Vaccination and allergy:  
EAACI position paper, practical aspects

Lennart Nilsson¹, Knut Brockow², Johan Alm³, Victoria Cardona⁴, Jean-Christoph Caubet⁵, Eva Gomes⁶, Maria C. Jenmalm⁷, Susanne Lau⁸, Eva Netterlid⁹, Jürgen Schwarze¹⁰, Aziz Sheikh¹¹, Jann Storsaeter¹², Chrysanthi Skevaki¹³, Ingrid Terreehorst¹⁴, Giovanna Zanoni¹⁵

¹Allergy Center, University Hospital, Linköping, Sweden; ²Department of Dermatology and Allergy Biederstein, Technical University Munich, Germany; ³Sachs’ Children and Youth Hospital, Stockholm, Sweden; ⁴Allergy Section, Department of Internal Medicine, Hospital Universitari Vall d’Hebron, Barcelona, Spain; ⁵University of Geneva, Division of Paediatrics, Genève, GE, Switzerland; ⁶CHP, Porto, Portugal; ⁷Unit of Autoimmunity and Immune Regulation, Department of Clinical and Experimental Medicine, Linköping University, Linköping, Sweden; ⁸Pediatric Pneumology and Immunology, Charité Universitätsmedizin, Berlin, Germany; ⁹Department of Occupational and Environmental Dermatology, Lund University, Malmö, Sweden and The Public Health Agency of Sweden; ¹⁰Child Life & Health and MRC-Centre for Inflammation Research, The University of Edinburgh, Edinburgh, UK; ¹¹Asthma UK Centre for Applied Research, Usher Institute of Population Health Sciences and Informatics. The University of Edinburgh, UK; ¹²Norwegian Institute of Public Health; ¹³Institute of Laboratory Medicine and Pathobiochemistry, Molecular Diagnostics, Philipps University Marburg, University Hospital Giessen and Marburg GmbH, Marburg, Germany, ¹⁴AMC, Dept ENT, the Netherlands; ¹⁵Immunology Unit, University Hospital, Verona, Italy

Dedicated to Christoph Grüber and Isil Barlan

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Abstract
Immunization is highly effective in preventing infectious diseases and therefore an indispensable public health measure. Allergic patients deserve access to the same publicly recommended immunizations as nonallergic patients unless risks associated with vaccination outweigh the gains.
Whereas the number of reported possible allergic reactions to vaccines is high, confirmed vaccine-triggered allergic reactions are rare. Anaphylaxis following vaccination is rare, affecting less than 1/100,000, but can occur in any patient. Some patient groups, notably those with a previous allergic reaction to a vaccine or its components, are at heightened risk of allergic reaction and require special precautions. Allergic reactions, however, may occur in patients without known risk factors and cannot be predicted by currently available tools. Unwarranted fear and uncertainty can result in incomplete vaccination coverage for children and adults with or without allergy.
In addition to concerns about an allergic reaction to the vaccine itself, there is fear that routine childhood immunization may promote the development of allergic sensitization and disease. Thus, although there is no evidence that routine childhood immunization increases the risk of allergy development, such risks need to be discussed.

Aims
This position paper provides expert advice on how to prevent and manage allergic reactions to vaccines against infectious diseases, and immunization in relation to the development of allergic diseases. Because systemic reactions can cause greater harm than local reactions, this paper focuses on the former.

Methods
Evidence and recommendations provided are based on currently available published data. In January 2013, articles in English, German, and Italian with data on hypersensitivity reactions to vaccines were identified by searching the Medline (National Library of Medicine) database. Additional articles were found through the reference lists of the identified articles, textbooks, publications of national registries or organizations, existing guideline articles, and a Medline search update covering January 2013-September 2016. Relevant articles were identified on the basis of title and abstract, retrieved and analysed. Evidence was discussed, and statements were adopted or amended by consensus among the authors.
I. Basic information

I A. Allergic reactions to vaccines.

**Statement:** Allergic reactions to vaccines are rare, mostly directed to additives. Knowledge of all ingredients is of importance when vaccinating an allergic individual.

Documented allergic reactions have been reported for all vaccines but account only for a minority of all adverse events following immunization (AEFI, abbreviations; see also Table 1). In addition to microbial antigens, vaccines may include stabilizers, adjuvants, preservatives, and residual contaminants from the production process. (1, 2, http://www.vaccinesafety.edu/components.htm and http://www.cdc.gov/vaccines/pubs/pinkbook/appendix/index.html). Although microbial antigens rarely cause allergic reactions, they have been described in recent papers for anaphylaxis associated with influenza vaccine and for a mutant diphtheria toxin (CRM197) in pneumococcal conjugated vaccine (PCV) (3, 4). Knowledge of all the ingredients in a vaccine is crucial to identifying the culprit allergen. The principal allergens in vaccines are listed below.

**Gelatine**, a vaccine stabilizer of bovine or porcine origin, has been reported to be responsible for anaphylaxis to some brands of measles, mumps, and rubella (MMR) and varicella vaccines, and also earlier in Japanese encephalitis and influenza vaccines.

Residual ovalbumin from **hen's egg** can be present in yellow fever (YF), influenza, MMR, tick-borne encephalitis (TBE) and some rabies vaccines in various concentrations (Figure 1). **Chicken protein** in YF vaccine has been reported to be a potential severe problem in chicken-allergic recipients. Very low concentration of **cow's milk** proteins may be present in some brands of diphtheria, tetanus and pertussis (DTP) vaccines, and oral polio vaccine (OPV) (5).

**Thiomersal, aluminium, and phenoxyethanol** can cause local reactions (mostly delayed-type hypersensitivity such as contact allergy and maculopapular rash), but have not been reported as a cause of proven anaphylaxis. Nowadays, thiomersal is rarely used as a preservative in vaccines, and its clinical importance as an allergen is doubtful (6). Local reactions can nevertheless be more frequent among sensitized recipients (7).

**Formaldehyde** is still used in vaccine preparation (8), but no IgE-mediated reactions to formaldehyde have been recently described.

Trace amounts of **antimicrobials** could theoretically cause anaphylaxis in sensitized patients; however, few reports are found in the literature. Although the association of neomycin sensitization and IgE-mediated allergic reactions to vaccines is poorly supported by the literature, a history of anaphylaxis to neomycin is considered a contraindication for immunization with vaccines containing neomycin (9). Contact dermatitis with neomycin is more frequent (9).
Vaccine vial stoppers or syringe plungers may contain natural latex rubber and pose a theoretical risk to latex-allergic patients (10). Incidence is, however low; only one report of an anaphylactic reaction in a latex-allergic patient was attributed to rubber in the stopper (11) of an Hepatitis B (HB) vaccine. Human papillomavirus vaccines (HPV) may contain residual yeast protein (Saccharomyces cerevisiae) from the production process. Rarely, an immediate reaction can happen after vaccination in yeast-allergic patients (12). Yeast is also used in the production of the carrier CRM197, and could theoretically be contained in PCV-13 and some meningococcal and oral typhoid vaccines. (1). Dextran has been implicated in allergic reactions to some vaccines that have been withdrawn from the market (1). Alpha-gal anaphylaxis minutes after immunisation with zoster vaccine (OKA VZV) has recently been suggested in a patient with a documented history of red meat allergy. It has been postulated that the patient has reacted to alpha-gal from porcine gelatin or bovine calf serum in the vaccine (13).

I B. Immune response to vaccines in relation to allergy

Statement: Determination of vaccine antigen-specific IgE is not recommended in the work-up of allergic reactions to vaccines, because IgE production can be part of the normal vaccine immune response and it is mainly not commercially available.

Specific IgE response to vaccine antigens can frequently be observed alongside IgG responses (14). After primary immunization, about 50% of infants have detectable IgE against D and T toxoids [14]; after booster, more than 90% of vaccines have detectable IgE against the vaccine antigens [15]. The IgE response to vaccine antigens, mediated by a Th2-type immune response, seems more pronounced among atopic individuals [14]. It has therefore been hypothesized that immunization of atopic children may be associated with clinical vaccine allergy. However, no relevant clinical allergic reaction to microbial antigens in vaccines has been reported before two recent papers, see 1A (3, 4). In young children, Th1-/IFN-associated and Th2-associated gene networks coexist in an apparent state of dynamic equilibrium, but atopic individuals have Th2-dominant allergen-specific responses, and their Th1/IFN networks are disrupted and down-regulated (16). Therefore, the optimal immunogenicity/reactivity balance of new vaccines will have to be specifically defined in this population.

I C. Systemic and local reactions

Statement: Anaphylaxis following vaccination is rare and has to be distinguished from vasovagal reaction. Local reactions are common and mainly due to non-allergic immune reaction.

Classification of hypersensitivity reactions to vaccines is challenging as the underlying mechanisms are poorly understood and no consensus exists in the literature. Several classifications have been proposed, based on the
extent, severity and timing of the reaction (17). In this paper, reactions after vaccination are categorized as systemic and local reactions according to WHO (18).

**Systemic reactions**

Among AEFI, systemic severe allergic reactions are rare but important. Anaphylaxis is an acute severe, potentially life-threatening emergency (19), Table 2. Symptoms usually start within the first hour after immunization (17). Reactions occurring more than two hours after exposure have been described, but are uncommon, and the causal relationship is unclear (20). The incidence of anaphylactic reactions to certain vaccines is listed in Table 3. In typical cases with multi-organ involvement and objectively measurable signs in the four organ systems (skin, gastrointestinal tract, respiratory tract and cardiovascular system), diagnosis can be easy and certain. In other cases, diagnosis may be difficult, and anaphylaxis has to be differentiated from vasovagal reaction after immunization, Table 4.

Anaphylactic reactions can be IgE-mediated or non-IgE-mediated; these can be difficult to differentiate clinically.

Non-allergic systemic reactions should be distinguished from systemic IgE-mediated reactions. Fever and nonspecific systemic symptoms, such as skin rash, irritability, malaise, diarrhoea, headache, muscle pains and syncope are the most common systemic events after vaccination. Skin rashes, delayed urticaria and/or angioedema or maculopapular skin rash often occur a few hours after vaccine administration. Nonspecific activation of the immune system and nonspecific degranulation of mast cells may be the cause (21).

**Local reactions**

Local reactions include pain, redness and/or swelling at injection site. Mild local reactions are attributed to nonspecific inflammation due to the injection itself and injection of foreign materials. Large local reactions are less common and usually occur within 24-72 hours after vaccine administration. However, after a fifth dose of DTaP vaccine in 4-5 year-olds, about 1/4 of the children will get a large local reaction, usually well tolerated and resolving within 1-2 weeks (22). Typical large local reactions and chronic subcutaneous nodules with itching and eczema are considered type IV reactions. Local reactions could also be Arthus type, i.e. type III hypersensitivity. For these, the administration technique is important; deeper injection is associated with a lower rate of local reactions, especially in children younger than 3 years (23). Injection in the arm is associated with higher incidence of reactions than injection in the thigh (24). Traces of antibiotics, thiomersal and formaldehyde can contribute to local reactions. The incidence of local reactions for certain vaccines is shown in Table 5.
ID. Possible development of allergy by immunization

**Statement:** Routine childhood immunization does not promote the development of allergic sensitization to common inhalant or food allergens or the development of allergic disease.

Immunizations have been widely suspected of promoting the development of allergies, with related concerns contributing to delayed or incomplete immunization (25).

Epidemiological studies have addressed a possible effect of immunization on allergy development in general. However, immunizations had no effect on allergic disease in several studies (26, 27). Higher cumulative vaccine antigen doses were associated with less allergic sensitization, allergic disease (28) and less severe infant eczema (29). In concordance, regional immunization rates were inversely associated with allergic disease (30). Pertussis immunization has been suspected as pro-allergic because P toxin, included in cellular and acellular vaccines, can enhance IgE formation. However, data from a randomized intervention trial failed to show an increased risk of allergic sensitization or allergic disease up to 7 years of age (31). In a large ecological study, there was no increased risk of requiring asthma medication in adolescents whether they had had P vaccination in infancy or not (32).

Lower rates of allergic symptoms and allergic sensitization have been found among children with measles, but no association was found between measles vaccination and allergic symptoms (33). DT immunization was associated with asthma in one study (34), but not in others. Importantly, several further studies could not find any effect of MMR (28, 35), *Haemophilus influenzae* type b (36) or DTP (27) vaccinations on allergic sensitization or allergic disease. Mycobacterial lipoproteins elicit particularly strong Th1 responses. Consequently, it has been suggested that BCG vaccine administered in infancy might protect against the development of Th2-mediated allergic disease. A systematic review and meta-analysis (37) suggested that BCG vaccination is unlikely to be effective in preventing allergic sensitization or eczema, but might offer transient benefits against developing asthma.

**Topic II Specific vaccines and adverse events**

**Topics II A. Diphtheria, tetanus, pertussis vaccines**

True allergic or immediate hypersensitivity reactions to routine vaccines are rare, estimated as 2 per million doses for DTaP (20). In Japan (1994-2004) the total incidence of anaphylaxis was 0.95 per million doses of DTaP, but the authors were unable to identify a causal relationship to any vaccine component (38). Neither skin prick tests (SPT) nor specific IgE analyses could predict these reactions.

Specific IgE antibodies to D, T and P vaccines are common after booster doses if primary vaccination was with an acellular P vaccine; this response was exaggerated in atopic children with clinical manifestations (39).
Elevated P toxin IgE levels are associated with local reactions (40). As the adjuvant effect of aluminium on IgE production is well known, controversy exists regarding the extent to which the toxoids cause the local reactions (22).

Casein, a cow’s milk protein, has been implicated as a cause of anaphylaxis to DTP-containing vaccines in children with severe milk allergy and high specific milk IgE levels (41). Whereas these data need to be confirmed, trace amounts of casein have been demonstrated in some brands of DTaP or dTaP-containing vaccines prepared in a medium derived from cow’s milk protein. However, it is important to recognize that most patients with even severe milk allergy tolerate childhood vaccines, so no changes to vaccine recommendations have resulted from these case reports (42).

II B. Influenza vaccination

Vaccines for influenza prevention include the trivalent and quadrivalent inactivated influenza vaccines (IIVs), recombinant subunit vaccine (RIV), and live attenuated three and quadrivalent influenza vaccines (LAIVs).

According to the Centers for Disease Control and Prevention (CDC) and WHO, individuals from six months of age should be vaccinated against seasonal influenza [http://www.cdc.gov/mmwr/volumes/65/rr/rr6505a1.htm?s_cid=rr6505a1_w; August 26, 2016]. IIVs have generally been found to be safe for adults and children with asthma [43, 44], including those with severe disease [44]. Medically significant wheezing was increased in children 6–23 months of age who had received LAIVs but not in children 2–5 years of age [45]. Moreover, a recent Cochrane review did not show any significant increase in acute asthma exacerbations immediately following IIVs in adults or children older than 3 years of age [46]. In addition, data support the safety and efficacy of LAIVs among children aged 2-17 years with mild to moderate asthma or with a history of wheezing [47], but data regarding individuals with severe asthma/active wheezing are limited.

Recent studies provide robust evidence that IIVs with low ovalbumin content (< 0.12 µg/mL), can be administered safely in egg allergic patients, even in those with severe reactions (48 - 50). Data regarding the safety of LAIVs in egg allergy is emerging. The upper ovalbumin content of LAIVs is, reported on the package insert, 0.24 µg per 0.2 mL dose, but independent laboratories found it to be very low, between 0.00013 and 0.0017 µg per 0.2 mL dose (17). The ovalbumin content is published prior to the influenza season each year (https://www.gov.uk/government/collections/vaccine-update). The recent SNIFFLE studies combined found no systemic vaccine reactions and only 17 (1.6%) mild self-limiting reactions in 1,242 LAIV doses given to 1,061 egg-allergic children, including 335 with previous anaphylaxis to egg (49, 50). Based on these results, UK immunization recommendations no longer consider egg allergy a contraindication to LAIV, unless a child has had life-threatening anaphylaxis requiring intensive care treatment (51).
II C. MMR vaccine

MMR vaccination has been considered a problem in egg-allergic children because the attenuated viruses are cultured in hen’s embryonic fibroblasts, and the vaccines could contain traces of ovalbumin. However, several studies revealed that MMR vaccination is safe in infants and children with egg allergy (52). There are, however, reports of allergic reactions to gelatine (53).

Recent data confirm that infants and children allergic to hen’s egg can be vaccinated in GP settings and do not have to be referred to specialized centres. A review of the Irish paediatric emergency department vaccination programme for patients at risk of allergy/anaphylaxis analysed the clinical outcome of 374 children referred due to a history of allergy or anaphylaxis after 446 vaccine doses, including 310 (69.5%) MMR doses, were administered to 374 patients. Only six patients (1.3%) experienced a minor immediate reaction to a vaccination (54). In the Danish Childhood Vaccination Programme, 32 patients with sensitization to hen’s egg displayed no reaction to MMR vaccine (Priorix®) (55).

The British Society for Allergy and Clinical Immunology (BSACI) guidelines for the management of egg allergy recommend that children with egg allergy should receive routine MMR vaccination in primary care (56).

II D. Pneumococcal and meningococcal vaccines

There are no contraindications to pneumococcal or meningococcal vaccines for patients with allergy except for those with other known hypersensitivity to vaccine components including D (or CRM 197) or T toxoids present as carriers in conjugated vaccines, or previous severe reaction to the vaccine.

II E. BCG vaccine

Most adverse reactions after BCG vaccination are infectious. Hypersensitivity reactions are mostly mild injection site reactions and lymphadenitis, whereas systemic reactions, such as the immune reconstitution inflammatory syndrome, are rare (57).

II F. Polio vaccination

A theoretical risk of hypersensitivity reactions exists due to trace amounts of streptomycin, neomycin and polymyxin B in both injectable and oral polio vaccine. The latter may also contain cow’s milk proteins (5) (see I A). Confirmed anaphylaxis is extremely rare. Data from the UK, Canada and the US indicate rates of 0.65-3 anaphylaxis events per million doses of vaccine administered (58).

II G. Hepatitis B vaccination

Hepatitis B (HB) vaccines are manufactured in yeast cells, and residual Saccharomyces cerevisiae antigens can be present in the product. Anaphylaxis in children with HB vaccine has been rarely reported; it has been related to possible hypersensitivity to yeast (52). Anaphylaxis has been reported in a further HB vaccine
recipient with the causative agent most likely being latex (11). The package components have been changed, and latex is currently not present.

II H. Yellow fever vaccine

Demand for the vaccine is increasing, with more than 60 million doses administered annually (59). The YF vaccine Stamaril (UK) contains 0.13 to 0.61 μg/ml of egg protein (60) and YF-VAX contains 2.43 to 4.42 μg/ml of egg protein (59), used in US. Compared to the recommendations for egg protein in TIV, egg protein in Stamaril is not high. However, no large studies about egg allergy in YF vaccines exist. Anaphylaxis risk from YF vaccine ranges from 0.42 to 1.8/100,000 doses (60). With the low ovalbumin content in the present YF vaccine, desensitization will probably not be necessary henceforth. However, egg allergic persons should be evaluated by an allergist before YF vaccination (See III C).

II I. HPV vaccine

IgE mediated anaphylaxis to quadrivalent HPV vaccine is rare, 2.6/100,000 (61). An expert panel classifying suspected cases using the Brighton Collaboration (BC) case definition of anaphylaxis found eight cases. The panel rejected the possibility that these could have been vasovagal episodes or somatic conversion disorder misdiagnosed as anaphylaxis. The anaphylaxis rate was higher than in previous vaccination programs. However, there was no anaphylactic shock.

Allergenicity of the vaccine is biologically plausible for HPV virus-like particles, which are highly immunogenic when injected (62). Any residual amounts of yeast proteins might cause allergic reactions (12); the quadrivalent vaccine also contains polysorbate 80 as a stabilizer, which might trigger anaphylaxis (63).

II J. TBE – tick-borne encephalitis vaccine

In the 1990s, the TBE vaccine (Encepur, Chiron Vaccines) caused an immediate allergic reaction in approximately 1/50,000 doses and was modified in 1998. The stabilizer polygeline (a gelatine) was replaced with human serum albumin, and the immediate reactions decreased to 0.08-0.24/100,000 doses (64).

III. Diagnostic aspects of severe reactions

In the setting of vaccination reactions, different definitions and grading systems for anaphylaxis have been proposed. Our group prefers the case definition of anaphylaxis established at an NIH consensus conference and subsequently endorsed by WAO and EAACI (Table 2, NIH criteria for anaphylaxis). The definition is widely accepted by allergists.
III A. Diagnostic tests of severe reactions

Serum mast cell tryptase (MCT) levels have been used as a marker of anaphylaxis [65], although its predictive value for vaccine-associated anaphylaxis has not been formally established. We recommend MCT level determined within 2 hours after a systemic vaccine reaction, as well as serum baseline tryptase evaluated at least 48 hours afterwards. A significant increase in MCT level from baseline is a strong indicator of a systemic mast-cell-mediated hypersensitivity reaction.

If a patient has had a suspected allergic reaction to a vaccine, identification of the culprit allergen is important, because it may permit the use of a vaccine formulation without the offending allergen for subsequent doses and also to avoid other products containing these allergens.

**Statement:** Pre-immunization allergy tests (skin test, specific serum IgE) as screening do not reliably predict or exclude future allergic vaccine reactions and are not recommended.

Testing serum IgE to microbial components is frequently unhelpful in preventing allergic vaccine reactions because the IgE response is part of the regular immune response and does not predict an allergic reaction to a vaccine (see section I B). Specific IgE tests are not commercially available for most microbial components. For some other constituents (e.g. ovalbumin and gelatin), the predictive capacity for reaction to vaccines is rather low. False positive tests may occur as many more individuals are allergic and sensitized to a given allergen than those reacting clinically on exposure to the minute amounts of this allergen encountered during immunization.

**Statement:** After a vaccine reaction preferably specific IgE to egg/gelatin/latex/yeast should be analysed when suspected; otherwise skin test is recommended. However, lack of data on the sensitivity and specificity of skin test to vaccines in different concentrations makes them unreliable in predicting or excluding future allergic vaccine reactions (48). More studies are needed to establish thresholds for the prediction of anaphylaxis to a vaccine.

Skin testing can provide additional information about sensitization and the probability of a hapten/allergen being the culprit. This could help evaluate severe vaccine reactions. Skin testing should start with SPT (undiluted), a positive reaction being a sign of an allergic reaction. Skin prick testing sensitivity to vaccines itself is low. If negative, intradermal testing (0.02 mL) should follow (1:100 dilution, 1:10 dilution, see Figure 2). Undiluted intradermal testing is discouraged because of the high rate of irritant (non-relevant) reactions. False positive reactions may also occur at 1:10 dilution especially with influenza, MMR and varicella vaccines, and were even described for 1:100 dilutions in 5% of controls for DT and DTaP, and 15% for influenza (66). Thus, positive reactions should be regarded as indicative rather than confirmatory, and further studies are needed. Positive and negative controls are mandatory.
In non-immediate local reactions, contact dermatitis or subcutaneous nodules, type IV hypersensitivity to preservatives, aluminium, or antibiotics may be assessed by patch testing. Although patch testing is not essential for therapeutic decisions, it could help in choosing alternative vaccines if available.

III B. Local aluminium reactions

**Statement:** Aluminium-allergic persons can be vaccinated with aluminium-containing vaccines without inducing severe reactions, although new itching nodules may appear (67).

Aluminium compounds, such as aluminium phosphate and aluminium hydroxide, are used as vaccine adjuvants and can induce type IV hypersensitivity (contact allergy) (68). Contact hypersensitivity to aluminium was demonstrated in 77% of the children with itching nodules and in 8% of the symptomless siblings who had received the same vaccines, i.e. not a specific test for symptoms. Subcutaneous nodules may develop and persist for months before they gradually disappear (67). Risk factors for aluminium sensitization at vaccination seem to be the dose of aluminium, the number of vaccinations, and the aluminium compound, where aluminium hydroxide seems more liable to induce sensitization than aluminium phosphate.

In a prospective study of 4,758 children, 0.66% (n=38) developed an itching granuloma after Pentavac® (DTaP-Hib-Polio vaccine). When Prevenar® (conjugated pneumococci vaccine) was added, the percentage was 1.2%, and most of them had positive patch tests to aluminium (69). Patch tests with aluminium chloride hexahydrate 2% and elemental aluminium have been suggested, but some cases may be missed unless tested with aluminium chloride hexahydrate 10% (70). Patch tests should be read after three or four days and after one week (71). An itching granuloma and a positive epicutaneous test are illustrated in figure 4 and 5.

III C. Identification of patients at risk and contraindications to immunization

Currently available tools cannot predict most of the severe allergic reactions following immunization. Patients who manifested a severe allergic reaction following immunization are considered at high risk for the next immunization and merit special precautions (72) (see IV B).

Patients who reacted clinically to an allergen contained in the vaccine are at increased risk of allergic vaccine reactions. Although specific sensitization can increase the risk of allergic reaction to vaccines, atopy in general does not seem an important risk factor (73).

**Statement:** Atopy and family history of allergy or asthma are not *per se* contraindications for immunization.

Few real contraindications for routine immunizations exist. Patients are often falsely labelled as allergic although, in most cases, administration of another dose is well tolerated. Patients with anaphylaxis or other...
severe (life-threatening) adverse events following immunization should not be re-immunized with the same vaccine before allergological investigations are completed. Most patients can be immunized safely (see introduction).

**Statement:** Local reactions to antibiotics are not a contraindication for immunization.

Previous localized delayed-type reactions to thiomersal, neomycin or aluminium are not considered absolute reasons for withholding vaccines because the risks of not being immunized outweigh problems caused by local reactions.

Patients with mastocytosis, particularly children, are at increased risk of mast cell-mediated reactions after various triggers including routine vaccination. Therefore, we recommend administering vaccines in single injections, avoiding co-administrations, under medical supervision for at least 30 minutes (74).

**IV. Practical aspects**

As it is important to evaluate whether there is an evident risk of allergic reactions, patients should be asked whether they experienced allergic symptoms following previous vaccinations. Also, underlying uncontrolled diseases must be ruled out.

**Statement:** Expertise and equipment for treating anaphylaxis should always be available when immunizing.

All vaccinating units need to have adrenaline, antihistamine and oral steroids at hand and in most countries beta-2-inhalers. For patients at risk, also parenteral steroids, oxygen and a defibrillator should be available close to where the vaccinations are administered.

**IV A. Immunization of patients at increased risk**

**Statement:** A history of a previous allergic reaction to a vaccine or to one of its constituents should be ascertained before immunization.

Identification of increased risks through clinical history is essential for risk minimization. Patients with a positive history should be investigated for type I hypersensitivity to the vaccine and its ingredients, and vaccination should be managed following specific recommendations for subjects allergic to vaccine components, see III A.
Statement: Immunization under standard conditions (standard vaccine, full dose, no mandatory observation time) is recommended for patients with:

- Allergic sensitization but without a clinical reaction to an allergen contained in the vaccine;
- Allergic disease not related to a vaccine;
- Family history of allergy.

Statement: If, based on a positive benefit/risk balance, an additional dose is needed after an anaphylactic vaccine reaction, a vaccine preparation without the offending ingredient should be preferred.

Statement: Egg-allergic patients can be MMR-immunized under standard conditions.

Data from clinical studies suggest that the small amount of residual egg protein in MMR vaccines represents an exceptionally uncommon risk for egg-allergic patients (75).

Statement: Patients with manifest egg allergy who intend to be influenza-immunized should only be vaccinated with low egg (<0.12 µg/mL) vaccines:

A) Previous non-anaphylactic reactions to egg: can be influenza-vaccinated under standard conditions

B) Previous anaphylaxis to egg: single-dose vaccination with a personal staff experienced in recognizing and treating anaphylactic reactions under observation (minimum 1 hour).

Gelatine-allergic patients could most often receive an alternative vaccine without gelatine as a stabilizer. Otherwise, SPT with the vaccine should be performed and, if positive, fractionated vaccine doses administered (17).

IV B Fractionated immunization or graded desensitization. Management of allergic reactions to vaccines.

Patients sensitized to a vaccine or its components with previous anaphylaxis to this vaccine should be revaccinated only if absolutely necessary. If at all possible, a vaccine without the offending allergen should be chosen. Where this is not possible, two pragmatic (not evidence-based) approaches have been used:

Assuming that a smaller vaccine dose does less harm than a full dose, patients with negative skin tests to the vaccine but with a history of anaphylaxis or other severe allergic reaction can be immunized with split-dose

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vaccination. Initially, 10% of the dose is given, followed 30 min later by the remaining 90% provided that no allergic reaction has occurred after the initial dose.

As in rapid desensitization, immunization in graded doses may reduce the risk of anaphylaxis. Increasing vaccine doses are administered every 15-30 minutes provided that there are no signs of allergic reaction (0.05ml of 1:10 dilution, then 0.05ml, 0.1ml, 0.15ml, 0.2ml, of a 0.5ml full-strength vaccine) (17). Importantly, this protocol only leads to transient desensitization, and patients undergoing this protocol successfully must still be considered allergic to the vaccine. These vaccination approaches must only be used in a controlled setting where prompt treatment of anaphylaxis by experienced staff is available, see Figure 3.

IV C. Delay of routine immunization

**Statement:** Delay of routine immunizations is not recommended. Delay withholds protection from vaccine-preventable disease, and there is no justifiable evidence that it would prevent allergic reactions or development of allergic disease.

One study reported that delaying primary DTP immunization beyond 2 months of age was associated with a 50% risk reduction of recorded asthma by age 7 years (76). This effect could not be replicated (77) and may have been reporting bias. A further study of children with ≥ 2-month delay in the 3rd DTP dose reported a 20% risk reduction in hay fever at school age (78). In contrast, a recent large Swedish study did not show any increased risk of requiring asthma medication whether the first DTaP vaccine was administered at 2 months or at 3 months of age (32). Studies on the effects of delaying other immunizations are lacking. The risk of vaccine-preventable disease outweighs a doubtful risk reduction in allergic disease.

V. Strategic aspects

VA. Surveillance

**Statement:** EAACI should make efforts to register severe vaccine adverse events.

Strategies to monitor AEFI need to be developed, particularly those that may have an underlying allergic aetiology. Here, EAACI can play an important role by encouraging the sharing of best practice and insights gained within and between member countries, and through fostering common surveillance approaches to assess beneficial and adverse impacts of immunization strategies. Greater use of electronic health record systems is likely to be the key to such efforts in the future.

Concerning paediatric patients, adverse reactions to vaccines are already the most common reactions reported to pharmacovigilance systems.
V B. Risk communication

Public interest in the field of risk communication and vaccines is growing, fuelled by contemporary debate about perceived adverse events and easy access to information via the internet, which, however, increases the risk of misinformation. Although public confidence in vaccines is may be decreasing [79, 80], the public’s trust in health care workers remains well documented. Therefore, it is important to properly educate and train vaccine providers to maintain public acceptance of immunizations [81].

The extensive scientific literature on risk communication includes several publications on immunization and allergy, but apart from advice on egg allergy [56], few studies on risk communication specifically address allergy in connection with immunization. The general literature on risk communication highlights the value of transparency, sensitivity and respect, with trust and confidence as essential elements [80, 82]. There is no reason for other strategies when communicating risks concerning immunizations and allergy. Denying or diminishing known risks is unethical and can lead to a higher risk perception among the target group [83].

V C. Education and information for health professionals

To communicate effectively with patients/carers and members of their teams, health care professionals need accurate, authoritative and accessible information on the potential benefits and risks of immunizations. It is unrealistic to expect busy professionals to read, digest and interpret the substantial body of epidemiological and health services research on this subject. They also need tools to communicate these benefits/risks in an open, non-coercive way to foster relationship-building and trust between health providers and patients/carers. As a respected professional body throughout Europe, EAACI can play an important leadership and coordinating role by ensuring the consistency of key messages being transmitted to health professionals throughout Europe and by eliciting information on professional concerns and hitherto unanswered questions.

V D. Future vaccine development and use

Vaccination stimulates different types of Th cells and IgE production. Immunological effects can be considerable, particularly when adjuvants are used. When trials of new vaccines or vaccine components are planned, aspects of clinical allergy and its immunological features should be integrated into research protocols. Also, both stabilizers and adjuvants in new vaccine compositions should be evaluated. New vaccines without egg protein and gelatine would be preferable.

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V E. Research needs

A validated test predicting clinical reactions following vaccination would be of major benefit. Such a study could examine whether graded desensitization has a role in these situations, and the results could be further studied, potentially through a network within EAACI.

Aluminium gives local itchy granuloma from paediatric vaccinations in approximately 1% of cases. A change of adjuvant might be advisable.

Although extensive scientific research has not concluded that vaccination promotes allergic diseases, new data from ongoing studies, and new environmental factors and vaccine constituents will require us to conduct retrospective and prospective studies in the future.

References


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Figure 1: Contamination by culture media in the preparation of vaccines

Figure 2: Diagnostic algorithm in case of suspected allergic reaction to vaccine or vaccine component

Figure 3: Pre-immunization testing and immunization in patients who had a suspected previous allergic reaction to a vaccine

Figure 4: Local reaction after vaccination at 3, 5 and 12 months of age with DTaP-Hib-polio

Figure 5: Epicutaneous test with aluminium 2% in a 2-year-old child
Table 1. Abbreviations:

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>AEFI</td>
<td>adverse event following immunization</td>
</tr>
<tr>
<td>BC</td>
<td>Brighton Collaboration</td>
</tr>
<tr>
<td>CDC</td>
<td>Centers for Disease Control and Prevention</td>
</tr>
<tr>
<td>D</td>
<td>Diphtheria</td>
</tr>
<tr>
<td>DTaP</td>
<td>Diphtheria - Tetanus - Acellular Pertussis</td>
</tr>
<tr>
<td>DTP</td>
<td>Diphtheria - Tetanus - Pertussis</td>
</tr>
<tr>
<td>EAACI</td>
<td>European Academy of Allergy and Clinical Immunology</td>
</tr>
<tr>
<td>HBV</td>
<td>Hepatitis B vaccine</td>
</tr>
<tr>
<td>Hib</td>
<td>Haemophilus Influenzae type b</td>
</tr>
<tr>
<td>IIV</td>
<td>Trivalent and quadrivalent inactivated influenza vaccine</td>
</tr>
<tr>
<td>IPV</td>
<td>Inactivated polio vaccine</td>
</tr>
<tr>
<td>RIV</td>
<td>Recombinant subunit influenza vaccine</td>
</tr>
<tr>
<td>LAIV</td>
<td>Live attenuated trivalent and quadrivalent influenza vaccine</td>
</tr>
<tr>
<td>MCT</td>
<td>Mast Cell Tryptase</td>
</tr>
<tr>
<td>MMR</td>
<td>Measles - Mumps – Rubella</td>
</tr>
<tr>
<td>OPV</td>
<td>Oral Polio Vaccine</td>
</tr>
<tr>
<td>P</td>
<td>Pertussis</td>
</tr>
<tr>
<td>PCV</td>
<td>Pneumococcal conjugated vaccine</td>
</tr>
<tr>
<td>T</td>
<td>Tetanus</td>
</tr>
<tr>
<td>TBE</td>
<td>Tick-borne encephalitis</td>
</tr>
<tr>
<td>TIV</td>
<td>Trivalent inactivated influenza vaccine</td>
</tr>
<tr>
<td>WAO</td>
<td>World Allergy Organization</td>
</tr>
<tr>
<td>WHO</td>
<td>World Health Organization</td>
</tr>
<tr>
<td>YF</td>
<td>Yellow fever</td>
</tr>
</tbody>
</table>
Table 2: Clinical criteria for diagnosing anaphylaxis (NIAID and EAACI)

Anaphylaxis is highly likely when any of the following 3 criteria are fulfilled:
1. Acute onset of an illness (minutes to several hours) with involvement of the skin, mucosal tissue, or both (e.g., generalized hives, pruritus or flushing, swollen lips-tongue-uvula)
   AND AT LEAST ONE OF THE FOLLOWING
   a. Respiratory compromise (e.g., dyspnoea, wheeze-bronchospasm, stridor, reduced PEF, hypoxemia)
   b. Reduced BP or associated symptoms of end-organ dysfunction (e.g., hypotonia [collapse], syncope, incontinence)

2. Two or more of the following that occur rapidly after exposure to a likely allergen for that patient (minutes to several hours):
   a. Involvement of the skin-mucosal tissue (e.g., generalized hives, itch-flush, swollen lips-tongue-uvula)
   b. Respiratory compromise (e.g., dyspnoea, wheeze-bronchospasm, stridor, reduced PEF, hypoxemia)
   c. Reduced BP or associated symptoms (e.g., hypotonia [collapse], syncope, incontinence)
   d. Persistent gastrointestinal symptoms (e.g., crampy abdominal pain, vomiting)

3. Reduced BP after exposure to known allergen for that patient (minutes to several hours):
   a. Infants and children: low systolic BP (age-specific) or greater than 30% decrease in systolic BP*
   b. Adults: systolic BP of less than 90 mm Hg or greater than 30% decrease from that person’s baseline

Reproduced with permission from Hugh Sampson, see Muraro et al. PEF, Peak expiratory flow; BP, blood pressure.
*Low systolic blood pressure for children is defined as less than 70 mm Hg from 1 month to 1 year, less than (70 mm Hg + [2 x age]) from 1 to 10 years, and less than 90 mm Hg from 11 to 17 years.

From: Hugh Sampson, and used in the position paper in Allergy 2014 (19)
Table 3: Anaphylaxis after vaccination, rates; from NcNeil et al, 2016 (20)
Brighton Collaboration case definition

<table>
<thead>
<tr>
<th>Vaccine</th>
<th>Rate/million doses</th>
<th>Total doses administered (in millions)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hib</td>
<td>0</td>
<td>1.14</td>
</tr>
<tr>
<td>Hepatitis B</td>
<td>0</td>
<td>1.29</td>
</tr>
<tr>
<td>Influenza (TIV)</td>
<td>1.59</td>
<td>8.83</td>
</tr>
<tr>
<td>MMR</td>
<td>5.14</td>
<td>0.58</td>
</tr>
<tr>
<td>Pertussis (dTap)</td>
<td>2.89</td>
<td>3.12</td>
</tr>
<tr>
<td>Pertussis (DTaP)</td>
<td>2.07</td>
<td>1.45</td>
</tr>
<tr>
<td>Pneumococcal (PCV13)</td>
<td>0</td>
<td>0.74</td>
</tr>
<tr>
<td>IPV</td>
<td>1.65</td>
<td>1.22</td>
</tr>
<tr>
<td><strong>All vaccines</strong></td>
<td><strong>1.31</strong></td>
<td><strong>25.17</strong></td>
</tr>
</tbody>
</table>
Table 4 Differentiation of anaphylaxis and vasovagal reaction.

<table>
<thead>
<tr>
<th>Possible symptoms</th>
<th>Anaphylactic reaction</th>
<th>Vasovagal reaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Onset from time of</td>
<td>Few minutes delay, typically within 30 minutes</td>
<td>During or shortly after injection</td>
</tr>
<tr>
<td>immunization</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Respiratory</td>
<td>Wheezing, stridor</td>
<td>Normal or hyperventilation</td>
</tr>
<tr>
<td>Cardiovascular</td>
<td>Tachycardia, hypotension</td>
<td>Self-limited bradycardia, hypotension</td>
</tr>
<tr>
<td>Skin</td>
<td>Flushing, itchy rash, angioedema, urticaria</td>
<td>Pale, sweaty, cold, clammy</td>
</tr>
<tr>
<td>Gastrointestinal</td>
<td>Abdominal cramps</td>
<td>Nausea, vomiting</td>
</tr>
<tr>
<td>Neurological</td>
<td>Loss of or altered consciousness, little response</td>
<td>Self-limited loss of consciousness, good</td>
</tr>
<tr>
<td></td>
<td>to prone positioning</td>
<td>response to prone positioning</td>
</tr>
</tbody>
</table>


Table 5: Common, minor local vaccine reactions

<table>
<thead>
<tr>
<th>Vaccine</th>
<th>Local adverse events (pain, swelling, redness)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Measles/MR/MMR</td>
<td>1 of 20 (mild rash)</td>
</tr>
<tr>
<td>Pertussis (DTaP)</td>
<td>1 of 4* (redness or swelling)</td>
</tr>
<tr>
<td>Pneumococcal conjugate (PCV 13)</td>
<td>1 of 3 (swelling)</td>
</tr>
<tr>
<td>Pneumococcal unconjugated</td>
<td>1 of 2 (redness or pain)</td>
</tr>
<tr>
<td>Tdap</td>
<td>1 of 5 (redness or swelling) (3 of 4 pain)</td>
</tr>
<tr>
<td>Varicella</td>
<td>1 of 5 (soreness or swelling)</td>
</tr>
<tr>
<td>HPV (quadrivalent)</td>
<td>1 of 3 (redness or swelling)</td>
</tr>
</tbody>
</table>

*More often after the 4th and 5th dose

Source: http://www.cdc.gov/vaccines/vac-gen/side-effects.htm

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Figure 1:

Contamination by culture media in the preparation of vaccines

- Human diploid cell (HDC) culture
  Rabies
  No ovalbumin

- Chicken fibroblast cell culture
  Measles-mumps-rubella, TBE, rabies
  ≤1ng per dose

- Chicken embryonated eggs
  Influenza
  ≤1.6ug per dose

- Chicken embryos
  Yellow fever
  ≤16ug per dose
Figure 2: Diagnostic algorithm in case of suspected allergic reaction to vaccine or vaccine component

- **Skin prick test**
  
  *(vaccine or vaccine component)*
  
  **undiluted**

  **If negative**

- **Intradermal test**
  
  *(vaccine)*
  
  **1:100 dilution**

  **If negative**

- **Intradermal test**
  
  *(vaccine)*
  
  **1:10 dilution**

  **Cave:** local irritant reaction possible
Figure 3: Pre-immunization testing and immunization in patients who had a suspected previous allergic reaction to a vaccine

<table>
<thead>
<tr>
<th>Allergic reaction to previous vaccine dose</th>
<th>Skin test result</th>
<th>Vaccine administration</th>
<th>Precautions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Local reaction</td>
<td>Not needed</td>
<td>Full dose</td>
<td>No observation period</td>
</tr>
<tr>
<td>Anaphylaxis, systemic reaction</td>
<td>Negative</td>
<td>Allergen avoidance if possible, split dose</td>
<td>60 minutes observation, IV line</td>
</tr>
<tr>
<td>Anaphylaxis, systemic reaction</td>
<td>Positive</td>
<td>Allergen avoidance if possible, graded doses</td>
<td>60 minutes observation, monitoring, IV line</td>
</tr>
</tbody>
</table>

*Allergen avoidance does not mean no vaccination, but using an allergen-free vaccine or a low allergen content vaccine, if available.
Figure 4: Local reaction after vaccination at 3, 5 and 12 months of age with DTaP-Hib-polio
Figure 5: Epicutaneous test with aluminium 2% in a 2-year-old child