype (i.e. intermittent vs. persistent) (116).

The success of vaccination programmes was affected by complex factors such as the nature of healthcare provision (e.g. free care or insurance based), education levels, and cultural acceptance of the vaccine. Therefore, national health surveys with information on vaccination status linked with medical and socio-demographic details, (such as that of Spain with data spanning more than two decades (92-95, 117) can aid in assessing the impact of influenza vaccination, at population level.

Simple interventions such as allowing for year-round scheduling of appointments for vaccination can be effective (118). Nevertheless, it is likely that only major policy shifts will lead to significantly higher vaccination rates, due to more clear and consistent messaging about the importance and acceptability of influenza vaccination to patients and healthcare workers alike. This is illustrated by the increase in influenza vaccination uptake among children with asthma from 36% (2005–2006) to 50% (2010–2011) after the US Centre for Disease Control and prevention (CDC) advised vaccinating all children, regardless of risk (100). With its favourable safety and cost-benefit profile (18, 119, 120) there is a need for health campaigns to raise awareness amongst the public and clinicians of the benefits of influenza vaccination and to improve implementation of guidelines.
4. Immunogenicity of influenza vaccines in asthma

There is uncertainty about the level of protection that influenza vaccines afford in people with asthma. Influenza vaccine immunogenicity, a prerequisite for protection, is usually assessed by haemagglutinin inhibition assay (HI) measuring seroprotection rate (percentage of subjects with antibody titres ≥1:40), seroconversion rate (percentage of subjects with ≥4-fold increase in antibody titre), antibody geometric mean titre (GMT), and GMT ratio (GMT post vaccination/GMT pre-vaccination).

We identified limited data on influenza vaccine immunogenicity in asthma. Only one study of TIV in children aged 0.5–3 years compared healthy controls to recurrent wheezers. It found good and similar seroconversion and seroprotection rates to all three vaccine strains (H1N1, H3N2, and B) in both groups (after 1 month), with higher pre-and post-vaccination GMTs in children with wheeze, but a higher GMT ratio in healthy controls (121). In children with asthma, TIV and a virosomal inactivated subunit influenza vaccine induced good seroprotection and seroconversion at 1 month in both pre-school and school-age children, with some reduction in GMTs by 6 months (122, 123). A 2009 H1N1 vaccine given to adolescents and adults induced equally good seroprotection if 15 µg or 30 µg vaccine were given in those with mild-to-moderate asthma. In contrast, in severe asthma the higher dose resulted in better seroprotection, which in elderly severe asthma patients (≥60 years) was only achieved with the high 30 µg dose (124). In the latter group and in young children with recurrent wheeze, corticosteroid treatment or corticosteroid dose did not influence influenza vaccine immunogenicity (121, 124).

5. Efficacy of influenza vaccines in asthma

Data on the efficacy of influenza vaccination in asthma are limited, with only two randomized studies addressing the efficacy of TIV in patients with stable asthma. One study, conducted in Turkey, included 128 adults with asthma who received either TIV or no vaccination (125). No significant difference in the frequency of upper respiratory tract infections (URTI) or AAE rate was found between vaccinated and non-vaccinated participants; however, this study had significant limitations (see Table 1). In a larger randomized, double-blind, placebo-controlled trial from the Netherlands (81) in 6–18 year-old children with asthma, TIV had no significant effect on the number, severity, or duration of AAEs caused by confirmed influenza infection, or on the respiratory symptoms recorded; however, it moderately improved quality of life during the weeks of influenza-positive illness (126). In contrast, a conference abstract (127) reported significantly lower AAE rates in children <5 years with asthma following TIV (13%) compared with placebo (53%).

For LAIV, we found no efficacy data in asthma compared to non-vaccination. A large European multi-center, randomized, open-label trial compared intranasal cold-adapted LAIV to intramuscular TIV in children and adolescents aged 6–17 years with asthma, and reported a 34.7% reduction in culture-confirmed influenza illness after LAIV (128). A re-analysis of efficacy data from two randomized, multinational trials (129, 130) comparing LAIV to TIV, demonstrated that 2–5 year old children with mild-to-moderate asthma or prior wheezing, receiving LAIV had fewer cases of culture-confirmed influenza illness than TIV recipients (131).

6. Effectiveness of influenza vaccination in asthma

While efficacy studies reflect a treatment's performance in tightly controlled clinical trials, effectiveness studies reflect how well a treatment works in practice and are more representative of real life. Influenza vaccination may be less effective than expected
due to differences between the characteristics of patients vaccinated in the general population and those recruited into efficacy trials. Furthermore, the influenza vaccine may not contain the prevalent virus strain, and vaccine coverage may be suboptimal.

There are limited data on the effectiveness of influenza vaccination in patients with asthma or high-risk populations that included asthma. Six studies assessed the effectiveness of seasonal influenza vaccines, while one other studied the H1N1 pandemic vaccine (132).

In children, two retrospective cohort studies found that influenza vaccination was associated with a reduction in the rate of asthma exacerbations (133) and fewer acute respiratory problems (only in those less than 6 years of age) (134). A survey found that vaccination in children with asthma was associated with less use of rescue prednisolone (135), while a nested case-control study suggested that influenza vaccination was associated with fewer acute general practice visits for respiratory problems in children with long-term conditions (136). However, another case-control study suggested that vaccination was associated with an increased risk of unscheduled attendance for asthma care (137). In this study, the cases and controls were poorly balanced in terms of ethnicity and likelihood of vaccination in previous years. Although an adjusted analysis was undertaken, this may not have taken into account all the differences between the groups.

Only one study focused on adults with asthma (132). In this study, the H1N1 influenza vaccine was found to be protective against influenza in those aged under 61 years, but not in older patients. Also, it did not protect against AAE. Two other studies enrolled adults with long-term (136) or high-risk conditions (138); the first demonstrated a reduction in general practice visits, hospitalization and deaths from respiratory problems, while the second demonstrated a protective effect against influenza and influenza related hospitalization. Asthma was not analysed separately in these studies and it is thus not clear if influenza vaccination was beneficial for people with asthma.

A recent systematic review reports a 59-78% reduction in asthma attacks leading to emergency visits and/or hospitalizations after influenza vaccination and a vaccine effectiveness of 45% (95% CI 31 to 56) for laboratory-confirmed influenza in people with asthma based on a metanalysis of two test-negative design case-control studies (139).

7. Safety of influenza vaccines in asthma

Large retrospective and prospective studies have demonstrated the safety of influenza vaccines in children and adults with asthma, for both TIV and LAIV (131, 140-142). In these studies, vaccinated patients did not have more frequent asthma exacerbations than unvaccinated patients, and no other adverse event was more common in asthmatics. A study of adult asthma patients who reported bronchospasm after influenza vaccination, indicated that these symptoms were not related to vaccine hypersensitivity (143).

A common concern is the safety of egg-derived influenza vaccines in patients with egg-allergy, a frequent co-morbidity in children with asthma. The last case of anaphylaxis after influenza vaccination occurred more than 25 years ago at a time when the egg content of vaccines was much higher. The ovalbumin content of most current vaccines is less than 1.2 μg/mL, and an egg-free influenza vaccine is also available. Several studies demonstrated that influenza vaccines containing low levels of egg protein (≤1.2μg/mL) were safe even in egg-allergic patients (143-149). This also seemed to be true in egg-allergic patients with associated asthma. The recent SNIFFLE studies of LAIV in egg allergic children, including 634 children with asthma/recurrent wheeze (79.7% on daily inhaled corticosteroid treatment), found no immediate systemic allergic
reactions (150, 151). For 6.6% of children with asthma/recurrent wheeze, the parents reported wheeze within 72 hours of LAIV, but most wheeze episodes only required routine treatment; there was no hospitalization and no increase in respiratory symptoms in the four weeks after LAIV.

Based on the robust evidence that influenza vaccines are safe in asthma patients with and without egg allergy, British immunization guidelines (152) now recommend administration of LAIV in the community to children from 2 years of age, including those with asthma, irrespective of egg allergy, unless they have severe or acutely exacerbated asthma. US and Canadian guidelines remain more conservative and still suggest that vaccines containing egg proteins (e.g. LAIV) should not be used in children with egg allergy and/or asthma (153-155).

8. Antiviral treatment against influenza in patients with asthma

For asthma patients with suspected influenza infection, treatment with antiviral medication as early as possible, prior to laboratory confirmation of influenza, is currently recommended (156-160). Since amantadine and rimantadine are limited by drug resistance, they have been replaced by neuraminidase inhibitors (NI). The currently marketed NIs are zanamivir and oseltamivir. However, post-marketing surveillance has revealed bronchospasm and worsening of lung function in a few cases when asthma patients were treated with zanamivir inhalation (161). Therefore, the manufacturer does now not recommend zanamivir for patients with chronic respiratory disease, including asthma (157, 158). Oseltamivir was not associated with any significant respiratory adverse effects and is recommended for treatment and chemoprophylaxis of influenza in children and adults with high-risk chronic conditions, including asthma (162-164). However, large-scale telephone surveys in the USA showed that having asthma, in contrast to diabetes, was not associated with increased rates of antiviral treatment of influenza cases (165).

Data on the efficacy of NI in asthma patients (162, 166-168) are limited. A benefit of oseltamivir treatment was found in children in an RCT that showed significantly improved pulmonary function and fewer AAEs (168). A meta-analysis, based on a single RCT, reported improved FEV1 after oral oseltamivir given to children with asthma and influenza infection, but no effects on peak flow or AAEs (169). Two recent systematic reviews found that oseltamivir relieved influenza symptoms in otherwise healthy children, but had no effect on children with asthma who have influenza-like illness (170, 171), a finding that is currently controversial.

As high rates of resistance limit the effectiveness of the currently available antiviral drugs, new NI (laninamivir, peramivir) and an RNA-dependent RNA polymerase inhibitor (favipiravir) are currently undergoing clinical trials (172). Laninamivir showed similar efficacy and safety to oseltamivir in the treatment of influenza in asthma patients (173).

9. Conclusions

We observed significant variation in the reported rates of influenza detection among AAE episodes in both children and adults, with higher rates observed in the latter age group. This may be attributed to major differences in inclusion criteria, disease definitions, virus detection methodology, small population sizes and the limited number of available studies. Therefore, it is presently unclear to what degree influenza causes exacerbation of underlying asthma.

All European Union countries currently recommend influenza vaccination of people with asthma, with TIV being the most widely used vaccine. Nonetheless, vaccination
coverage varies between regions, and is significantly lower among children with asthma. Several factors and misbeliefs have been shown to predict non-adherence to influenza vaccination, while healthcare worker endorsement highly promotes vaccination. Major policy shifts and health campaigns may help in raising awareness and improve guideline implementation.

The limited immunogenicity data suggest that good seroprotection and seroconversion can be achieved in children and adults with asthma. However, higher vaccine doses may be required in elderly patients. Available studies suggest increased efficacy of LAIV in reducing influenza illness in children/adolescents with asthma compared to TIV, but do not provide evidence of efficacy of TIV in reducing influenza illness in asthmatics concurring with a recent systematic review (18).

There are some limited data to suggest that influenza vaccination reduces asthma-related morbidity in children in real-life settings, as has been concluded in another recent systematic review (139). The only data to suggest that vaccination may have a beneficial impact in adults with asthma are from studies of patients with long-term or high-risk conditions, not just asthma. The effectiveness outcomes of these studies are difficult to interpret due to lack of an agreed definition for asthma and/or influenza, and insufficient information on the type of vaccination used. Lack of randomization is also a limitation, as differences in the characteristics of those who do and do not receive the vaccine, such as disease severity and socioeconomic status, may independently impact on the likelihood of asthma morbidity, and analyses may not control for all these differences. Despite these limitations, population-based effectiveness studies are the only way to assess the impact of vaccination in real-life healthcare settings.

Inactivated influenza vaccines appear to be safe for people with asthma, and, provided they have low egg protein content (≤1.2 μg/mL), also for those with egg allergy. LAIV is also safe in asthma and egg allergy with the exception of severe or acutely exacerbated asthma, where LAIV cannot be given. Oseltamivir is the NI that is currently used in asthma. Although it has moderately beneficial effects on influenza illness in healthy people, there are insufficient data to determine its efficacy and effectiveness in people with asthma. However, novel NIs are currently in development.

10. Recommendations and future needs

People with asthma are at high risk of more severe forms of influenza, with increased hospitalization and admission at healthcare facilities, therefore:

- Education campaigns for both the general population and healthcare providers are needed to promote an increase in annual influenza vaccination, early case recognition and consideration of early empirical antiviral therapy, and to reduce influenza-attributable healthcare utilization among asthmatics.

- Large cohort and case control studies using standardized definitions (for AAE), sensitive influenza detection methods, such as PCR, and comprehensive assessment of respiratory pathogens are necessary in both children and adults with asthma.

- There is a need to develop more internationally co-ordinated registries, e.g. for Europe, that accurately collate vaccination status along with medical and socio-demographic details to assess the impact of public health interventions over time.

- Multiple sources of data must be used to record vaccination status (e.g. self-reported as well as medical notes, vaccination card or billing data).
- Targeted vaccination of children and at-risk groups is needed.
- Further studies of influenza vaccine immunogenicity, comparing healthy and asthmatic populations, are required to determine if people with asthma have normal influenza vaccine responses or if they require influenza vaccines with enhanced immunogenicity. The discovery and development of improved immune correlates of protection against influenza will enable more meaningful testing of vaccine immunogenicity.
- In view of the uncertainty around the degree of protection that influenza vaccination affords in people of all ages with asthma, further efficacy data need to be collected in this population. Data should be stratified by age and asthma severity, using adequately powered, randomised, placebo-controlled trials that compare both TIV and LAIV to non-vaccinated controls.
- Further studies are needed to evaluate the effectiveness of influenza vaccination in patients with asthma, but must be designed to minimize any risk of bias.
- Safety data in asthma with and without egg allergy are required for new influenza vaccines, and, depending on egg protein content, for new LAIV batches.
- There is a need to collect more safety data on zanamivir in patients with chronic respiratory disease, and efficacy data on oseltamivir in children with asthma.
- Development and testing of new anti-influenza drugs among asthmatics is needed.

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Author contributions

All authors are members of the EAACI Influenza and Asthma Task Force. Skevaki and Schwarze conceived and designed the study. Aziz Sheikh developed the search protocol. Ather Akhlaq, Ulugbek Nurmatov performed the systematic literature search, reviewed and selected the relevant publications. All authors except the former three performed data extraction, compiled the summary tables, contributed to interpretation of results and contributed to writing the first draft of the manuscript, which was reviewed by Skevaki and Schwarze and then revised by all authors. All authors approved the final version of the manuscript.

Author conflict of interest

The authors declare no conflict of interest.
Table 1. Studies included in the systematic review, by section

**Section 2a. Influenza-attributable health care utilization in asthma patients**

| Authors, Year       | Country     | Study Period                  | Type of study                        | Intervention | Outcome measure                                                                 | Key finding                                                                                                                                                                                                 | Population characteristics                                                                 | Limitations/ comments                                                                 |
|---------------------|-------------|------------------------------|--------------------------------------|--------------|---------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------|-------------------------------------------|
| Chacon et al. 2015  | Guatemala   | May 2009 - June 2010         | Epidemiological retrospective study   | NA           | Demographic and clinical characteristics of deaths associated with influenza A (H1N1) pdm09. | History of Asthma in 11% of the 112 dead patients with a pre-existing medical condition (7% children < 5 years, 11% people aged 5-59, 16% people aged 60+ and 18% of the 14 dead pregnant or postpartum women) | 183 children and adults aged 0-60+ years. SARI decedents who tested positive for influenza. | Study performed in seven Central America Countries. |
| Dawood et al. 2014  | USA         | 2003-2009 (seasonal) 2009-2010 (H1N1) | Epidemiological retrospective, population-based study | NA           | Frequency and severity of influenza complications in hospitalized children.     | History of asthma in 31% of children hospitalized for seasonal influenza vs. 42% in children hospitalized for influenza A (H1N1). Asthma exacerbation as a complication in 27% of children with influenza A vs. 18% in children with seasonal influenza (overall, 22% of total). | 6769 children and adolescents aged <18 years hospitalized for both seasonal and H1N1 influenza. |                                                                                          |
| Hagerman et al. 2015| Switzerland | June 2009 - Jan 2010         | Prospective/retrospective study performed in 11 hospitals | NA           | Clinical characteristics of the influenza-hospitalized children.               | Asthma 12% among the comorbidities of the 126 influenza-hospitalized children.                                                                                                                        | 326 patients aged <18 years (189 <5 years; 126 with comorbidities hospitalized in eleven children’s hospitals in Switzerland with a positive influenza A/H1N1/09 RT-PCR from a | Study performed in 11 hospitals in Switzerland. |
| Authors, Year | Country | Study Period | Type of study | Intervention | Outcome measure | Key finding | Population characteristics | Limitations/ comments |
|--------------|---------|--------------|---------------|--------------|-----------------|-------------|--------------------------|----------------------|
| Jiang et al. 2015 (71) | China | Dec 2008 - June 2014 | Epidemiological retrospective study | NA | Clinical characteristics and severe case risk factors for adult inpatient cases of confirmed influenza. | Out of the 240 (7.8%) cases of laboratory-confirmed influenza, (H1N1, pdm2009 and H3N2), asthma was the most important risk factor for severe cases (RR = 15.200, 95% CI: 1.157-199.633). | 240 adults aged ≥15 years consistent with SARI case definition who were monitored by SARI sentinel hospitals in ten cities in China. | Data extrapolated from abstract in English. Study performed in ten provinces of China. |
| Kwon et al. 2014 (176) | South Korea | May 2010 - Apr 2011 | Retrospective study | NA | Prevalence of year-round respiratory viral infection in children with LRTI and the relationship between respiratory viral infection and allergen sensitization in exacerbating asthma. | Influenza virus was detected in 7 (25%) of the 28 asthma patients hospitalized for asthma exacerbation. 18 asthmatics (64.3%) had atopic sensitization. Influenza virus responsible for asthma exacerbations with an OR 11.9 (1.5-90.4) for atopic asthma), 0.9 (0.1-12.3) for non-atopic asthma, 4.0 (1.2-13.8) for asthma in general. | 309 children (median age 26 months) who were hospitalized for acute LRTIs (pneumonia, bronchitis, bronchiolitis or asthma). |
| Park et al. 2013 (177) | South Korea | Jan - Dec 2009 | Multi-site case-control study | NA | Association of asthma status and severity of H1N1 influenza in adults. | Of the 91 cases, 12(13%) and 3 of 54 controls (5%) were asthmatics (p=NS); of those without other comorbidities except asthma, 8 of 56 cases (14%) and 2 of 49 (4%) | 91 cases (47.3±21.1 years) with a positive PCR for H1N1 influenza and were admitted to the ICU or general ward with a |
| Authors, Year | Country | Study Period | Type of study | Intervention | Outcome measure | Key finding | Population characteristics | Limitations/ comments |
|---------------|---------|--------------|---------------|--------------|----------------|-------------|-----------------------------|----------------------|
| Placzek et al. 2014 (178) | USA | Oct 2008 - Apr 2009 (seasonal influenza)  
Apr 2009 - Sept 2009 (H1N1) | Epidemiological retrospective study | NA | Association of age and comorbidity on 2008-2009 seasonal influenza and 2009 Influenza A (H1N1)-related ICU stay. | controls had asthma (p=NS). | | diagnosis of H1N1 influenza.  
56 controls (45.0±20.6 years) with a positive PCR for H1N1 influenza, but were not admitted to hospitals. |
| Stripeli et al. 2015 (179) | Greece | Oct 2009 - Feb 2010 and Jan 2011 - May 2011 (pH1N1)  
2002-2003 and 2004-2005 (Influenza A) | Epidemiological retrospective study | NA | Epidemiological and clinical characteristics of children hospitalized with pH1N1 compared with those of children hospitalized with seasonal influenza A in previous years. | Asthma was the most prevalent risk factor for hospitalization, occurring in 28% (n=148) of the pH1N1 population, and 24% (n=174) of those with seasonal influenza. | | 1254 children and adults aged <5-64 years with seasonal and H1N1 influenza admitted to an ICU. |
| Taha 2014 (57) | Saudi Arabia | June 2009 - Feb 2010 | Retrospective cohort study | NA | Characteristics of pregnant Women admitted with 2009 H1N1 Influenza in a referral maternity hospital. | Bronchial asthma was present in 15 (45.5%) of pregnant women admitted with influenza. | | 33 pregnant women: mean age 27.7±5.6 years. |
| Authors, Year | Country | Study Period | Type of study | Intervention | Outcome measure | Key finding | Population characteristics | Limitations/ comments |
|---------------|---------|--------------|---------------|--------------|-----------------|-------------|--------------------------|----------------------|
| Tam et al. 2014 (180) | USA | 2007-2011 | Retrospective population-based study (chart reviews) | NA | Influenza-related hospitalization of adults associated with low census tract, socioeconomic status and female sex. | Asthma was increasingly prevalent among cases with progressively higher census tract poverty status (16.4% in low poverty vs. 37.7% in high poverty). Females overall and in each age group were more likely to have asthma (overall, 27.6% vs. 16.7%; P<0.001). | 1094 hospitalized adults aged 18-65+ years with laboratory-confirmed influenza. |

ICU intensive care unit; NS not significant; LRTI lower respiratory tract infection; RR risk ratio; CI confidence interval; OR odds ratio; RT-PCR reverse transcriptase polymerase chain reaction; SARI severe acute respiratory infection.
## Section 2b. Influenza-induced acute asthma exacerbations

| Authors, Year       | Country   | Study Period          | Type of study | Intervention | Outcome measure                                    | Key finding (absolute number of positive samples) | Population characteristics | Limitations/ comments*            |
|---------------------|-----------|-----------------------|---------------|--------------|----------------------------------------------------|-------------------------------------------------|-----------------------------|---------------------------------|
| Azevedo et al. 2003 (75) | Brazil    | ND                    | Retrospective | NA           | Laboratory confirmed IFV/ culture and IFA          | 0.0% (0)                                        | 36 children aged 2-14 years    |                                 |
| Thumerelle et al. 2003 (83) | France    | 1 Oct 1998 - 30 June 1999 | Prospective   | NA           | Laboratory confirmed IFV/IFA, serology or both    | 3.6% (3)                                        | 82 inpatients aged 2-16 years   |                                 |
| Biscardi et al. 2004 (86) | France    | 1 Jan 1999 - 30 June 2001 | Prospective   | NA           | Laboratory confirmed IFV/culture and IFA          | 3.4% (4)                                        | 119 inpatients aged 2-15 years   | 0.0% (0)                        |
| Bueving et al. 2004 (81) | Netherlands | 1999-2001            | Double-blind, randomized, placebo-controlled trial | Inactivated influenza vaccine vs placebo | Laboratory confirmed IFV/ culture, IFA and RT-PCR | 7.2% (17)                                        | 235 children aged 6-18 years    | Low number of IFV positive samples in first season compensated by recruitment of higher number of subjects in second season |
| Johnston et al. 2005 (79) | Canada    | 10-30 Sept 2001       | Prospective   | NA           | Laboratory confirmed IFV/PCR                        | 0.0% (0)                                        | 52 children aged 5-15 years     |                                 |
| Ketsuriani et al. 2007 (80) | USA       | Mar 2003 - Feb 2004   | Prospective   | NA           | Laboratory confirmed IFV/PCR                        | 0.0% (0)                                        | 65 children aged 2-17 years     | Modest sample size              |
| USA 2002-2004         | Prospective | NA                   |               |             |                                                    | 15.5% (38)                                       | 245 outpatients                |                                 |
| Authors, Year | Country | Study Period | Type of study | Intervention | Outcome measure | Key finding (absolute number of positive samples) | Population characteristics | Limitations/ comments* |
|---------------|---------|--------------|---------------|--------------|----------------|-----------------------------------------------|---------------------------|------------------------|
| Miller et al. 2008 (63) |        |              |               |              | Laboratory confirmed IFV/PCR or culture | 3.7% (19) | 513 inpatients aged 6-59 months |                        |
| Vallet et al. 2009 (84) | France  | 1 Oct 2005 - 30 Nov 2007 | Prospective | NA | Laboratory confirmed IFV/IFA | 2.4% (4) | 166 inpatients aged 2-15 years |                        |
| Kwon et al. 2013 (176) | Korea  | Apr 2010 - Apr 2011 | Prospective | NA | Laboratory confirmed IFV/RT-PCR | 25.0% (7) | 28 inpatients aged 60±34 months (Mean±SD) | Modest sample size, selection of the study population, short duration (1 year) of the study, single medical center, and subjects were limited to children with LRTIs severe enough to require hospitalization. |
| Mandelcwajg et al. 2010 (82) | France  | Nov 2005 - Mar 2009 | Retrospective | NA | Laboratory confirmed IFV/IFA | 14.0% (15) | 107 outpatients | 2.6% (6) |
| Wark et al. 2001 (72) | Australia/UK  | 2 Feb - 1 Dec 1998 | Prospective | NA | Laboratory confirmed IFV/RT-PCR, serology, culture and IFA | 24% (12) | 49 outpatients aged 16-74 years |                        |

**ADULTS**
| Authors, Year         | Country       | Study Period | Type of study | Intervention                                      | Outcome measure                                             | Key finding (absolute number of positive samples) | Population characteristics | Limitations/ comments*                                                                 |
|----------------------|---------------|--------------|---------------|--------------------------------------------------|------------------------------------------------------------|------------------------------------------------------|--------------------------------|-------------------------------------------------------------------------------|
| Simpson et al. 2003  | Australia/UK  | ND           | Prospective   | NA                                               | Laboratory confirmed IFV/PCR or serology or IFA            | PCR screening: 24.0% (12) Serology screening: 18.4% (9) IFA screening: 2.0% (1) | 49 inpatients aged 16-74 years | Uneven sex distribution (prevalence of females in acute asthma group). |
| Tan et al. 2003      | Singapore     | ND           | Prospective   | NA                                               | Laboratory confirmed IFV/RT-PCR, dot blot and DIG-probe hybridization | 20.7% (6)                                            | 29 inpatients aged 42±15 years (Mean±SD) |                                                                              |
| Iikura et al. 2014   | Japan         | May 2011-Dec 2012 | Prospective   | NA                                               | Laboratory confirmed IFV/RT-PCR                            | 10.0% (5)                                            | 50 inpatients aged 28-92 years | Although 98% of patients had a common cold history receding their AAE, a pathogen was detected from nasopharyngeal swabs in only 70% of cases. Overall, 15 major viruses and 6 major bacteria were evaluated by multiplex PCR analysis. It is possible that other minor pathogens may be involved in AAE. Other potential |


| Authors, Year | Country | Study Period | Type of study | Intervention | Outcome measure | Key finding (absolute number of positive samples) | Population characteristics | Limitations/ comments* |
|--------------|---------|--------------|---------------|--------------|----------------|-------------------------------------------------|---------------------------|-------------------------|
| Teichtal et al. 1997 (77) | Australia | Aug 1993-July 1994 | Prospective | NA | Laboratory confirmed IFV/Culture and serology | 19.0% (15) | 79 inpatients aged 16-66 years | reasons for the low yield include inappropriate timing of sampling, lower respiratory tract infections, misinterpretation of allergic rhinitis symptoms, and single testing upon admission. |
| Atmar et al. 1998 (73) | USA | 6 Dec 1991-3 May 1994 | Prospective | NA | Laboratory confirmed IFV/Culture, serology and RT-PCR | 37.9% (11) | 29 inpatients aged 19-50 years | |
| | | 9 Dec 1992-31 May 1993 | Cross-sectional | | | 9.8% (12) | 122 outpatients aged 17-77 years | |

*as reported in the original article
# Current influenza vaccine recommendations for people with asthma

| Authors, year | Country | Study Period | Type of study | Intervention | Outcome measure | Key findings | Population characteristics | Limitations/comments |
|---------------|---------|--------------|---------------|--------------|----------------|--------------|---------------------------|----------------------|
| Grohskopf et al. 2014 (181) | USA | 2013-2014 | National Recommendations after reviewing literature based on GRADE approach | LAIV and IIV in asthma | Efficacy and safety | Advise that children between 2-4 years who have asthma or wheezing in previous 12 months should not be given LAIV. | Adults and children including those with asthma. Population size unavailable. |
| Grohskopf et al. 2015 (153) | USA | 2014 - 2015 | National Recommendations after reviewing literature based on GRADE approach | LAIV and IIV in asthma | Efficacy and safety | - | All adults and children including those with asthma. Population size unavailable. | No change from 2014 - 2015 |
| Byington et al. 2014 (182) | USA | 2013 - 2014 | National Recommendations from American Academy of Paediatrics | LAIV and IIV in asthma | Efficacy and safety | Annual vaccination (trivalent or quadrivalent) recommended for everyone with special effort made towards asthmatics. Asthma present in 26% hospitalized children with confirmed influenza during the 2013-14 season. | All children aged >6 months including those with asthma. Population size unavailable. |

LAIV live attenuated influenza vaccine; IIV inactivated influenza vaccine
### Section 3b. Current influenza vaccine coverage for people with asthma

| Authors, year   | Country           | Study Period         | Type of study                                      | Intervention                                                                 | Outcome measure                                                                                      | Key findings                                                                                          | Population Characteristics | Limitations/ comments                                                                                   |
|-----------------|-------------------|----------------------|---------------------------------------------------|-----------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------|-----------------------------|----------------------------------------------------------------------------------------------------------|
| Koshio et al. 2014 (132) | Japan             | Sept-Oct 2010        | Questionnaire-based cross-sectional survey        | NA                                                                          | Receipt of H1N1 vaccine and infection status as well as recording of ACT score.                      | Incidence of H1N1 pdm09 infection 6.7% (95% CI 5.7–7.6). Vaccine receipt in 63.9% (95% CI 62.1–65.8%). Infected patients (n=170) had significantly lower rates of vaccination, were younger and had experienced more night symptoms and sleep disturbance in the preceding 2 weeks. | 2,555 asthma patients, aged ≥16 years. | Good characterization of asthmatics but no objective definition of influenza infection. |
| Pennant et al. 2015 (183) | USA               | 2009-2015            | Prospective interventional quality improvement study | Physician reminders, patient letters, nurse-driven                          | Change in vaccination uptake rate.                                                                 | Vaccine receipt improved from 59% to 64% between 2011 and 2014 influenza seasons.                        | 1142 asthma patients in an allergy clinic aged ≥13 years (85 declined vaccine). |                                                                                                           |
| Gnatiu et al. 2015 (184) | Global (23 countries) | 2008 onward          | International cross sectional survey comparing high income and low-middle income countries | Participants with doctor diagnosed asthma underwent reversibility testing with salbutamol | Use of respiratory medicines and uptake of influenza vaccine in preceding 12 months.                | Overall influenza vaccination uptake rates: 1.4% in LMIC vs. 28.4% in HIC ($P = 0.0001$).              | 15,590 adults with asthma and COPD, aged ≥40 years (unclear what proportion are asthmatic). | Rare example of research comparing multiple countries from across the income spectrum The age criteria for the study likely excluded a large number of patients. |
| Authors, year | Country | Study Period | Type of study | Intervention | Outcome measure | Key findings | Population Characteristics | Limitations/ comments |
|---------------|---------|--------------|---------------|--------------|-----------------|--------------|----------------------------|------------------------|
| Vernacchio et al. 2014 (185) | USA | 2009-2012 | Learning collaborative model | Regular face to face meetings and electronic communication between healthcare teams | Uptake of influenza seasonal vaccination. Uptake of asthma action plans. | Inconsistent improvements in rates of vaccine uptake. Significant increase in implementation of asthma action plans. | 594 patients with persistent asthma, aged 5-17 years. |
| Blackwell et al. 2015 (186) | USA | 2009-2010 | Cross-sectional survey | Telephone survey | Receipt of vaccine in previous 12 months. | Asthmatics receiving H1N1 vaccine: OR 1.13 (CI 0.97-1.33). | 667 children, aged 4 months to 17 years, including asthmatics. |
| Aigbogun et al. 2015 (187) | Global | 2014 | Systematic review | NA | Uptake of vaccination rates in high-risk children. | Reminder letters useful in improving uptake but less evidence for telephone recall. | 9 studies in children with asthma, aged 6 months to 19 years. |
| Mazurek et al. 2014 (188) | USA | 2006-2009 | Cross-sectional survey | Telephone survey | Influenza vaccination uptake in the preceding 12 months. | 42.7% uptake rate. Higher rates in those aged 50-64 years, unemployment and in those requiring urgent treatment for worsening symptoms. | 28,809 adults aged 18-64 years with work-related asthma. |
| O'Halloran, 2015 (189) | USA | 2012-2013 | Cross-sectional survey | Telephone survey | Influenza vaccination uptake in the preceding 12 months. | Uptake rate of 29% (95% CI 26.5-31.7%). | 8,831 adults, aged 18-64 years with "pulmonary disease". |
| Lin et al. 2015 (190) | USA | 2010-2013 | Prospective interventional quality | Utilized the "4 pillars | Influenza vaccination rates pre and post intervention. | Uptake rates increased between 2010-11 to 2012-2013 seasons in: Non-White 8.250 asthmatic children aged 6 months-18 years. This intervention reduced the racial... |
| Authors, year | Country | Study Period | Type of study | Intervention | Outcome measure | Key findings | Population Characteristics | Limitations/ comments |
|--------------|---------|--------------|---------------|--------------|-----------------|-------------|---------------------------|----------------------|
| Shoup et al. 2015 (191) | USA | 2012-2013 | Pragmatic 3-arm randomized control trial | Three arms: 1. Postcard reminder 2. Telephone reminder 3. Postcard & telephone reminder | Change in influenza vaccination rate. | No difference in intervention groups. Telephone (interactive voice response) calls were most cost-effective. | 12,285 adults aged 19-64 years with asthma or COPD. | Disparity seen in vaccination uptake rates. |
| Buyuktiryaki et al. 2014 (109) | Turkey | 2010 | Cross-sectional survey of parents | Questionnaire | H1N1 and seasonal influenza vaccination rates with analysis of demographic factors, asthma control parameters and parental attitudes towards vaccination. | 16.8% received H1N1 vaccine; 45.7% received seasonal vaccine. Significant factors influencing uptake included low educational background of parents, previous vaccination with seasonal influenza and having a family member vaccinated against Influenza A/H1N1. Asthma control parameters had no influence on uptake of the vaccine and physician recommendation (84.8%) was important in the decision-making process for immunization. | 625 children and adolescents aged 6-18 years with asthma. | |
| Skupin et al. 2015 (192) | USA | 2013 | Cross-sectional survey | Questionnaire | Influenza vaccination uptake in preceding year. | Black patients had much lower vaccination rates | 472 adults attending internal | |
| Authors, year | Country | Study Period | Type of study | Intervention | Outcome measure | Key findings | Population Characteristics | Limitations/ comments |
|---------------|---------|--------------|---------------|--------------|-----------------|--------------|----------------------------|---------------------|
| Hart et al. 2014 (193) | USA | 2013 | Cross-sectional survey | Structured interview | Vaccination uptake in previous year and parental intent to vaccinate in upcoming flu season. | (62.1%) than White (83.8%) or Asian (93.3%) patients. Belief in efficacy in vaccine and physicians recommendation important determinants in higher vaccination rates. | medicine and allergy clinics. | 224 children, aged 6 months-17 years, attending paediatric emergency department. Unclear what proportion are asthmatic at baseline. |
| Lo, 2014 (194) | USA | 2013-2014 | Prospective interventional quality improvement study | Implementation of influenza immunization bundle | Vaccination rate pre- and post-intervention. | Vaccination rate improvement from 19% (6-26%) to 39% (9-55%). | 117 children attending hospital with asthma exacerbation. |
| Regan, 2014 (195) | Australia | 2013 | Cross-sectional survey of electronic outpatient records | NA | Assessment of proportion of people immunized and estimate of vaccine effectiveness. | 29.7% vaccination rate in asthmatics vs. 52.2% in those with coronary heart disease. In all patients, combined clinical and pathological data suggested vaccination had effectiveness of 56% in preventing infection: OR 0.44 (95% CI 0.24-0.79). | 2731 general practice patients with clinical and pathological data. |

ACT asthma control test; LMIC low-middle income countries; HIC high-income companies; COPD chronic obstructive pulmonary disease OR odds ration; CI confidence interval; NA not available;
## Section 4. Immunogenicity of influenza vaccines in asthma

| Authors, year | Country | Study period | Type of study | Intervention | Outcome measure | Key finding | Population characteristics | Limitations/comments |
|---------------|---------|--------------|---------------|--------------|-----------------|-------------|---------------------------|----------------------|
| Bae et al. 2013 (121) | Korea | 2011-2012 influenza season | Phase 4, multi-center, open-label study | TIV (H1N1, H3N2, B) (7.5 μg per strain) Two doses in vaccine naïve children; one dose in previously vaccinated interval between first and second dose not stated. | Short-term seroconversion rates, seroprotection rates, GMT, and GMT ratio; HI assay Safety: local and systemic adverse events. | Seroconversion and seroprotection rates in wheezers were similar to healthy children and significant for all 3 strains. The GMT was higher in the one dose-group vs. the two-dose group both in healthy children and in wheezers. The GMT ratio was significantly higher in healthy children than wheezers for the H1N1, H3N2 strains. No difference between steroid-treated and non-treated wheezers. | 68 healthy children and 62 recurrent wheezers aged 6 months to 3 years. | Chronic and high dose ICS excluded. No adjustment for ICS dose. Age significantly lower in healthy controls (1-2 y) than in wheezers (2-3 y). |
| Busse et al. 2011 (124) | US | 2009 influenza autumn season | Randomized, open-label study | Monovalent H1N1 tested at two doses (15 vs. 30 μg) twice at 21 days interval | Short term rates and GMT of seroconversion and seroprotection; HI assay and microneutralization. | The 2 dosing groups did not differ in seroconversion rates. In mild-to-moderate asthma the two doses offered equal seroprotection. In severe asthma seroprotection was significantly higher with the 30 μg dose. In severe asthmatics >60 years old the seroprotection was achieved only with the 30 μg dose The second dose provided no additional benefit. | Adolescents and adults (12-79) with asthma; special subgroup > 60 years with mild-to-moderate and severe asthma (including use of OCS). | No healthy controls for comparison. |
| Authors, year   | Country | Study period              | Type of study                              | Intervention                                                                 | Outcome measure                                                                 | Key finding                                                                                                                                                                                  | Population characteristics                           | Limitations/comments                                                                                     |
|----------------|---------|---------------------------|--------------------------------------------|------------------------------------------------------------------------------|--------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------|-----------------------------------------------------------------------------------------------------------|
| Pedrosa et al. 2009 (122) | Mexico | 2001-2002 influenza season | Double-blind, placebo controlled, randomized study | Inactivated TIV (H1N1, H3N2,B); dose 15 μg per strain; twice at 28 days interval | Safety: local and systemic adverse events; lung function; short-term rate of seroconversion; HI assay. | Significant rate of seroconversion after the first and second dose for all 3 strains.                                                                                                        | Children 5-9 years with mild intermittent and moderate asthma. | No healthy controls for comparison. Study funded by industry. Asthma severity assessed only as LF; no data on exacerbations. Subjects with a history of allergy to egg protein excluded. Previous influenza vaccination status not reported. |
| Zuccotti, et al. 2007 (123) | Italy  | 2005-2006 influenza season | Observational prospective open label multi-centre study | Virosomal-adjuvanted TIV (H1N1, H3N2,B); dose 15 μg per strain; single dose | Short (at one month) and long-term (at 6 months) seroconversion (≥1:40) and seroprotection (≥1:10); HI assay; both rate and fold-increase in GMT. Seroconversion and seroprotection in relation to pre-existing antibodies. | Seroconversion: Both rates and GMT significantly high at both 1 and 6 months for all 3 strains. Better GMT increase for the B strain in children 6-9 years compared to 3-5 years. Seroprotection: At one month satisfactory in both age groups, decreased | Children with asthma aged 3-9 years (subgroups 3-5 years and 6-9 years). | No data on asthma severity. No healthy controls. More than 50% of children had pre-existing antibodies at baseline. |
| Authors, year | Country | Study period | Type of study | Intervention | Outcome measure | Key finding | Population characteristics | Limitations/comments |
|--------------|---------|--------------|---------------|--------------|----------------|-------------|----------------------------|---------------------|
|              |         |              |               |              | Safety: local and systemic adverse events. | seroprotection for H3N2 for children aged 6-9 years. At 6 months: >80% in children aged 6-9 years; significantly lower (<70%) in children aged 3-5 years. Both seroconversion and seroprotection to any specific strain significantly lower without pre-existing antibodies. | Baseline GMTs were relatively high for all 3 strains. Subjects with history of egg allergy and/or severe atopy were excluded. | BMI body mass index; ICS inhaled corticosteroids; LF lung function; GMT geometric mean titer; HI hemagglutinin inhibition; OCS oral corticosteroids; SPT skin prick test; TIV trivalent influenza vaccines |
## Section 5. Efficacy of influenza vaccines in asthma

| Authors, year | Country | Study Period | Type of study | Intervention | Outcome measure | Key finding | Population characteristics | Limitations/ comments |
|---------------|---------|--------------|---------------|--------------|-----------------|-------------|----------------------------|----------------------|
| Abadoglu et al. 2004 (125) | Turkey | Recruitment: Sept - Nov 2001. Follow-up until Mar 2002 | Randomized, controlled trial | TIV vs. no vaccine | Frequency of URTI infections and asthma exacerbations during the winter following vaccination. | URTI and asthma exacerbation rates similar between vaccinated and non-vaccinated groups = no protective effect. | Asthmatic adults aged 22-84 years. In total: 128 Vaccination: 86 Nothing: 42. | Lack of knowledge on rate of influenza epidemic in tested period or on the cause of URTI. Relatively low number of subjects enrolled. |
| Bueving et al. 2004 (126) | The Netherlands | 1999-2000 2000-2001 | Double-blind, placebo controlled, randomized study | TIV vs. placebo | Symptom scores (to detect infection); pharyngeal swab and spirometry; quality of life (QoL). | Influenza vaccination did not result in a significant reduction of the number, severity, or duration of asthma exacerbations caused by influenza. | Asthmatic children aged 6-18 yrs. In total: 696 (296 in 1999 -2000 and 400 in 2000 - 2001) 347: influenza vaccine and 349 placebo. | Relatively few influenza-positive weeks of illness. |
| Bueving et al. 2004 (81) | The Netherlands | 1999-2000 2000-2001 | Double-blind, placebo controlled, randomized study | TIV vs. placebo | Symptom scores (to detect infection) Pharyngeal swab. Blood samples for influenza virus-specific antibodies (immunogenicity). | Influenza vaccination did not result in a significant reduction of the number, severity, or duration of asthma exacerbations caused by influenza. | 696 asthmatic children aged 618 years. |
| Fayyaz Jahani et al. 2012 (127) | Iran | ND | Randomized, placebo-controlled study | TIV | Asthma exacerbations. | Lower asthma exacerbation rates | 140 asthmatic children aged 6 to 60 months. | Conference abstract, thus |
| Authors, year | Country | Study Period | Type of study | Intervention | Outcome measure | Key finding | Population characteristics | Limitations/ comments |
|--------------|---------|--------------|---------------|--------------|----------------|-------------|---------------------------|----------------------|
| Fleming et al. 2006 (128) | Multicenter (UK & Europe) | Oct 2002 - May 2003 | Reactive-CT to examine relative efficacy of CAIV-T vs. TIV | CAIV-T vs. TIV | Surveillance for influenza-like symptoms. Nasal swab viral culture culture-confirmed influenza illness. Primary efficacy endpoint: incidence of culture-confirmed influenza illness caused by a community-acquired subtype antigenically similar to that in the vaccine. | CAIV-T had a significantly greater relative efficacy of 35% compared with TIV. There was no evidence of a significant increase in adverse pulmonary outcomes for CAIV-T compared with TIV. CAIV-T was well tolerated in children and adolescents with asthma. | Asthmatic children aged 6 to 17 years. CAIV-T (n=1114) TIV (n=1115). No placebo group, therefore the absolute efficacy of each vaccine cannot be calculated. | limited information. |
| Ambrose et al. 2012 (131) | | Meta-analysis of two randomized, multinational trials | CAIV-T vs. TIV | | Culture confirmed influenza illness during the influenza surveillance period. Rate of wheezing, hospitalization, adverse events. | Children receiving LAIV had fewer cases of culture-confirmed influenza illness than TIV recipients. | 1940 children aged 2-5 years with asthma or prior wheezing. | Post hoc definition of study cohorts, but this is in part addressed by the sub-population analysis. Identification of subjects asthma or history of wheezing relied on investigator judgment without validation. | |

TIV trivalent influenza vaccines; URTI upper respiratory tract infection; QoL quality of life; CAIV-T trivalent, cold-adapted influenza vaccine; LAIV live attenuated influenza vaccine
| Authors, year        | Country   | Study Period | Type of study                      | Intervention                        | Outcome measure                                      | Key finding                                                                 | Population characteristics                          | Limitations/ comments                                                                                                                                 |
|---------------------|-----------|--------------|------------------------------------|-------------------------------------|------------------------------------------------------|-----------------------------------------------------------------------------|--------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------|
| Christy et al. 2004 (137) | USA       | 1996-1997    | Retrospective nested case control study | Influenza vaccine                   | Asthma-related hospitalization, clinic and emergency department visits. | Adjusted OR (95% CI) associated with vaccine for asthma related: Hospitalization 1.9 (0.9-3.9). Clinic visit 2.9 (2.0-4.1). Emergency department visits 2.0 (1.2-3.1). | Primary care cohort. 400 randomly selected children with asthma who had been vaccinated, and 400 who had not. Age range 1-19 years. | Two groups were not balanced in terms of ethnicity and likelihood of receiving vaccine in previous year, but this was taken into account in adjusted analysis. Study suggests that vaccine associated with worse outcomes. Patients with more severe asthma or those more likely to present may be more likely to have been vaccinated although controlling for these did not alter the results. |
| Hak 2005 (136)      | Netherlands | 1999-2000    | Nested case control study          | Influenza vaccination               | All cause mortality, hospitalization, general practice visits for | High-risk children <18 years, 43% (10%-64%) of visits were prevented by | All age groups with long term medical | No separate analysis for asthma. |
| Authors, year | Country | Study Period | Type of study | Intervention | Outcome measure | Key finding | Population characteristics | Limitations/ comments |
|---------------|---------|--------------|---------------|--------------|----------------|-------------|---------------------------|----------------------|
| Herrera et al. 2007 (138) | USA | 2003-2004 | Case-control study | Influenza vaccine | Laboratory confirmed influenza report. | Influenza: vaccine “efficacy” was 60% (43-72%) and 48% (21-66%) among those without and with high-risk medical conditions. Influenza-related hospitalization: vaccine “efficacy” was 90% (68-97%) and 36% (0-63%) against for persons without and with high-risk conditions. | 330 cases, 1055 randomly recruited controls. Age range 50-64 years. | No separate data for asthma. Apparent effectiveness may be inflated by potential tendency to not test vaccinated people and miss an influenza diagnosis on a vaccinated patient. Authors say “efficacy” but study set up as an effectiveness study. Vaccine strains did not closely match |
| Authors, year | Country | Study Period | Type of study | Intervention | Outcome measure | Key finding | Population characteristics | Limitations/ comments |
|---------------|---------|--------------|--------------|--------------|-----------------|-------------|----------------------------|----------------------|
| Koshio et al. 2014 (132) | Japan | Sept-Oct 2010 | Survey | Vaccination against “new type of influenza”: H1N1 pdm09. | Report of a “new type of influenza”: H1N1 pdm09. | For influenza, OR (95% CI) for previous vaccination was 0.62 (0.42-0.90) for younger patients ≤61 years; among older patients OR was 1.38 (0.66-2.89). For infection-induced asthma exacerbation, OR (95% CI) for previous vaccination was 1.67 (0.60-4.66) for younger patients and 1.71 (0.50-5.83) for older ones. | 2555 adults with asthma. | There may have been other differences between vaccinated and non-vaccinated patients that explained the difference in risk of influenza. Influenza cases were not laboratory proven. This study looked at pandemic rather than ordinary influenza. |
| Kramarz et al. 2001 (133) | USA | 1993-1996 | Population based, retrospective cohort study | Influenza vaccine | Exacerbation of asthma. | Adjusted incidence rate ratios of asthma exacerbations after vaccination were 0.78 (0.55-1.10), 0.59 (0.43-0.81), and 0.65 (0.52-0.80) compared with the period before vaccination during the 3 influenza seasons. | Children aged 1-6 years with asthma in 4 large health maintenance organizations. (>70,000 patients). | As a database-based analysis, the asthma diagnostic labelling and recording of asthma exacerbations may not be completely accurate. |
| Authors, year     | Country  | Study Period | Type of study          | Intervention                  | Outcome measure                                                      | Key finding                                                                                                                                                                                                                                                                                                                                                       | Population characteristics | Limitations/ comments                                                                                                                                                                                                                     |
|-------------------|----------|--------------|------------------------|-------------------------------|----------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Ong et al. 2009 (135) | USA      |              | Survey                 | Influenza vaccine             | Exacerbation of asthma defined by oral corticosteroid, hospital visit or emergency department visit | In the multivariate analyses, current influenza vaccination status was independently associated with significantly decreased OR of using oral steroids in the previous 12 months (0.29, 0.10-0.84). There was no association with hospitalization (1.39; 0.340-5.67) nor emergency department visits.                                                                                                           | 80 children with asthma in a paediatric clinic. | Differences in patients receiving vaccination may have explained the difference in exacerbations. Adjusted for asthma severity.                                                                                                           |
| Smits et al. 2002 (134) | Netherlands | 1995-1997 | Retrospective cohort study | Inactive influenza vaccination | Physician diagnosed acute respiratory illness. | Vaccine effectiveness was 27% (-7 to 51%) after adjustments.                                                                                                                                             | Children with asthma aged 0-12 years in primary care. Over 300 patients vaccinated may have been different to                                                                                                                               |                                                                                                                                  |
| Authors, year | Country | Study Period | Type of study | Intervention | Outcome measure | Key finding | Population characteristics | Limitations/ comments |
|--------------|---------|--------------|---------------|--------------|-----------------|-------------|----------------------------|---------------------|
| Jaiwong et al. 2015 (196) | Thailand | June 2012 - Aug 2013 | Cross-sectional non-randomized study | TIV vs. no vaccine | Respiratory illnesses and asthma-related events. | The immunized group had significantly reduced: acute respiratory tract illnesses; asthma exacerbations; ER visits; bronchodilator usage and systemic steroid administrations; number of hospitalizations (p<0.001) and their duration (p <0.034). | 93 mild persistent asthmatic children aged 1-14 years (48 patients: 2 doses of inactivated influenza). | Non-randomized study. Small sample size. Lack of info related to other confounding factors in asthmatic exacerbations such as compliance and controller usage, other virus and bacterial infections, pollution, emotional conditions of the patient. |
| Yokoushi et al. 2014 (197) | Japan | May 2009 - Feb 2010 | Retrospective, cross-sectional, questionnaire-based survey | Inactivated, split-virus, non-adjuvanted | Vaccination rates. Vaccine effectiveness against physician-diagnosed influenza A infection, and | Comparison of positive influenza diagnosis rates between vaccinated and unvaccinated children with asthma. | 460 children with asthma (n=144) and without asthma (n=316). | Other environmental factors that may cause asthma |
| Authors, year | Country | Study Period | Type of study | Intervention | Outcome measure | Key finding | Population characteristics | Limitations/ comments |
|---------------|---------|--------------|---------------|--------------|-----------------|-------------|--------------------------|----------------------|
|               |         |              |               | monovalent vs. no vaccine | consecutive asthma exacerbations. | and without asthma showed that: Unvaccinated children with asthma had an elevated odds ratio (13.235; 95% CI: 5.564-32.134). Treatment for asthma exacerbations was needed in a larger proportion of unvaccinated children. Vaccine effectiveness against physician diagnosed influenza was 87% (95% CI, 78-93%) overall, 92% (95% CI, 81-96%) in children with asthma and 81% (95% CI, 63-91%) in children without asthma, respectively. | exacerbation were not considered. No data on atopic comorbidities. Retrospective study. Physician diagnosed influenza infection with or without a positive rapid antigen test result for influenza A during the pandemic period. |
### Section 7. Safety of influenza vaccines in asthma

| Authors, year       | Country | Study Period            | Type of study                                      | Intervention                                                                 | Outcome measure                                      | Key findings                                                                 | Population characteristics          | Limitations/ comments                                                                 |
|---------------------|---------|-------------------------|---------------------------------------------------|-----------------------------------------------------------------------------|------------------------------------------------------|-----------------------------------------------------------------------------|--------------------------------------|---------------------------------------------------------------------------------------|
| Greenhawt et al, 2012 (146) | USA     | Oct 2010 - Mar 2012     | Phase 1: randomized, controlled trial. Phase 2: retrospective, observational trial | Phase 1: Group A: graded 2 dose TIV challenge Group B: sham (saline) 1st dose followed by full TIV dose Phase 2: participants who declined study and received TIV from their allergist or had already received TIV in primary care. | Safety of TIV in egg-allergic children with severe allergy. | No allergic reactions were recorded. | Egg-allergic children with severe allergy (mean age 12 years). All patients had previous confirmed severe allergic reactions to egg. |
| Howe et al. 2011 (147) | USA     | Phase 1: Oct 2004-Feb 2009 Phase 2: 2009-2010 | Phase 1: Retrospective, observational study. Phase 2: Prospective observational study. | TIV | Safety of TIV in egg-allergic and non-allergic patients. | No severe allergic reactions have been recorded. | Children |
| Miller et al. 2003 (143) | USA     | Double-blind, placebo controlled trial. | Influenza vaccine | T-cell immune response to vaccine components in patients with asthma who experience bronchospasm. | Increased influenza (but not egg) antigen-induced mononuclear cell proliferation, unrelated to asthma exacerbations after vaccine. No differences in cytokine production in response to either influenza or egg antigen in | Adults, mean age 43 years. |

- **TIV**: Trivalent inactivated vaccine.
| Authors, year          | Country   | Study Period | Type of study                     | Intervention                        | Outcome measure                                                                 | Key findings                                                                                           | Population characteristics | Limitations/ comments |
|-----------------------|-----------|--------------|-----------------------------------|-------------------------------------|---------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------|-----------------------------|-----------------------|
| Chung et al. 2010 (144) | USA       | 2002-2009    | Retrospective, observational study | Influenza vaccine                  | Tolerability of the vaccine, defined as the lack of localized or systemic adverse reactions, in egg-allergic and non-allergic patients. | Association with asthma exacerbations. Bronchospasm after immunization is not related to vaccination. | 95% of egg-allergic patients who received the vaccine skin test tolerated the influenza vaccine without any serious adverse reactions. | Children, 6 months - 18 years. |
| Gagnon et al. 2010 (145) | Canada    | Oct-Dec 2009 | Prospective, observational study  | Adjuvanted monovalent 2009 pandemic influenza A/H1N1 influenza vaccine | Risk of anaphylaxis in children with egg allergy administered an adjuvanted monovalent 2009 pandemic influenza A/H1N1 influenza vaccine. | No patient had an anaphylactic reaction. | No patient had an anaphylactic reaction. | Children and adults. |
| Greenhawt et al. 2010 (149) | USA       | Oct 2009-Feb 2010 | Controlled, prospective trial | H1N1 influenza vaccine            | Safety of administering H1N1 to egg-allergic patients compared with non-egg-allergic, paediatric, control patients. | No significant allergic reactions were recorded. | No significant allergic reactions were recorded. | Children and adults aged 6 months to 24 years. |
| Esposito et al. 2008 (148) | Italy     | Nov-Dec 2007 | Prospective study                | Virosomal adjuvanted influenza vaccine | To evaluate whether the virosomal adjuvanted influenza vaccine that has been shown to have the lowest egg protein content could be administered to children with even severe egg | Administration of the whole vaccine was safe and well tolerated by all of the children, with no differences between those with and without egg allergy, or between those with egg allergy of different severity. | Administration of the whole vaccine was safe and well tolerated by all of the children, with no differences between those with and without egg allergy, or between those with egg allergy of different severity. | Children >3 years. |
| Authors, year       | Country | Study Period  | Type of study                                                                 | Intervention          | Outcome measure                                                                 | Key findings                                                                                                                                                                                                 | Population characteristics | Limitations/ comments |
|---------------------|---------|---------------|--------------------------------------------------------------------------------|-----------------------|---------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------|-----------------------|
| Castro et al. 2001 (140) | USA     | Sep-Nov 2000  | Multicenter, randomized, double blind, placebo-controlled, crossover trial | TIV or placebo        | Safety of TIV in adults and children with asthma.                                | The frequency of asthma exacerbations was similar in the two weeks after the TIV or placebo injection. The exacerbation rates were similar in subgroups defined according to age, severity of asthma, and other factors. | Children and adults aged 3-64 years.                                                                                     |                         |
| Baxter et al. 2012 (141) | USA     | Oct 2003 - March 2008 | Prospective observational post-marketing study | LAIV, TIV or placebo. | Safety of LAIV, compared to TIV and placebo in children.                        | Asthma/wheezing major adverse events were not statistically increased in LAIV recipients. No anaphylaxis events occurred.                                                                                       | Children aged 5-17 years.                                                                                                   |                         |
| Ambrose et al. 2012 (131) | USA     | Study 1: 2004-2005 Study 2: 2002-2003 | Post-hoc analysis of two randomized, multinational trials of LAIV and TIV in children aged 6 - 71 months | LAIV vs. TIV          | Safety of LAIV in asthmatic children younger than 6 years.                     | Wheezing, lower respiratory illness, and hospitalization were not significantly increased in children receiving LAIV compared with TIV. Increased upper respiratory symptoms and irritability were observed in LAIV recipients. | Children aged 2-5 years.                                                                                                    |                         |
| Gaglani et al. 2008 (142) | USA     | 1998-2002     | Open-label field trial                                                          | Trivalent LAIV        | Safety of the intranasal LAIV3 in children with intermittend wheezing/asthma. | No increased risk for medically-attended acute respiratory illnesses, including asthma exacerbation.                                                                                                       | Children aged 1.5-18 years.                                                                                                  |                         |
| Authors, year | Country | Study Period | Type of study | Intervention | Outcome measure | Key findings | Population characteristics | Limitations/ comments |
|--------------|---------|--------------|---------------|--------------|----------------|--------------|---------------------------|---------------------|
| Haber et al. 2015 (198) | USA | LAIV reports: 2013-2014 LAIV3 reports: 2010-2013 | Analysis of reports received at the "Vaccine Adverse Event Reporting System" | LAIV4 vs. LAIV3 | Medical records were reviewed for non-manufacturer serious reports (i.e., death, hospitalization, prolonged hospitalization, life-threatening illness, permanent disability) and reports of selected conditions of interest. | No new or unexpected safety concerns with LAIV4. The safety profile of LAIV4 was similar to LAIV3. Just over 94% of LAIV4 reports were non-serious and the most common adverse events were mild, self-limited conditions (e.g., fever, headache and cough). | Children and adults aged 2-49 years. 12.7 million doses of LAIV4 were distributed. | General limitations of passive surveillance. |
| Turner et al. 2015 (151) | UK | Sept 2013 - Jan 2014 | Prospective, multi-centre, open-label, phase IV intervention study | LAIV | Clinical symptoms after LAIV administration to estimate incidence of 1. Allergic reaction within 2 hours, 2. Delayed allergic symptoms within 72 hours, 3. Non-allergic adverse events; and change in nasal airway patency (acoustic rhinometry). | LAIV appeared to be safe in children with egg allergy and additional asthma or recurrent wheeze. No systemic allergic reactions reported. Eight children had mild, self-limiting, potentially IgE-mediated allergic symptoms. 26 (9.4%; 95% CI for population, 6.2% to 13.4%) children experienced lower respiratory tract symptoms within 72 hours, including 13 with parent-reported wheeze. None of these episodes required medical intervention beyond routine treatment. | 282 children with egg allergy aged 2-17 years. | The egg protein content of LAIV batches may vary, thus data from this study may not be applicable to future LAIV stocks in which egg content may be higher. |

TIV trivalent influenza vaccine; LAIV live attenuated influenza vaccine; CI confidence interval
### Section 8. Antiviral treatment against influenza in patients with asthma

| Authors, year | Country | Study Period | Type of study | Intervention | Outcome measure | Key finding | Population characteristics | Limitations/ comments |
|---------------|---------|--------------|---------------|--------------|-----------------|-------------|---------------------------|---------------------|
| Johnston et al. 2005 (168) |         |              | Randomized study | Oseltamivir (2 mg/kg) or placebo twice daily, as a syrup | Effects of oseltamivir among influenza-infected children with asthma. | Oseltamivir is safe and well tolerated among asthmatic children, may reduce symptom duration and helps improve lung function and reduce asthma exacerbations during influenza infection. | Asthmatic children aged 6-12 years. Analysis was performed for both the intent-to-treat infected (n=179) and per protocol (n=162) populations. |                      |
| Lin et al.2006 (162) | China | 2002-2003 | Randomized, open-label, controlled trial | Oseltamivir 7.5 mg twice daily for 5 days (oseltamivir group), or symptomatic treatment (control group) within 48 h after symptom onset | Duration and severity of illness in influenza-infected patients, incidence of complications, antibiotic use, hospitalization and total medical cost. | Oseltamivir is effective and well tolerated in high-risk patients with chronic respiratory or cardiac diseases. It can reduce the duration and severity of influenza symptoms and decrease the incidence of secondary complications and antibiotic use, without increasing the total medical cost. | 118 Chinese patients (mean age 48.1 years) with chronic respiratory diseases (chronic bronchitis, obstructive emphysema, bronchial asthma or bronchiectasis) or chronic cardiac disease. | Only 8 patients with bronchial asthma. |
| Cass et al. 2000 (166) | UK | 2000 | Double-blind, randomized, placebo controlled, two way cross-over study, 7 day wash out period. | 10 mg zanamivir as a dry powder (265 mg) or a placebo, twice daily on day 1 and then four times daily from day 2 to day 14. | Safety of Zanamivir in asthmatics. | Zanamivir inhaled as a dry powder does not significantly affect the pulmonary function and airway responsiveness of subjects with mild/moderate asthma and therefore its use in such patients/subjects is not precluded. | 13 subjects, aged 19-49 years, with asthma. |                      |
| Authors, year | Country | Study Period | Type of study | Intervention | Outcome measure | Key finding | Population characteristics | Limitations/ comments |
|---------------|---------|--------------|---------------|--------------|-----------------|-------------|---------------------------|----------------------|
| Shun-Shin et al. 2009 (169) | | | Systematic review and meta-analysis of randomized controlled trials | NA | Effects of the neuraminidase inhibitors oseltamivir and zanamivir in treatment of children with seasonal influenza and prevention of transmission to children in households. | Oseltamivir did not reduce asthma exacerbations or improve peak flow in children with asthma. | Children aged ≤12 years in the community (not admitted to hospital) with confirmed or clinically suspected influenza. | Based on only one trial. |
| Jefferson et al. 2014 (170) | | | Systematic review | NA | Potential benefit and harm of oseltamivir by reviewing all clinical study reports (or similar document when no clinical study report exists) of randomized placebo controlled trials and regulatory comments. | Oseltamivir had no effect in children with asthma, but there was an effect in otherwise healthy children. | Children and adults who either were healthy before exposure to respiratory agents or had a chronic illness (such as asthma, diabetes, hypertension) but excluding people with immune suppression. | |
| Jefferson et al. 2014 (171) | | | Cochrane review | NA | Potential benefit and harm of neuraminidase inhibitors for influenza in all age groups by reviewing all clinical study reports of published and unpublished randomized, placebo-controlled trials and regulatory comments. | For the treatment of adults, oseltamivir reduced the time to first alleviation of symptoms by 16.8 hours (95% CI 8.4-25.1, p<0.0001). This represents a reduction in the time to first alleviation of symptoms from 7 to 6.3 days. There was no effect in asthmatic children, but in otherwise healthy children there was a reduction by | All age groups. 53 trials in Stage 1 and 46 in Stage 2 (formal analysis), including 20 oseltamivir (9623 participants) and 26 zanamivir trials (14,628 participants). |
| Authors, year | Country | Study Period | Type of study | Intervention | Outcome measure | Key finding | Population characteristics | Limitations/comments |
|---------------|---------|--------------|---------------|--------------|----------------|-------------|---------------------------|----------------------|
| Watanabe, 2013 (173) | Japan | 2009 | Double-blind, randomized controlled trial | Laninamivir group: single inhalation of 40 mg laninamivir and placebo oseltamivir capsules twice daily for 5 days. Oseltamivir group: one capsule (75 mg) of oseltamivir orally twice daily for 5 days and laninamivir placebo powder once on day 1. | Efficacy and safety of laninamivir octanoate, an inhaled neuraminidase inhibitor for the treatment of influenza patients with chronic respiratory diseases. | Laninamivir octanoate showed similar efficacy and safety to oseltamivir in the treatment of influenza, including that caused by influenza A(H1N1)2009, in patients with chronic respiratory diseases. | 203 patients aged ≥20 years. Most patients had underlying bronchial asthma. |
| Kohno 2011 (172) | Japan | Jan-May 2009 | Multicenter, uncontrolled, randomized, double-blind study | Peramivir intravenously administered at 300 or 600 mg/day for 1 to 5 days, as needed | Efficacy and safety of the novel anti-influenza virus drug peramivir in high-risk patients. | In the 600 mg group about 50% shorter duration of illness. In 33% of patients adverse drug reactions, described as “not particularly problematic clinically” with quick recovery in all cases. | 37 influenza patients, ≥20 years, with risk factors, including asthma. 5 patients with asthma were also taking steroids. |
| Biggerstaff et al. 2014 (199) | USA | 2009-2010 | USA state-based, random-digit-dialled | none | Influenza antiviral treatment prescription in influenza like illness (ILI). | 26% of respondents with ILI received clinical influenza diagnosis. Of these 36% received | 216,431 adults ≥18 years, non-institutionalized Data in this study are self-reported and subject to |
| Authors, year | Country | Study Period | Type of study | Intervention | Outcome measure | Key finding | Population characteristics | Limitations/ comments |
|---------------|---------|--------------|---------------|--------------|-----------------|-------------|----------------------------|----------------------|
| Biggerstaff et al. 2014 (165) | USA | 2010-2011 | USA state-based, random-digit-dialled telephone survey | none | Influenza antiviral treatment prescription in influenza like illness (ILI). | Among adults with a physician diagnosis of influenza, 34% received antiviral treatment. Having asthma was not associated with antiviral treatment. | 75,088 adults (18- >65 years) and 15,649 children (0-17 years) were interviewed. | recall and social desirability bias. Diagnosis and treatment are not verified. |

COPD chronic obstructive pulmonary disease; NAI neuraminidase inhibitor; CI confidence interval.
Figure 1. Global seasonal influenza vaccination policies for people with asthma. Source data obtained from following references: Europe\textsuperscript{1,2}; Americas\textsuperscript{3,4}; Africa\textsuperscript{5,6}; Asia\textsuperscript{2,7-9}; Oceania\textsuperscript{7}. The following countries have vaccination policies for chronic disease, which includes asthma in the majority of cases: Argentina, Bahamas, Belize, Bolivia, Brazil, Chile, Colombia, Costa Rica, Cuba, Dominican Republic, Ecuador, Egypt, El Salvador, Fiji, Grenada, Ivory Coast, Mauritius, Mexico, Morocco, Panama, Paraguay, Peru, Philippines, Saint Lucia, Thailand, Trinidad and Tobago, Uruguay and Venezuela. This graphic is licenced by a Creative Commons Attribution-ShareAlike 3.0 Unported License: [http://gunn.co.nz/map](http://gunn.co.nz/map).

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Appendix 1

Search strategy: Medline format

1. Respiratory Hypersensitivity/ or Bronchial Hyperreactivity or Asthma/ or Status Asthmaticus/
2. (asthma or asthma exacerbation or asthma attack or acute asthma or acute asthmatic attack or asthmatic children or wheez*).mp.
3. (bronchial disorder or airway reactivity or airway hyperresponsiveness or bronchial hyperreactivity or bronchial hyperresponsiveness or lung function or ventilatory function or FEV or FEF or FVC or PEF).mp.
4. or/1-3
5. Influenza, Human/ or Influenza vaccines/ or Influenza A virus or Influenza B virus or Influenzavirus C/ or Influenza A Virus, H5N1 Subtype/ or Influenza A Virus, H1N1 Subtype/ or Influenza A Virus, H3N2 Subtype/
6. exp Epidemiologic Studies/ or exp Intervention Studies/ or exp Evaluation Studies/ or exp Comparative Studies/ or exp Follow-up Studies/ or exp Prospective Studies/ or exp Cohort Studies/ or exp Case-Control Studies/ or exp Cross-Sectional Studies/
7. (prospectiv* or cohort or birth cohort or analytical stud* or cohort stud* or case-control stud* or cross-sectional stud*).mp.
8. Review literature as Topic/ or systematic review.mp.
9. Clinical trial/ or Controlled Clinical Trial/ or Randomized Controlled Trial/
10. exp Placebos/ or exp Random Allocation/ or exp Double-Blind Method/ or exp Single-Blind Method/
11. (trial or clinical trial or controlled clinical trial or random* or randomi?ed controlled trial or double-blind design or single blind design).mp.
12. exp Survey/ or survey.mp.
13. or/6-12
14. 4 AND 5 AND 13
15. limit 14 to yr="2000-2014"
16. limit 15 to human

Search Strategy: free-field format

(Asthma or acute asthmatic attack or status asthmaticus or wheez* or respiratory hypersensitivity or bronchial hyperreactivity or bronchial hyperresponsiveness or airway hyperreactivity or airway hyperresponsiveness)
AND
(Influenza, human or influenza vaccin* or influenza A virus or influenza B virus or influenzavirus C or influenza A virus subtype H1N1 or Influenza A virus subtype H3N2 or Influenza A virus subtype H5N1)
AND
(Analytical stud* or epidemiologic* or compar* or evaluat* or follow-up or observation* or cohort stud* or case-control stud* or cross-sectional stud* or systematic review or intervention* or trial* or RCT or double-blind or single-blind or control* or random* or prospective or retrospective)