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Conflict of Interest

Dr. Diamant reports personal fees from ALK, AstraZeneca, Boehringer Ingelheim, GlaxoSmithKline, HAL Allergy, Merck Sharp & Dohme, and Sanofi-Genzyme-Regeneron and acted as Research Directorat QPS-NL, an institution which received research support from several bio-pharmaceutical companies, esp within respiratory: HAL Allergy, Foresee Pharmaceuticals, Patara Pharma (now Respivant), Novartis; Dr. Jesenak received honoraria, consultancy and speaker fees from Sanofi-Pasteur, Pfizer, GlaxoSmithKline, Merck Sharp&Dohme, Takeda, ALK, Stallergenes-Greer, Angelini, Novartis, Mundipharma and Berlin-Chemie and acted as principal investigator in trials sponsored by BioCryst, Pharming, Octapharma, and Baxalta; Dr. Vieths reports personal fees from Swiss Society for Allergy and Immunology, SchattauerAllergologieHandbuch, Elsevier Nahrungsmittelallergien und Intoleranzen, and Karger Food Allergy: Molecular Basis and Clinical Practice and also reports non-financial support from German Research Foundation, European Directorate for the Quality of Medicines and Health Care, European Academy of Allergy and Clinical Immunology, German Chemical Society (GDCh), AKM Allergiekongress, International Union of Immunological Societies, and from Spanish Society for Allergy and Clinical Immunology (SEAIC); Dr. Agache is an Associate Editor at Allergy and PAI; Dr. Chinthrajah reports grants from NIAID, CoFAR, Aimmune, DBV Technologies, Astellas, Regeneron, and FARE and is an advisory board member of Alladapt Therapeutics, Genentech, Novartis, and Sanofi; Dr. Eiwegger acts as local PI for company sponsored trials by DBV and sub-investigator for Regeneron, holds grants from Innovation fund Denmark, CIHR. He is Co-Investigator or scientific lead in three investigator initiated oral immunotherapy trials supported by the Food Allergy and Anaphylaxis Program SickKids and serves as associate editor for Allergy. He/his lab received unconditional/kind contributions from Macro Array Diagnostics and ALK. He holds advisory board roles for ALK; Dr. Barber reports grants from ALK, ALLERO therapeutics, personal fees from ALK and Aimmune; Dr. Ollert reports personal fees from Hycor Biomedical and is Scientific cofounder of TolerogenicsSarL; Dr. Palomares received research grants from Inmunotek S.L., Novartis and MINECO, received fees for giving scientific lectures or participation in Advisory Boards from Allergy Therapeutics, Amgen, AstraZeneca, Diater, GlaxoSmithKline, S.A, Inmunotek S.L, Novartis, Sanofi-Genzyme and Stallergenes; Dr. Pfaar reports grants and personal fees from ALK-Abelló, Allergopharma, Stallergenes Greer, HAL Allergy

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Abstract

Vaccines are essential public health tools with a favorable safety profile and prophylactic effectiveness that have historically played significant roles in reducing infectious disease burden in populations, when the majority of individuals are vaccinated. The COVID-19 vaccines are expected to have similar positive impacts on health across the globe. While serious allergic reactions to vaccines are rare, their underlying mechanisms and implications for clinical management should be considered to provide individuals with the safest care possible. In this review, we provide an overview of different types of allergic adverse reactions that can potentially occur aftervaccination and individual vaccine components capable of causing the allergic adverse reactions. We present the incidence of allergic adverse reactions during clinical studies and through post-authorization and post-marketing surveillance and provide plausible causes of these reactions based on potential allergenic components present in several common vaccines. Additionally, we review implications for individual diagnosis and management and vaccine manufacturing overall. Finally, we suggest areas for future research.

Keywords: allergy, anaphylaxis, COVID-19, SARS-CoV-2, vaccine

Word count: 5162

Introduction

The rapid development and the launch of several novelCOVID-19 vaccines for severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) is an extraordinary and remarkable accomplishment of modern science. ThePfizer-BioNTech BNT162b2 was the first vaccine to be granted temporary authorization for emergency use by the Medicines and Healthcare Products Regulatory Agency (MHRA) in the U.K for the treatment of COVID-19on Dec 2, 2020.¹ Soon after, on Dec 11, 2020, it also received emergency use authorization (EUA) by the U.S. Food and Drug Administration (FDA).²EUA is a mechanism to facilitate the availability of vaccines during public health emergencies, such as the current COVID-19 pandemic. Under an EUA, the FDA may allow the use of unapproved medical products (including vaccines) to prevent serious or life-threatening disease when certain statutory criteria have been met and no adequate and/or approved alternatives are available. The authorization of BNT162b2 was followed by an EUA fora second COVID-19 vaccine, the Moderna mRNA-1273 on Dec 18, 2020.3 This was followed by the authorization of mRNA-1273 for use by other regulatory agencies such as the European Commission, UK MHRA, Israel Ministry of Health, and others.⁴ On December 30, a third COVID-19 vaccine, the Oxford/AstraZeneca recombinant adenoviral AZD1222orChAdOx1-S was authorized for use by the UK MHRA.⁵In addition to the above authorized COVID-19 vaccines, a number of other novel COVID-19 vaccines are currently indifferent phases of clinicals development. Currently used platforms in COVID-19vaccines include classical and novel platforms, such as RNA- and DNA-based, viral vectors (nonreplicating), protein subunits, virus like particles and inactivated viral platform (Table 1).⁶⁻⁸

With the authorization of COVID-19 vaccines, vaccination campaigns have been initiated in many areas throughout the world. Within the first few days of public vaccinations, however, BNT162b2 was associated with a few severe cases of anaphylaxis.⁹ While severe allergic reactions may pose a potential risk with any vaccine (or systemic medications), the benefits of vaccination outweigh the potential risks of receiving the vaccine for the vast majority of individuals. However, the fear of allergic reactions may lead to vaccine hesitancy, which could compromise herd immunity and limit efforts to contain the pandemic. It is therefore critical that we understand the immunopathological changes in COVID-19,¹⁰risk of severe allergic reactions, and their mechanisms

in order to improve individual safety and issueproper guidance with the goal of vaccinatingthe maximum number of individuals. Here, we review adverse events associated with vaccine-related allergic and non-allergic reactions to COVID-19 and other vaccines, mechanisms associated with allergic adverse reactions, and rates of severe allergic reactions with existing vaccines and with the novel COVID-19 active vaccines currently being administered to large parts of the world.

Section I:Vaccine-associated allergic vs non-allergic reactions

There are a number of vaccines currently used with proven safety and efficacy. Vaccines are potentially associated with adverse events. An adverse event is defined asany untoward medical occurrence associated with the use of a drug in humans, whether or not considered drug related.¹¹ Adverse events can present as local or systemic, immediate or non-immediate, and immune or non-immune mediated reactions. While all allergic reactions are immune-mediated, not all immune-mediated reactions are allergic.Local non-immediate reactionsthat are not allergenic are common and may includeswelling and erythema at the injection site. These reactions can occur hours or days after administration and are not always mediated through the immune system. Systemic non-allergic reactions include mild fever and vasovagal reactions such as hypotension, nausea, and syncope are also relatively common.Neither the local nor the vasovagal reactions pose any serious risk. Although some of these reactions are immune-mediated, they are not allergic reactions. Rather, soreness at the injection site or fatigue are consequences of activation of the innate immune system.¹²⁻¹⁵

As of Feb 10, 2021, 44.77 million people had taken one or two doses of a COVID vaccine in the US. 653 deaths and 12,697 total adverse events had been reported to following COVID-19 vaccinations to the CDC and Prevention's VAERS.¹⁶Adverse events, including allergic reactions, are graded according to severity as mild, moderate, and for purposes of this review, serious. Typical signs of an allergic reaction include bronchoconstriction, conjunctivitis, rhinorrhea, gastrointestinal symptoms, and/or characteristic skin lesions such as generalized urticaria and/or angioedema. These can occur in combination or alone, and onset can be immediate, within minutes, or up to several hours post-vaccination. Examples of mild allergic reactions are swelling with itching at the injection site, conjunctivitis, or rhinorrhea. Examples of moderate allergic reactions are bronchoconstriction that can be adequately treated with nebulized beta-agonists or generalized urticaria that may be treated with an

antihistamine. Serious adverse events (SAE) are those events that are life-threatening, require inpatient hospitalization or prolongation of existing hospitalization, cause a persistent or significant incapacity or substantial disruption in the ability to conduct normal life functions, a congenital anomaly/birth defect, or death. Two examples of SAE that are allergic reactions are bronchospasm that requires intensive treatment and life-threatening anaphylaxis.^{11,17}

Anaphylaxis, animmediate systemic multi-organ reaction, is rare but can be lifethreatening.Organs affected include the cutaneous, gastrointestinal, respiratory, and cardiovascular systems. Anaphylaxis can be either immunological, non-immunological, or idiopathic. Idiopathic anaphylaxis is diagnosed through exclusion of other known causes and may mask a clonal mast cell disorder.¹⁸⁻²³Non-immunological anaphylaxis was previously termed anaphylactoid reactions, but the World Allergy Organization (WAO) in 2004 suggested replacing anaphylactoid reactions with nonimmunological anaphylaxis.²⁴ The change in terminology is to reinforce the risk and potential fatality of all types of anaphylaxis, regardless of the underlying mechanism. All three mechanisms of anaphylaxis produce the same clinical picture. (see **Section IV** on mechanisms below). Distinguishing between systemic vasovagal reactions and anaphylaxis during immunization is critical to ensure that appropriate and immediate treatment can be administered (**Table 2**). Vasovagal reactions usually occurimmediately or up to 30 minutes of vaccine administration. Similar to anaphylaxis, organs affected include the cutaneous, gastrointestinal, respiratory, neurological, and cardiovascular systems.^{25,26}

Anaphylactic reactions areconsideredadverse events of special interest (AESI)²⁷, i.e. adverse events that are of significant medical and scientific concern for which immediate medical action with ongoing monitoring and rapid communication by the investigator or sponsor is required.AESI reporting is a critical aspect of pharmacovigilance forcharacterization of the safety profile of a drug or vaccine in context of previous reports of the vaccine or of other vaccines with similar manufacturing processes, formulation, immunogenicity, and novelty. AESIs alert regulators to potential risks. Particularly in mass vaccination programs where a large number of adverse reactions may be reported, identification and assessment of AESIs are a high priority because they highlight potential risks that may alter risk-benefit profile and may require immediate investigation, regulatory action, and prompt communication with the public.^{28,29}**Table 3** lists Pharmacovigilance Practices that follow authorization of a vaccine.

Section II: Allergic reactions to vaccines

In the last 120 years, global vaccination programs haveeradicated or vastly reduced the incidence of debilitating infectious diseases such as smallpox, polio, and measles.³⁰According to the Institute of Medicine, epidemiologic and mechanistic evidence support a causal relationship between anaphylaxis and several vaccines, including those for measles, mumps, and rubella (MMR), varicella, influenza, hepatitis B, meningococcus, human papillomavirus, and the combined diphtheria, tetanus, pertussis (DTaP or TdaP) vaccine.³¹Although approved vaccines have been rigorously tested for safety, anaphylactic reactions, although rare, can occur in individuals.³²An analysis of reported anaphylaxis to the Vaccine Adverse Reporting System (VAERS) in the United States over a 26-year period found that out of the almost 500,000 reports to VAERS, only 828 were classified as anaphylaxis based on either on physician's diagnosis or in according to the Brighton Collaboration case standards.³³ A 2003 study analyzing over eight million routine vaccinations in the Vaccine Safety Datalink (VSD) Project³⁴ found the risk of anaphylaxis ranged between 0.65 and 1.53 cases/million doses. They also noted that most anaphylactic episodes occurred when multiple vaccines were administered during the same visit.³⁵Similarly, a 2016 study used health data from VSD and found 33 confirmed cases of anaphylaxis after 25,173,965 vaccine doses or an anaphylaxis rate of 1.31 per million vaccine doses.³⁶ The study also found that 85% of cases of anaphylaxis had pre-existing atopic disease, which was consistent with earlier reports emphasizing coexisting atopic disease, particularly asthma, as being clinical risk factors for anaphylaxis.³⁷In Asian children, analysis of a large-linked database found risk of anaphylaxis to be 1.21 cases/million doses.³⁸A study conducted in Australia found that estimated incidence rate of anaphylaxis for DTaP vaccines was 0.36 cases per 100,000 doses, and 1.25 per 100,000 doses for MMR vaccines.³⁹ Overall, rates of reported anaphylaxis occurs at arate of about 1 per 100,000 to one per 1,000,000 depending on the vaccine.⁴⁰Figure 1 and Table 4shows the frequency of anaphylaxis for specific vaccines. Table 5 provides population-specific considerations for common vaccines.⁴¹⁻⁴⁴

Section III: Allergic reactions to COVID-19 mRNA vaccines

Development of the SARS-CoV-2 mRNA vaccine occurred in record time. So far, three candidate vaccines (mRNA BNT162b2, mRNA-1273 and adenovirus vectored ChAdOx1) have been authorized for COVID-19 in the European Union and the United Kingdom. Of these, mRNA BNT162b2 and mRNA-1273 are authorized for emergency use in the United States. Additional candidates have entered or are completing the pivotal stage of clinical development programs. At the time of this review, there are 64 vaccines in several stages of clinical development and 173 in pre-clinical development worldwide.⁴⁵Details of their composition, mode of action, and developmental stagecan be found in the review by Rodriguez-Coiraet al.⁴⁶

Even after formal authorization, vaccine rollout in the United Kingdom, United States, and European Union has been and continues to be challenging. Navigating complex distribution logistics, determining ethical allocation of a limited resource, and de-mystifying widespread news coverage of anaphylactic events that perpetuate vaccine hesitancy are among the most pressing. While anaphylaxis after routine vaccination is very rare, it is important for the scientific community to be informed and prepared to treat adverse vaccine reactions to increase safety and acceptability. During the COVID-19 pandemic, vast quantities of vaccine are expected to be administered over a very short period of time, with increased public awareness and surveillance. In this situation, the probability of reporting anaphylaxis likely increases without the added context of the denominator, which is the millions of individuals receiving the vaccine.

Of the vaccines authorized for use in Europe and the US, the first vaccine to be administered and distributed was the mRNA BNT162b2 vaccine in the UK. Following two reports of allergic reactions in the UK on December 8, 2020, the MHRA updated its guidelines to state that individuals should not receive the vaccine if they have had previous anaphylaxis to a vaccine, medicine, or food. In addition, MHRA recommended that individuals who experience anaphylaxis after their first dose of the mRNA BNT162b2 vaccine should not receive the second dose, and that everyone should be monitored for a minimum of 15 minutes after vaccination. However, on Dec 30, after a review of additional data, the guidelines were further updated to indicate that anyone with a previous history of allergic reactions to the ingredients of the vaccine should not receive it, but those with any other allergies such as a food allergy can receive the vaccine.⁴⁷

A CDC report of the VAERS monitoring database indicates that between December 14–23, 2020, 1,893,360 first doses of the mRNA BNT162b2 COVID-19 vaccine were administered, and 21 cases of anaphylaxis were reported (11.1 cases per million doses). Seventy one percent of these reactions occurred within 15 minutes of vaccination.⁴⁸ Similarly, the CDC reported that between December 21, 2020–January 10, 2021, 10 cases of anaphylaxis were reported after administration of 4,041,396 first doses of the mRNA-1273 vaccine (2.5 cases per million doses administered). In nine cases, onset occurred within 15 minutes of vaccination. No anaphylaxis-related deaths were reported.⁴⁹This suggests that the incidence of anaphylaxis in the mRNA BNT162b2 (11.1 cases per million doses) and mRNA-1273 COVID-19 vaccines (2.5 cases per million doses) may be about 2 to 8.5 times as high as the incidence reported in the 2016 VSD study for all vaccines (1.31 per million doses). The US FDA authorized labeling currently lists past severe allergic reactions (e.g. anaphylaxis) to any component of the vaccine as a contraindication to the mRNA BNT162b2 vaccine. Yet, the CDC also recommends that individuals with a history of immediate allergic reactions to other vaccines weigh their risk of exposure, risk of severe disease or death due to COVID-19, and consider whether they were previously infected with COVID-19 (because of lower rates of reinfection in the three-month period after infection), when deciding whether to delay or forego vaccination.^{50,51}

After the mRNA BNT162b2 vaccine received central European marketing approval on December 21, 2020, the Paul-Ehrlich-Institut, Federal Institute for Vaccines and Biomedicines in Germany, and the European Medicines Agency (EMA) released its own guidance for vaccination of people with allergies. These guidelines recommended that only individuals in the European Union with an allergy to a specific vaccine component not receive the vaccine.⁵²TheEuropean Academy of Allergy and Clinical Immunology (EAACI) position paper on diagnosis, management, and prevention of severe allergic reactions to the COVID-19 vaccines states that the vaccines are contraindicated only when there is an allergy to one of the vaccine components or if there was a severe allergic reaction to the first dose. The paper also provides a simplified algorithm of prevention, diagnosis and treatment of severe allergic reactions and a list of recommended medications and equipment for vaccine centers.⁵³

Section IV.General mechanisms in pathways of immune-mediated and non-immune-mediated reactions.

Immunological hypersensitivity is either IgE-mediated (hypersensitivity reaction type 1),⁵⁴ IgG- mediated (hypersensitivity reaction type 2), immune complex and/or complement-mediated (hypersensitivity reaction type 3) or cell-mediated (hypersensitivity reaction type 4) (**Figure 2**).⁵⁵ The mechanisms underlying IgE-mediated hypersensitivity are best understood. Individuals with IgEmediated allergic reactions first undergo an initial sensitization phase during which allergen specific IgE antibodies bind to high affinity FccRI receptors on mast cells and basophils. Subsequent allergen exposure can cross-link the cell-boundIgE antibodies and triggers degranulation of mast cells and/or basophils, with subsequent release of histamine and other inflammatory chemical mediators (cytokines, interleukins, leukotrienes, and prostaglandins) into the surrounding tissue causing several systemic effects, such as vasodilation, mucous secretion, tissue eosinophilic infiltration, and airway smooth muscle contraction.⁵⁶

Type II IgG-mediated immune vaccine reactions are rare but have beenobserved with an MMR vaccine containing dextran before those preparations were taken off the market.⁵⁷The occurrence of large, local injection-site reactions with TdaP vaccines have been reported to be due to a local immune complex type III hypersensitivity reaction.⁵⁸ Type IV hypersensitivity reactions occur when an individual's T cells provoke an inflammatory response against allergens, leading to T cell activation and the release of cytokines and chemokines.⁵⁹Type IV reactions to vaccines induce local eczema, which may start between 2 hours to up to 2 days after vaccinations. These reactions are typically observed following vaccines containing aluminum and anti-microbial agents. The occurrence of such an event is not a contraindication for further vaccinations.⁶⁰

Non-immunologic anaphylaxis is caused by agents or events that induce sudden, massive mast cell and/or basophil degranulation in the absence of immunoglobulins. These reactions may be due to activation of complement by nanoparticles, colloidal solutions, or liposomes without immune complex formation, commonly termed complement-activation-related pseudoallergy or CARPAs⁶¹ (e.g., medications containing Cremophor EL⁶²), direct mast cell and basophil activation resulting in histamine release (vancomycin⁶³ and opiate medications⁶¹), or other mechanisms (activation of the kallikrein-kinin pathway⁶⁴). Nonsteroidal anti-inflammatory drugs (NSAIDs)⁶⁵, local anesthetics⁶⁶, monoclonal antibodies, and chemotherapeutic agents have also been reported to induce non-

immunologic anaphylaxis. Recently, transient receptor potential cation receptor, subfamily V (TRPV4) channel has been implicated as a driver of IgE-independent mast cell-dependent bronchospasm via cysteinyl leukotrienes release.⁶⁷In addition, MRGPRX2, a Mas-related G protein-coupled receptor, has been identified as a mechanism for mast cell degranulation. Since 2015, evidence has accumulated that off target occupation of this receptor by different therapy regimens, such as neuromuscular blocking agents (NMBA) and opioids could constitute an additional mechanism of non-immune immediate drug hypersensitivity. ⁶⁸⁻⁷⁰While radiocontrast agents have traditionally been considered to be non-immunologic, some of the newer, low-osmolar agents may induce IgE-mediated reactions.²¹

Although the clinical presentations of both IgE-mediated anaphylaxis and non-immunologic anaphylaxis are similar, measurements of tryptase and SC5b-9may assist in differentiating the two types.^{71,72} Tryptase is a marker of mast cell activationwhich is released following mast cell degranulation while SC5b-9 is a marker of complement activation and is a terminal complement complex.^{71,73}As both tryptase and SC5b-9 are transiently elevated soon after an anaphylaxis episode, blood should ideally be collected between 30 and 90 minutes after the onset of reaction.⁷¹Acute serum total tryptase should be at least 20% plus 2 ng/ml over the baseline tryptase level.⁷³Another novel emerging biomarker is hereditary α -tryptasemia which is present in mastocytosis and may be useful for determining the individual patient's risk of developing severe anaphylaxis.⁷⁴Figure 3 depicts the mechanisms of IgE-mediated and non-immunological anaphylaxis.⁷⁵⁻⁸¹

Section V: Proven and suspected allergenic components of vaccines

Anaphylaxis to vaccines israre and occurs primarily among individuals who have histories of allergies to the components of the vaccines.¹⁹Allergic reactions after vaccination can be due to any of the vaccine components such as antigens, adjuvants, stabilizers, preservatives, emulsifiers, leached packaging components, residual antibiotics, cell culture materials, and inactivating ingredients (**Box** 1).⁸²**Table 6** listscomponents that have been implicated in allergenic reactions and related adverse events. Here we discuss some of the most common allergenic or potentially allergenic components of vaccines.

Many vaccines contain small amounts of the egg protein ovalbumin. Influenza, yellow fever, and rabies vaccines tend to have higher concentrations of ovalbumin because they are cultured in embryonated chicken eggs.⁸³ Vaccines cultured in chicken embryo fibroblasts, such as the MMR vaccine, have lower concentrations of egg protein than those cultured in embryonic eggs.⁸⁴While egg allergy is common in childhood, studies have shown that vaccinating egg-allergic children with MMR and influenza vaccines is well tolerated and risk of an allergic reaction is similar in the general population.^{85,86}Specifically,egg-allergic children, including those who have had anaphylaxis, were successfully vaccinated with yellow fever⁸⁷vaccines with no serious adverse events reported. Since severe allergic reactions to egg-based influenza vaccines are rare, the CDC and its Advisory Committee on Immunization Practices (ACIP) guidelinesstate that individuals with mild egg allergy can receive any licensed and recommended age-appropriate flu vaccine and no longer need to be observed for 30 minutes after receiving the vaccine. However, inthose with severe egg allergy, the vaccines should only be givenunder the supervision of a health care provider who is capable of recognizing and managingserious allergic conditions.^{88,89}

Gelatin, a protein derived from bovine or porcine sources, is added to both live and inactivated vaccines as a stabilizing agent.⁶⁰Sensitivity to gelatin was confirmed with both skin-prick tests and by immunoassay in a 17-year old female who had an anaphylactic reaction to an MMR vaccine.⁹⁰A retrospective case-control study which interviewed and collected sera from individuals who had suffered anaphylaxis after receiving MMR found that 27% of them had anti-gelatin IgE. In comparison, anti-gelatin IgE was not detected in any of the vaccinated subjects who did not present with adverse events.⁹¹It was subsequently shown that patients who have anaphylaxis to MMR were sensitized to gelatin present in the DTaP vaccine^{92,93}, and that cellular immunity to gelatin from the DTaP vaccine can persist for more than three years.⁹⁴ However, sensitization may also persist due to exposure to gelatin in foods or through cross-reactivity to other allergens such as egg, chicken and cow's milk ⁹⁵. Gelatin is also a source of alpha-gal, an carbohydrate allergen that causes meat allergy.⁹⁶Anaphylaxis was observed after vaccination with MMR, Varicella, and DTaP/IPV in pediatric subjects with alpha-gal allergy.⁹⁷Removal of gelatin from vaccines has dramatically reduced allergic reactions to these vaccines.⁹⁸

Milk proteins are used as growth media in DTaP and Tdap vaccines. Although bovine casein is present in nanogram quantities in these vaccines, they rarely cause anaphylaxis. Kattan et al reported eight children with severe cow's milk allergy who reacted with anaphylaxis to the booster dose of DTaP or Tdap vaccine and suggested casein present in the vaccines may play role in the induction of anaphylaxis in atopic children.^{99,100}However, the methods used in this report were questioned¹⁰⁰ and to our knowledge, there have been no subsequent data that support a causative role for DTaP or Tdap vaccines in the induction of allergic disease. It is the position of EAACI (or subcommittee) that these vaccines do not contribute to the pathogenesis of allergic disease and that atopy is not a contraindication to these vaccines.¹⁰¹

As stated above, dextran present in one preparation of MMR vaccine was responsible for numerous cases of anaphylaxis, but this brand of MMR vaccine has since been withdrawn from the market.¹⁰² It was used as a medium nutrient or as a stabilizer. Similarly, during Brazil's national MMR vaccination campaign in 2004, the rate of hypersensitivity following MMR vaccination was unexpectedly high while its case-control study showed no association with a history of allergy.¹⁰³ However, subsequent studies implicated dextran as the likely cause of these hypersensitivity events.¹⁰⁴

Many vaccines contain antigens that are created in cell lines. For example, hepatitis B vaccines and the human papillomavirus (HPV) vaccinescontain antigens that are recombinant proteins expressed in Baker's yeast.¹⁰⁵ Purification removes most of the cellular material, but it is impossible to remove all trace components. Between 1990 and 2004, only 15 reports were identified of probable or possible anaphylaxis following vaccination of individuals with a reported history of yeast allergy. Elevenof these occurred after administration of the hepatitis B vaccine, which contains trace amounts of yeast proteins. Because these subjects were not tested for yeast allergy, it cannot be confirmed that sensitivity to yeast caused these adverse reactions. These data indicate that recombinant yeast-derived hepatitis B vaccine poses minimal risk of allergic reactions in yeast-sensitive individuals. Therefore, evaluation by an allergist is recommended for people who have a history of severe yeast allergy before administration of hepatitis B and HPV vaccines.¹⁰⁵ According to VAERS, there were 107 reports of adverse events in those with a history of yeast allergy present prior to vaccination; of these 11 recipients of hepatitis B vaccine had probable or possible anaphylaxis events.¹⁰⁶By contrast,

another study found no episodes of anaphylaxis in a large cohort of women who had positive skin tests to yeast extract after the HPV vaccine.¹⁰⁷

Antibiotics such as neomycin, streptomycin, polymyxin B, kanamycin and gentamicin are well known to cause mild to life-threatening allergic reactions. Anindividual receiving an MMR vaccinecontaining neomycin was reported to have experienced anaphylaxis shortly after vaccination.¹⁰⁸ Although a skin test to neomycin alone could not be performed in this individual due to a lack of a commercial preparation suitable for skin testing, patient history indicated systemic sensitivity on topical application of neomycin during infancy to disrupted skin. In another case, a history of previous reaction and positive skin test to neomycin was not associated with immediate or delayed hypersensitivity reactions following MMR vaccination.¹⁰⁹In a report of anaphylaxis after rabies vaccination, the presence of residual kanamycin in the vaccine and a positive kanamycin result to an antibiotic skin sensitivity test suggested that kanamycin was the likely cause of the adverse event.¹¹⁰Finally, there is one report of anaphylaxis after applying eye drops containing polymyxin B, an excipient used in DTaP and other vaccines.^{43,111}To our knowledge, no other antibiotics have been associated with vaccine-associated anaphylaxis. 2-Phenoxyethanol is widely used as preservative in cosmetics and vaccines due to its large spectrum of antimicrobial activity, and is considered as one of the most well-tolerated preservatives.¹¹²

Natural latex allergy is well characterized among healthcare personnel and latex content in vaccines as vial stopper or syringe plunger may pose safety concerns in this population. However, Smith et al could not detect latex allergens in adult vaccines.¹¹³In 2004, ananalysis of VAERS revealed only 28 cases of immediate hypersensitivity with latex allergy in vaccine recipients among 160,000 reports of vaccine-associated adverse events.¹¹⁴

Thimerosal, which is approximately 50% mercury by weight, has been one of the most widely used preservatives in vaccines to prevent growth of harmful microbes. All vaccines routinely recommended for children 6 years of age and younger in the U.S. are available in formulations that do not contain thimerosal.¹¹⁵ A risk assessment study revealed that except for local hypersensitivity reactions, there is no evidence of harm caused by thimerosal in vaccines.¹¹⁶Thus, while thimerosal is

the most prevalent preservative inducing contact dermatitis, it is considered irrelevant to vaccineinduced anaphylaxis.¹¹⁷

Formaldehyde and beta-propiolactone (BPL) have been used to inactivate viruses during the production of vaccines. However, approximately 6% of individuals who receive a booster dose of the rabies human diploid cell vaccine (HDCV) develop an immune complex-like reaction in the 2–21 days that follow.^{118,119}These reactions have been associated with the presence of BPL altered human albumin contained in the HDCV.^{120,121}Currently, SinoPharm' BBIBP-CorV¹²²and Sinovac Life Sciences's CoronaVac's¹²³COVID-19 vaccine are using BPL to inactivate SARS-CoV-2. Both vaccines are approved for use in China. Although formaldehyde is only found in residual quantities in vaccine preparation, it has been reported to aggravate eczematous dermatitis following hepatitis B vaccination.¹²⁴It is used to inactivate virus or for the detoxification of bacterial toxin. Formaldehyde-specific contact dermatitis had also been reported following formaldehyde-containing influenza vaccine.¹²⁵ It is hypothesized that the introduction of carbonyl groups on antigens by formaldehyde in vaccines profoundly affects its immunogenicity, thus explaining adverse effects due to formaldehyde-containing vaccines.^{126,127}

Adjuvants are incorporated into some vaccines to boost T-cell immunity and increase helper T-cell function. The most commonly used adjuvants in vaccines are aluminum hydroxide and aluminum phosphate. Despite its long-standing use as an adjuvant in vaccines, aluminum has always been the target of controversy. Although no association between direct toxicity of aluminum and vaccines has been established, several delayed type hypersensitivity reactions have been reported.¹²⁸⁻¹³⁰ In Denmark, 39 out of 42 children with persistent skin reactions following vaccination had positive patchtests for aluminum.¹³¹In another study, vaccination-induced granulomas and contact allergy to aluminum was reported in 60 out of 63 Swedish children receiving DTP vaccines.¹³²In contrast, an *in vivo* study in a mouse model of peanut allergyfound that the severity of peanut-hypersensitivity was reduced by an alum/CpG-adjuvanted vaccine while exposure to endotoxin and alum did not influence allergic symptoms.¹³³Other novel adjuvants such as MF59, AS03, AF03, which are squalene-based are also used in vaccines.¹³⁴ Although safety concerns were raised due to presence of antibodies to squalene, clinical evidence clearly suggested that squalene is poorly immunogenic and that low titrses of antibodies to squalene are found in healthy individuals. Further, neither the presence of anti-

squalene antibodies nor their titre is significantly increased by immunization with vaccines containing squalene.¹³⁵Details on the mechanism of these and other vaccine adjuvants under clinical investigation are detailed in the review by Shi et al.¹³⁶

Polysorbate 80 is an emulsifier which has beenwidely used to solubilize agents in foods and medicines, including vaccines. This nonionic detergent induces local and systemic allergic reactions, including IgE-mediated and non-immune anaphylaxis. The hydrophilic polymer polyethylene glycol (PEG) is structurally similar to polysorbate 80 and PEG and its derivatives are frequently found in household products including toothpaste, cosmetics, pharmaceuticals and foods. In addition, PEG is often conjugated to biological therapeutics to form a depot agent. It is now well understood that sensitivity to PEG can cause IgE-mediated anaphylaxis after administration of PEG-conjugated biological therapeutics¹³⁷⁻¹⁴², and that severe allergic reactions to PEG have been associated with preexisting anti-PEG antibodies induced by PEG-containing household products.¹⁴³Polysorbate 80 is present in many vaccines and some foods as well. Polysorbates are obtained from PEG moieties but have lower molecular weights and thus may be much less likely to trigger an allergic reaction. PEG may also be cross-reactive with polysorbates, which are contained in some COVID-19 vaccines.^{142,144-} ¹⁴⁶However, measures of pre-existing anti-PEG antibodies vary widely, range from 0.2% to 72% of healthy individuals.¹⁴⁷This has become immediately relevant because PEG 2000, a high-molecular weight version of PEG, is a component in two of the three authorized COVID-19 vaccines. The mRNA BNT162b2 and mRNA-1273 COVID-19 vaccines are lipid nanoparticles containing mRNA that codes for the spike protein in the coronavirus.¹⁴⁸ The lipid nanoparticle delivery system prevents premature degradation of the genetic instructions necessary for individuals to eventually become protected against SARS-CoV-2.149 PEG 2000-lipid is a component of the mRNA-1273 vaccine. The lipid nanoparticles stabilize and improve the aqueous solubility of the two mRNA vaccines, and also act as an adjuvant. While more research is needed to determine the cause of the potentially increasedrate of anaphylaxis to COVID-19 compared to other vaccines, based on the experience with PEG-conjugated biologics, PEG 2000 in the vaccines is considered the most likely culprit.^{139,140,150-155} Therefore, individuals with a known allergy to PEG should be excluded from vaccination with these vaccines for the time being.¹⁵⁶In addition to PEG, other excipients present in COVID-19 vaccines, such as distearoyl phosphatidylcholine, tromethamol, polysorbate 80, and EDTA, should also be

evaluated as potential allergenic components **Table 7** lists the excipients present in the mRNA and ChAdOx1-S vaccines.¹⁵⁷

Section VI: Management of vaccine allergy

Treatment of anaphylaxis in the setting of vaccine administration is reviewed in Sokolowskaet al ¹⁵⁸and Castells et al.¹³⁸ Briefly, due to the possibility of an anaphylactic reaction to the vaccine, any professional administering the vaccine must be capable of managing an anaphylactic reaction and should have the necessary medications and tools on hand. There must be a mandatory observation period after vaccine administration of at least 15 minutes for all individuals, to allow for the administration of adrenaline in an adequate dose.¹⁵⁹Individuals with a suspected allergic reaction to the first dose of the vaccine should be followed up by an allergist so that administration of the second dose can be performed in a specialized setting equipped to treat anaphylaxis. One approach used successfully for many vaccine-allergic individuals, but which has not been evaluated for the COVID-19 vaccines, is to administer the vaccine in incremental doses. Any adverse allergic reactions should be promptly reported including any additional information including individual characteristics.

Section VII: Conclusions

Vaccinations benefit public health and help to reduce the risk of disease across the entire population. Despite early reports of waning of antibody responses over 20 days following SARS-CoV2 infection, evidence is now accumulating that similarly to other infections, recovered patients develop long-lasting immunity.¹⁶⁰Recently, Hartley et al. reported that patients who have recovered from SARS-CoV-2 infection have stable virus-specific memory B cells that recognize the spike or nucleocapsid proteins of the SARS-CoV-2 virus for at least eight months post infection.¹⁶¹Another study found that neutralizing antibody titers against the SARS-CoV-2 spike protein persisted for at least 5 months after infection.¹⁶² These findings support optimism that the currently licensed vaccines will be efficacious despite the emergence of highly infectious mutant viruses and are consistent with the limited reports of natural reinfection after confirmed illnesses. These mRNA vaccines are currently not available for children. Moreover, research by Pfizer and researchers from the University

of Texas have determined that antibodies from 20 recipients of the mRNA BNT162b2vaccine can neutralize the mutant strains *in vitro* (as yet not peer reviewed).¹⁶³

It is crucial that further research is conducted to better understand to which components of the currently available vaccines individuals are reacting and howto identify individuals who may be at risk of an adverse allergicreaction (Box 2). Individuals who may be at risk for an allergic response or who have a history of allergic responses to vaccinations (or their components) should be evaluated by an allergist.⁶⁰

The best mechanism to further our understanding is to study individuals who have had reactions through a variety of *in vitro* experiments and clinical testing. *In vitro* experiments including analysis of plasma IgE markers as well as, basophil and mast cell line activation tests can help to better characterize potential allergens and identifyindividuals atrisk of an anaphylaxis. Clinical testing, including skin prick and intradermal tests can also help identify allergens and at-risk individuals. All of these tests can involve testing the individual components of the vaccine to determine the reaction-inducing antigen. However, the critical and pragmatic evaluation of these tests' results in relationship to particular patient's clinical problems remains crucial.

Increased understanding of vaccine-related allergieswill help to further improve the manufacturing processes and safety of vaccines. Specifically, through identifying specific vaccine components that cause allergic reactions, vaccine manufacturers can either try to remove or create replacements for those components. This also has an impact on the management of vaccine distribution, specifically with regards to ensuring that those who need vaccination will not have an allergic reaction.

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Table 1: Key Characteristics of COVID-19 vaccines(approved and in phase 3 clinical trials)^{7,164}

Developers	Vaccine Name (s)	Vaccine Platform	Authorization Status	Phase 3 Efficacy	Prod. capacity	% of doses to HICs for 2021	Agreement with COVAX?	Dose #	Temp. for storage (Celsius)
Approved vaccines	1	Γ	I	1	1	1	1	1	1
AstraZeneca with	AZD122	Viral	Emergency use in	62%	3bn	27%	Yes	2	2-8
Oxford University	2	vector	U.K., E.U., other						
		(non-	countries.						
		replicating)							
Bharat Biotech	BBV152,	Inactivated	Emergency use in	-	700m	0%	No	2	2-8
International	Covaxin	virus	India.						
Limited									
BioNTech with	BNT162	RNA-based	Approved in	95%	2bn	77%	Yes	2	-70
Pfizer+ Fosum	b2,		several countries.						
Pharma	toziname		Emergency use in						
	ran,		U.S., E.U., other						
	Comirnat		countries.						
	у								
CanSino	Ad5-	Viral	Limited use in	65.7%	320m	0%	No	1	2-8
Biological	nCoV,	vector	China. Emergency						
Inc./Beijing	Convidec	(non-	use in Mexico,						
Institute of	ia	replicating)	Pakistan.						

Biotechnology									
Gamaleya	Sputnik	Viral	Early use in	92%	1bn	0%	No	2	-18
Research Institute	V	vector	Russia.						
and Health		(non-	Emergency use in						
Ministry of the		replicating)	other countries.						
Russian									
Federation									
Moderna +	mRNA-	RNA-based	Approved in	94%	1bn	97%	No	2	-20
National Institute	1273		Switzerland.						
of Allergy and			Emergency use in						
Infectious			U.S., U.K., E.U.,						
Diseases (NIAID)			others.						
Sinopharm with	BBIBP-	Inactivated	Approved in	79%	1bn	8%	No	2	2-8
China National	CorV	virus	China, U.A.E.,						
Biotec Group Co			Bahrain.						
and Beijing			Emergency use in						
Institute of			Egypt, other						
Biological			coutries.						
Products									
Sinopharm and	N/A	Inactivated	Limited use in	-	600m	8%	No	2	2-8
China National		virus	China, U.A.E.						
Biotec Group Co									
and Wuhan									
Institute of									

Biological									
Products									
Sinovac and	CoronaV	Inactivated	Approved in	50-91%	1bn	18%	No	2	Room
Development Co.,	ac	virus	China.						temp.
Ltd			Emergency use in						
			Brazil, other						
			countries.						
Vector Institute	EpiVacC	Protein	Early use in Russia	N/A	11m	-	No	2	2-8
	orona	subunit							
Vaccines in phase 3	trials (unapp	proved)	•	<u> </u>	<u> </u>	1	<u> </u>		
AnGesand Takara	AG0302-	DNA-	Not approved	-	-	-	No	2	-70
Bio and Osaka	COVID1	based							
University	9								
Anhui	ZF2001	Protein	Not approved	-	300m	-	No	2 or 3	2-8
ZhifeiLongcom		subunit							
Biopharmaceutica									
l and Institute of									
Microbiology,									
Chinese Academy									
of Sciences									
Biological E	N/A		Not approved	-	-	-	No	2	2-8
Limited and									
Dynavax and									
Baylor College of									

Chinese A 1		T	Not any 1						
Chinese Academy	IN/A	inactivated	Not approved						
of Medical		virus							
Sciences and									
Institute of									
Medical Biology									
Clover	COVID-	Protein	Not approved	-	1bn	-	No	2	2
Pharmaceuticals	19 S-	subunit							
Inc.,/GSK/Dynava	Trimer								
x									
Covaxx with	UB-612	Protein	Not approved	-	1bn	0%	No	2	2
Nebraska		subunit							
University and									
United									
Biomedical Inc									
CureVacand	CVnCoV	RNA-based	Not approved	-	300m	100%	No	2	5
Bayer									
Inovio	INO-	DNA-	Not approved	-	100m	-	No	2	2
Pharmaceuticals	4800	based							
and International									
Vaccine Institute									
andAdvaccine									
(Suzhou)									
Biopharmaceutica									
(Suzhou) Biopharmaceutica									_

Johnson &	Ad26		Not approved	66%	1bn	38%	Yes	1	2-
Johnson			TT TT						
Medicago, GSK	CoVLP	Virus-like particle	Not approved	-	80m	100%	No	2	2
Novavax	NVX CoV2373	Protein subunit	Not approved	89%	2bn	31%	Yes	2	2
RIBSP	QazCovi d-in®		Not approved	-	60m	-	No	2	2
Research Institute for Biological Safety Problems	QazCovi d	Inactivated virus	Not approved						
Sanofi with GlaxoSmithKline	N/A		Not approved	-	-	73%	Yes	2	2
SII with Max Planck Institute	VPM100 2		Not approved	-	-	-	No	-	-5
SK Biosciences	GBP510		Not approved	-	-	-	No	-	2
University of Hong Kong	N/A		Not approved	-	-	-	No	-	-5
Zydus Cadila	ZyCoV- D	DNA- based	Not approved					3 doses (by skin patch)	

1	Table 2:	Differentiating	features of	vasovagal	episode vs	. anaphylaxis ^{25,26}
		0		0	1	1 0

Vasovagal Episode	Anaphylaxis
On	iset
Immediately or up to 30 minutes after	Can be immediate or within minutes orup
vaccine administration	several hoursafter vaccine administration
Respi	ratory
Normal respiration - may be shallow, but not	Cough, wheeze, stridor, hoarseness,
labored	rhinorrhea
	Signs of respiratory distress (tachypnoea,
	cyanosis, rib recession)
	Upper airway swelling (lip, throat, tongu
4	uvula or larynx)
Cardiov	vascular
Bradycardia – weak/absent peripheral pulse-	Tachycardia, weak/absent central pulse
but with strong central pulse (carotid)	
	Hypotension – sustained and no
Hypotension – usually transient and corrects	improvement without specific treatment
in supine position	infants and young children, limpness and
	pallor are signs of hypotension)
Loss of consciousness – improves once	
supine or head- down position	Loss of consciousness – no improvement
	once supine or in head-down position
Sk	kin
Generalised pallor, cool clammy skin	Urticaria, angioedema, pruritus, erythema
Gastroin	ntestinal
Nausea or vomiting	Abdominal cramps, diarrhea, nausea or
	vomiting
Neuro	logical

Nausea or vomiting	Feels faint, light-headed, headache, dizziness
	blurred vision, restless, seldom: seizures
7	

Goal	Steps
Observed vs. expected (O/E) analysis of	-Collect background incidence rates of AESIs
AESIs during a mass vaccination program	-Create system to process and display real-time
	vaccination data
Routine pharmacovigilance	-Provide AESI standard case definitions
	-Present age-stratified data on AESI incidence
	rate in target population
Follow-up for an adverse reaction	Collect data on:
	-Patient
	-Adverse reaction
	-Vaccination history
4	-Vaccination and diluent (including
1	manufacturer, batch number, batch release
	specifications, expiry date, laboratory test results
5	about the batch)
	-Route of administration
	-Storage and handling conditions

8 Table 3: General pharmacovigilance practices for monitoring vaccine reactions¹⁶⁵

Table 4: Frequency of anaphylaxis after vaccines.

Vaccine	Anaphylaxis	Comment	Reference
	rate per 10 ⁶		
	doses		
Rabies	55.43 - 86.1		McNeil et al. JACI 2016 ³⁶
HPV	1.29 – 26	Different rates according	Brotherton et al. 2008 ¹⁶⁶
		to the type of HPV	Erlewyn-Lajeunesse et al. 2012
		vaccine	McNeil et al. JACI 2016 ³⁶
TBE	20	Polygeline-free TBE	Zent et al. 2003 ¹⁶⁸
		vaccine with lower rate of	
		anaphylaxis	
MMRV	19.8		McNeil et al. JACI 2016 ³⁶
BNT162b COVID-19	11		CDC MMWR 2021 ⁴⁸
mRNA 1273	2.5		CDC MMWR 202145
Varicella	1.2 - 10.3	Gelatin-free varicella	Sakaguchi et al. 2000 ¹⁶⁹
		vaccine with lower rate of	Ozaki et al. 2005 ¹⁷⁰
		anaphylaxis	Su et al. 2016 ³³
Herpes zoster	6.16 - 9.6		McNeil et al. JACI 2016 ³⁶
Pandemic A/H1N1 vaccine	6.8 - 8	Increased risk of	Rouleau et al. 2013 ¹⁷¹
		anaphylaxis compared to	

		seasonal influenza	
		vaccines	
Yellow fever	7.6		Kelso et al. 1999 ¹⁷²
MMR	0.6 - 5.14	Egg allergens no longer	D'Souza et al. 2000 ¹⁷³
		clinically relevant	Pool et al. Pediatrics 2002 ⁹¹
			McNeil et al. JACI 2016 ³⁶
			Su et al. 2016 ³³
MCV4	6.16		McNeil et al. JACI 2016 ³⁶
HAV	3.34		McNeil et al. JACI 2016 ³⁶
DTaP	0.51 - 3.6	During last two decades,	Cheng et al. 2015 ³⁹
		the content of gelatin in	
		this type of vaccine	
		decreased followed by	
		decreased frequency of	
		anaphylaxis	
PPSV23	0.2 - 2.48		McNeil et al. JACI 2016 ³⁶
			Su et al. 2016 ³³
Influenza	0.1 - 1.83	No significant differences	Kawai et al. 2014 ¹⁷⁴
		by types of vaccine or	Ropero-Alvarez et al. 2015 ¹⁷⁵
		manufacturer	McNeil et al. JACI 2017 ³⁶

			2-fold higher rate for	Halsey et al. 2013 ¹⁷⁶
			LAV compared to IIV	Su et al. 2016 ³³
			regarding immediate	
			hypersensitivity reactions	
5	HBV	0-1.67		McNeil et al. JACI 2016 ³⁶
				Duclos. 2003 ¹⁷⁷
	Japanese encephalitis	0-0.26	Anaphylaxis reported in	Li et al. 2014 ¹⁷⁸
			live attenuated vaccine	McNeil et al. JACI 2016 ³⁶
	Hib	0	Very rare event	McNeil et al. JACI 2016 ³⁶
	PCV13	0	Very rare event	McNeil et al. JACI 2016 ³⁶
	Rotavirus vaccines	0	Very rare event	McNeil et al. JACI 2016 ³⁶

BNT162b2 - mRNA vaccine against COVID-19

mRNA 1273 - mRNA vaccine against COVID-19

DTaP - diphtheria-tetanus-acellular pertussis vaccine

HAV – hepatitis A vaccine

Hib – Haemophilus influenza type b vaccine

IIV - inactivated influenza vaccine

LAIV - live attenuated influenza vaccine

MCV4 - 4-valent meningococcal conjugated vaccine

MMR - measles-mumps-rubellavaccine

MMRV- measles-mumps-rubella-varicellavaccine PCV13 – pneumococcalconjugated 13-valent vaccine PPSV23 – 23-valent pneumococcalpolysaccharidevaccine TBE – tick-borne encephalitis vaccine

D	Population	Considerations
	Healthy	A. Follow routine guidelines/schedule for
		vaccination.
		B. Monitor mRNA vaccine recipients for 15-
		30 minutes per local guidelines.
	History of food allergies	Egg: Proceed with vaccination under
		supervision.
(
		Yeast: Seek allergist evaluation prior to
		Hepatitis A, B, HPV, DTaP, Meningococcal,
		Pneumococcal vaccines.
	\prec	
		Gelatin: Seek allergist evaluation prior to
		MMR, Zoster, Influenza, Rabies, Yellow
		fever, Typhoid vaccines.
	History of immunosuppression	A. Defer live vaccination.
D		B. Administer vaccine prior to
		immunosuppression if possible.
	History of vaccine, drug, or antibiotic allergy	A. Refer to allergist.
		B. Identify specific components that may be
		similar in other vaccines.
		C. Graded administration of vaccine/drug after
9		discussion of risks and benefits.
P		

Table 5: Population-specific considerations for vaccination⁴¹⁻⁴⁴

Component Function Vaccine Allergicreactions Stabilizer MMR, Zoster, Influenza Gelatin Anaphylaxis, Rabies, Yellowfever, Typoid Urticaria, Local Albumin Residual Yellowfever, MMR, Influenza, Rabies, Anaphylaxis (eggprotein, médium, Influenza bovine/calf/fet Stabilizer al serum) DTaP, Meningococcal, Hib, Tdap, Casein Medium nutrient Anaphylaxis Influenza Adenovirus Antrax, DTaP, Hib, Aluminum Adjuvant Local Hepatitis A/B, HPV, JapaneseEncephalitisMeningococcalPhe umococcal, Tdap 2-Stabilizer, DTaP, Influenza, Polio, Tdap Local Phenoxyethan Preservative ol Thimerosal Preservative Influenza, Td Local Medium nutrient Hepatitis A, B, HPV, DTaP, Yeast Anaphylaxis Meningococcal, Pneumococcal, Typhoid Natural latex Pharmaceuticalcl Tdap, Meningococcal, Anthrax, Anaphylaxis, Hepatitis A, B, Influenza, DTaP, Urticaria osure Rotavirus, Td, Yellowfever Antimicrobial Polio, DTaP, Hepatitis A, Influenza, Neomycin Anaphylaxis MMR, Rabies, Polio, Smallpox, Varicella, Zoster Polymyxin B Antimicrobial DTaP, Influenza, Polio, Smallpox NA Streptomycin Antimicrobial DTaP, Polio NA

Table 6: Major components, function, and allergic reactions**

	Kanamycin	Antimicrobial	Meningococcal, Influenza	Anaphylaxis
	Gentamicin	Antimicrobial	Influenza	NA
	Amphotericin	Antimicrobial	Rabies	NA
	В			
	Dextran	Stabilizer	MMR*, Rotavirus	Anaphylaxis
		Medium nutrient		
	Formaldehyde	Inactivation of	Polio, DTaP, Hib, Hepatitis B,	Local
		virus,	Influenza, JapaneseEncephalitis,	
		Detoxification of	Meningococcal, Tdap, Thypoid	
		bacterial toxin		
		(Inactivatingagen		
		t)		
Y	Peptone (soy)	Medium nutrient	Pneumococcal	
			Hepatitis B	
	Polysorbate 80	Surfactant	DTaP, Hepatitis A, B, HPV, Influenza,	Non-
			Meningococcal, Pneumococcal,	immunologicalanaph
			Rotavirus, Tdap, Zoster	ylaxis, Local
	Polyethylenegl	Surfactant of	COVID-19	Anaphylaxis
	ycol	mRNA		

*Currently, MMR vaccines containing Dextran not on the market.

** All reasonable efforts have been made to ensure the accuracy of this information, but

manufacturers can change product contents before that information is reflected here. Some major components of vaccines have higher levels in food.

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Table 7: Excipients in the COVID-19 vaccinesmRNA BNT162b2, mRNA-1273, and ChAdOx1-Sandtheirfunctions.

Type of Excipient	Excipient	Vaccine
Lipid nanoparticles: sta	bility and transport of mRNA	
EGylated lipids	((4-hydroxybutyl)azanediyl)bis(hexane-	6 mRNA BNT162b2
	diyl)bis(2-hexyldecanoate) (ALC-0159)	
	1,2-dimyristoyl-rac-glycero-3-	mRNA-1273
	methoxypolyethylene	
	glycol-2000 (PEG2000 DMG)	
hospholipids	1,2-distearoyl-sn-glycero-3-	mRNA BNT162b2,
	phosphocholine (DSPC)	mRNA-1273
ipids	((4-	mRNA BNT162b2
	hydroxybutyl)azanediyl)bis(hexane-	
	6,1-diyl)bis(2-hexyldecanoate) (ALC-	
	0315)	
	lipid SM-102 (patented ionizable lipid)	mRNA-1273
	Cholesterol	mRNA BNT162b2,
		mRNA-1273
Suffer: stability of lipid r	nanoparticles / radical oxidation inhibition	
hosphate buffer	potassium dihydrogen	mRNA BNT162b2
	phosphate/disodium phosphate	
	dihydrate)	
Fromethamol	tromethamol / tromethamol	mRNA-1273
	hydrochloride	
cetic acid/acetate	acetic acid / sodium acetate trihydrate	mRNA-1273
nistidine	L-histidine / L-histidine hydrochloride monohydrate	ChAdOx1-S

	potassium chloride	mRNA BNT162b2
	sodium chloride	mRNA BNT162b2,
		ChAdOx1-S
	magnesium chloride hexahydrate	ChAdOx1-S
	Ethanol	ChAdOx1-S
	disodium edetate dihydrate (EDTA)	ChAdOx1-S
	polysorbate 802	ChAdOx1-S
Thermostabilisation		
	Sucrose	mRNA BNT162b2,
		mRNA-1273
		ChAdOx1-S

Adaptedfrom: Borgsteede S, Tjerk G, Tempels-Pavlica Z, Otherexcipientsthan PEG might cause serioushypersensitivityreactions in COVID-19 vaccines. Allergy. Accepted Jan 2021. In Press., (2021).

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Figure Legends

Figure 1: Vaccination-triggered anaphylaxis rates for major vaccines¹⁹

Figure 2: Types of hypersensitivity reactions

Figure 3: Immunological (IgE-mediated) and non-immunological anaphylaxis

Box 1: Description of common components and contaminants present in vaccine formulations^{43,82,179,180}

Vaccine Component Category	Function
Antigen or its genetic code (DNA, RNA)	Molecules of the pathogen that cause the formation of antibodies and development of specific immune protection (humoral, cellular)
Adjuvants	To stimulate, broaden and optimize immun response
Stabilizers	To keep the vaccine potent and safe during storage and transportation
Preservatives	To prevent contamination
Residual antibiotics	To prevent contamination by bacteria durin the vaccine manufacturing process
Residual cell culture materials	To grow enough of the virus or bacteria to make the vaccine
Residual inactivating ingredients	To kill viruses or inactivate toxins during the manufacturing process
Latex	Found in the vial and syringes used to cont and administer the vaccine. It is a potential contaminant.

BOX 2: Future Safety Research Objectives

COVID-19
-Compare various COVID-19 vaccines to each
other to understand key differences in safety
and efficacy, stratified by patient age and
health status.
-Compare intradermal PEG tests to oral
challenges in evaluating PEG sensitization.

Vaccination 1488 0 recpansion



Vaccine





Figure 3_Nadeau al.

all_14840_f3.jpeg