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# A comprehensive overview of vaccines developed for pandemic viral pathogens over the past two decades including those in clinical trials for the current novel SARS-CoV-2

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The unprecedented coronavirus disease 2019 (COVID-19) is triggered by a novel strain of coronavirus namely, Severe Acute Respiratory Syndrome-Coronavirus-2 (SARS-CoV-2). Researchers are working around the clock to control this pandemic and consequent waves of viral reproduction, through repurposing existing drugs as well as designing new vaccines. Several countries have hastened vaccine design and clinical trials to quickly address this outbreak. Currently, more than 250 aspirants against SARS-CoV-2 are in progress, including mRNA-replicating or non-replicating viral vectored-, DNA-, autologous dendritic cell-based-, and inactivated virus-vaccines. Vaccines work by prompting effector mechanisms such as cells/molecules, which target quickly replicating pathogens and neutralize their toxic constituents. Vaccine-stimulated immune effectors include adjuvant, affinity, avidity, affinity maturation, antibodies, antigen-presenting cells, B lymphocytes, carrier protein, CD4<sup>+</sup> T-helper cells. In this review, we describe updated information on the various vaccines available over the last two decades, along with recent progress in the ongoing battle developing 63 diverse vaccines against SARS-CoV-2. The inspiration of our effort is to convey the current investigation focus on registered clinical trials (as of January 08, 2021) that satisfy the safety and efficacy criteria of international wide vaccine development.

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research at Tamkang University, Taiwan. Then he was employed as a SERB-post-doctoral fellow at the CSIR-Central Leather Research Institute, Chennai, India. He has synthesized substituted alkynylated hybrid molecules and utilized them for NIR and EL applications; and liquid crystalline-, steroidal-based materials for sensor and biomedical applications; he also worked on the characterization of natural product transmycin.



Senthilkumar obtained his M.Sc. chemistry at Bharathidasan University, where he was involved in mechanistic studies of anticancer drugs. He then explored new drug discovery and development from marine and medicinal plants, and their synthesis which is been developed at IIT Madras, IISc Bangalore and IIT Kanpur. He developed variegated drug candidates at IIT Madras, such

as phosphate binder resin, sevcar (two patents); and tolvaptan, hyponatremia and cystagon; and drugs from Terminalia arjuna. He then engaged in the peptidomimetic domain for antitumor screening at IISc Bangalore. Then back at IIT Madras worked on transmycin, which is in phase-II pre-clinical trials. He has also discovered new blood, breast, and liver cancer drugs, and diabetic treatments. Recently, he also discovered, arjunetin, that is a better candidate for COVID-19 treatment than FDA approved drugs.



## Introduction

One of the most successful therapeutic strategies to prevent or control various diseases is by “vaccination” protocol.<sup>1,2</sup> Millions of lives have been saved because of vaccinations, which cover a number of diseases including certain types of cancer, HIV and many viral infections.<sup>3</sup> Vaccination was initially accomplished by Edward Jenner, who was the pioneer for smallpox in the late 18<sup>th</sup> century.<sup>4</sup> In the 1980s, the development of vaccines to fight against pathogenic microorganisms was tentatively introduced. Vaccines are now employed to improve and increase the protection capability (immunity) of the body to fight severe infection and disease,<sup>5</sup> and moreover, primarily intend to make immunity stronger and reduce resistivity of diseases by reducing the reproduction of target pathogens. The majority of vaccines actively exist in the immune system, since anti-bodies are constantly generated in the body to sustain a healthy immunity system.<sup>6,7</sup>

### General components of vaccines

A vaccine consists of an antigen, stabilizer, adjuvant, antibiotic, preservative, and chemical reagents such as formaldehyde.<sup>8</sup> The advantages and disadvantages of vaccinations<sup>9a</sup> are listed in Table 1.

**Antigen.** The component matching the structural array of disease-oriented organisms, wherein, they are identified by the immune system as ‘foreign’ and cause an active immune response.

**Stabilizer.** This module is employed to assist the vaccine by sustaining its efficiency during storage. Instability of the vaccine can lead to reduced antigenicity and decreased infectivity of live attenuated vaccine (LAV). Magnesium chloride (MgCl<sub>2</sub>) for oral polio vaccine (OPV), magnesium sulfate (MgSO<sub>4</sub>) in measles vaccines, and lactose and gelatin associated with sorbitol, are current representatives of stabilizing factors.

**Adjuvant.** They are responsible for enhancing the efficacy of the vaccine by motivating the generation of antibodies.

Chemically, adjuvants are a highly heterogeneous group of compounds (including Al salts).

**Antibiotics.** Used in lower amounts during the development phase, to circumvent bacterial infection during tissue culture cells where the viruses are grown. MMR (measles, mumps, rubella) and IPV (inactivated polio vaccine) associated vaccines have a minimum amount (<25 µg) of neomycin for each dose.

The MMR vaccine was developed by Maurice Hilleman in 1971. Mumps, like measles infections, are caused by an RNA based virus from the Paramyxoviridae family. Moreover, measles and mumps belong to the genus Rubulavirus, it is a human disease with no animal reservoirs. Generally, the MMR vaccine exhibits side effects of a painful arm from the shot, minor rashes, generally in teen/adult women who have no earlier immunity; then the rubella vaccine component can result in joint and tendon stiffness. This vaccine is also associated with the minor threat of seizures/jerking instigated by fever, but is not connected with any enduring effects. The threat of febrile seizures increases as infants get older, hence this vaccine is recommended at a young age. Some people may experience cheek/neck inflammation, impermanent low platelet counts that generally do not require treatment and are also not life threatening.<sup>9b</sup>

In 1955, Jonas Salk initiated an inactivated polio vaccine (IPV), after that, Albert Sabin further developed the live, OPV. Even though poliovirus has three serotypes, both vaccines are trivalent and offer good resistivity against poliomyelitis, a limited number of countries have continuously provided IPV. Sabin's OPV vaccine is underused in most countries due to minimising oral management, but it advances immunity in the intestine, which is capable of spreading to others, and is associated with lower cost.<sup>9c</sup>

**Preservatives.** They are added in multi-dose vaccines, which can be used to control bacterial and fungal growth, including thiomersal (sodium(2-carboxylatophenyl)sulfanyl-ethyl mercury), formaldehyde, or phenolic derivatives. In fact, formaldehyde is responsible for inactivating the viruses (*e.g.* IPV) and removing toxicity in bacterial strains, including the toxin employed in diphtheria and tetanus vaccines.

### Types of vaccines

Various kinds of vaccines are available and those which are administered to infants and adults can be classified (Fig. 1) as follows:

- Live-attenuated.
- Inactivated.
- Toxoid.
- Conjugate.
- Subunit.

**Live attenuated vaccines.** These vaccines are adapted from the existing bacteria or virus, they have been weakened, and thus will not result in serious disease in people with strong immunity. LAV vaccines are more similar to the actual infection. A few representative examples are the MMR and varicella (chickenpox) vaccines. Albeit, it is efficient, but not everyone



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Table 1 Advantages and disadvantages of vaccines

| Advantages                                               | Disadvantages                                                               |
|----------------------------------------------------------|-----------------------------------------------------------------------------|
| Protection of inhabitants against disease                | Uncertainty about complete protection                                       |
| Preventing epidemic and pandemic diseases                | May have some possible side effects                                         |
| Prevents diseases spreading to others                    | Requires NHS/individual outlay                                              |
| Avoids large cost for the treatment of infected patients | Injections are not pleasant/more immune booster injections are inconvenient |

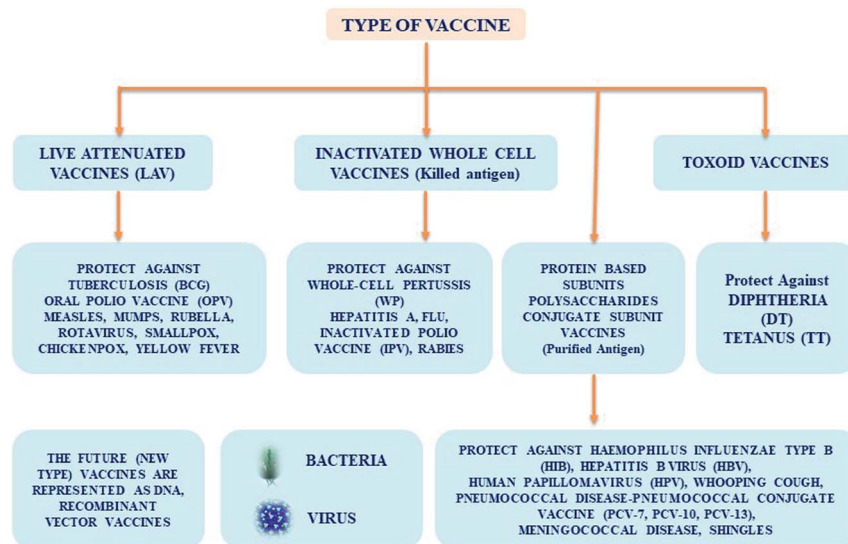


Fig. 1 Pictorial representation of vaccine types.

can be administered these vaccines, including children and patients undergoing chemotherapy as their immune system is too weak.

**Inactivated vaccines.** These vaccines are made from inactivated or dead organisms. The polio vaccine is an example of an inactivated vaccine, which generates an immune response through various routes different to the live attenuated vaccine. It requires a higher dosage to increase and/or sustain immunity.

**Toxoid vaccines.** This vaccine prevents the disease resulting from the toxins released by a virus into the body. Since the toxic substances administered are weak they are unable to cause the illness, while the immune system which encounters this toxoid vaccine, becomes able to repel the natural toxin. The DTaP vaccine for diphtheria and tetanus toxoids falls into this category.

**Conjugate vaccines.** They defend the system against different kinds of bacteria which have antigens that are surface coated with polysaccharide units. It is obvious, that this sugar coated unit masks the antigen. These vaccines append (or conjugate) themselves to the polysaccharide units on the antigens. *Haemophilus influenzae* type B (Hib) vaccine is an example of the conjugate type.

**Subunit vaccines.** These vaccines are made up of simple fractions of the virus/bacteria or subunits, rather than the complete germ. Since the subunit vaccines have an essential antigen only and are not the molecular array constituting the

germ, they have less side effects. The DTaP vaccine-pertussis (whooping cough) module is a paradigm of subunit vaccines.

### Common characteristics of vaccines

Live attenuated vaccines are typically grown in animal cell lines under poor development conditions. The development of an inactivated vaccine involves use of thermal or chemical methods in the beginning, and its mode of action involved in conferring immunity is not fully known. However, the live attenuated or execution of entire organism-supported vaccines have shown a lot of success in the control and inhibition of severe transmittable diseases in human, including animal infectious cattle plague, classic swine fever, equine infectious anaemia, measles, mumps, polio, rubella, smallpox, and so on. In recent times, the use of LAVs, subunit and peptide based vaccines have become possible because of progressive technologies in molecular biology. LAVs are based on the mechanism of action associated with the immune response. Inactivated vaccines, based on antibodies have been mostly used to prevent and manage microbial infectious diseases. LAVs introduce stronger cell immune responses that are decisive to remove several intracellular viral pathogens. However, these pathogens sometimes bypass inactivated vaccines<sup>9d</sup> by mutating peripheral antigens. On the other hand, subunit and peptide based vaccines are less efficient in drawing a strong CD8<sup>+</sup> immune response.



Progressive vaccination involves the use of non-viral distributed nucleic acid-supported vaccines, which imitate live microorganism infection- or immunization. This leads to T-helper cellular immune responses. In addition, this vaccine development<sup>9ef</sup> is harmless and consumes less. It does not require extreme infectious organisms, so is safe from infectivity through live transmittable agents and the discharge of harmful pathogenic organisms. These vaccines fill the gap between a virus outbreak and design of a desirable vaccine, wherein they are classified as DNA/RNA based pentose-carbon sugar motif. The remarkable growth of RNA-associated vaccines resulted in the growth of mRNA based vaccines. It is quite significant to note that mRNA vaccines provide many valuable benefits when compared to viral vectored- and DNA vaccines.<sup>10</sup>

## Proceedings of vaccination against viral diseases

In 1970, to evade the prevalent spread of foot and mouth disease, scientists discovered a vaccine using a single protein from the virus. Despite their achievements in virology, in particular vaccine studies and their development, the lack of understanding of immunological mechanisms of action during induced defensive immunity, has prevented the use of existing vaccines during global pandemic outbreaks of related diseases with similar viral pathogens. The immune system protects against various pathogens, including distinctive units of T-helper cells that are useful to protect against various unusual pathogens. Besides, the follicular T-helper cells (T<sub>FH</sub> cells) generate interleukins (ILs) and support the partition of B cells (they are lymphocytes, and take part in the humoral immune response) and generation of Memory B cells. Furthermore, Memory T cells can be sub classified as CD4<sup>+</sup>/CD8<sup>+</sup> T cells (Cluster of Differentiation) and their functionalities are (a) central memory and (b) effectors memory, which provide various responses upon vaccination against different pathogens.<sup>11</sup> Different vaccines attempted for identical pathogens depend on the perceptions of the scientist,<sup>12</sup> many healthcare professionals do not pay sufficient attention to vaccines, which may result in uncertainty in their efficacy, side effects and toxicity. Table 2 lists the various vaccines discovered that are available in the current market for protection against viral infections. They are listed on the basis of various factors,<sup>13,14</sup> including the apparent protection level, plausible mechanism of action, possibility of usage for other diseases.<sup>15,16</sup>

### Mode of action stimulated by vaccines

Normally antibodies prevent/minimize infections from extracellular pathogens:

- use enzymatic active sites to fuse to toxins to break their diffusion;
- prophylactic action preventing viral replication;
- facilitate opsonophagocytosis of extracellular bacteria;
- inducing the complement cascade.

CD8<sup>+</sup> T cells do not inhibit infection but work to minimize, regulate, and remove intracellular pathogens through:

(a) direct destruction of infected cells (discharge of perforin – a pore forming cytolytic protein present in cytotoxic T lymphocytes (CTLs) and natural killer (NK) cells), granzyme (serine proteases delivered through cytoplasmic granules inside cytotoxic T cells and NK cells);

(b) destroy infected cells with antimicrobial cytokine release.

CD4<sup>+</sup> T cells do not inhibit infection, however they contribute in the minimization, regulation, and refinement of extra- and intra-cellular pathogens with their control and cytokine-development capabilities. The main examples of CD4<sup>+</sup> T cells are:

(a) Follicular T-helper (T<sub>fh</sub>) cells yielding predominantly interleukin (IL)-21 and providing assistance to B-cells;

(b) T-helper 1 (Th1) effector cells yield interferon (IFN)- $\gamma$ , (TNF)- $\alpha$ /TNF- $\beta$ , IL-2, and provide a major role in controlling intracellular pathogens *e.g.*, viruses and bacteria such as *M. tuberculosis*;

(c) Th2 effector cells generate IL-4, IL-5, IL-13, and impact extracellular pathogens such as bacteria and helminths;

(d) Th9 effector cells generate IL-9 and also defend against extracellular pathogens;

(e) Th17 effector cells generate IL-17, IL-22, and IL-26 and participate in mucosal protection (for example against, *S. pneumoniae*, *B. pertussis*, *M. tuberculosis*).

### Main effectors of vaccine responses

Vaccines prevent disease by inducing effector modes of action in cells/molecules to reduce the development of pathogens and deactivate their toxic effects. Vaccine-stimulating immune effectors are resourceful antibodies generated by B lymphocytes' ability to interact to a particular toxin/pathogen. A pictographic representation of vaccine immunological function is shown in Fig. 2.

**Isotype switching.** Control of immunoglobulin (Ig) expression and production from IgM yielding IgG, IgA, or IgE that ensues through B-cell differentiation by DNA recombination.

**Marginal zone.** The zone between the spleen's red pulp and white pulp is known as marginal zone. Its main function is to catch particulate antigens from the circulation and distribute them to lymphocytes.

**Pattern recognition receptors.** These germline-encoded receptors sense the existence of infection through the identification of pathogenic microbe molecular arrays, and stimulate innate immune responses.

**Regulatory T cells-t.** T cells secrete cytokines (IL-10, transforming growth factor [TGF]- $\beta$ /surface markers) and react to reduce immune system response through different modes of action, this sustains immune homeostasis and tolerance to self-antigens.

**Resident memory T cells.** Effector memory T cells exist in particular tissues (such as the lungs, gut, and skin) and are an instant and early line of protection against various viral and bacterial pathogens.

**Somatic hypermutation.** This is a process which intercalates unsystematic mutations in the B-cell receptor region (*i.e.*,



**Table 2** The various antiviral vaccines available in the last two decades, and their characteristics

| Characteristic vaccines |                                            |                                                                                     |                                                                                                                                           |                                                                                                                                                                                                                                                                                                                                                         |                                                                                                                     |                       |
|-------------------------|--------------------------------------------|-------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------|-----------------------|
| S. no.                  | Generic name                               | Trade name                                                                          | Type                                                                                                                                      | Mode of action                                                                                                                                                                                                                                                                                                                                          | Treatment for other diseases                                                                                        | Year of approval Ref. |
| 1                       | AJ vaccines                                | Picovax IPV vaccine, Danish Medicines Agency                                        | Inactivated Polio Vaccine (IPV)                                                                                                           | IPV provides serum immunity to each poliovirus (three types) and defence against any ensuing paralysis causative disease (poliomyelitis). The mucosal immunity level in the intestine is less than that afforded by OPV, this difference may slightly show in the pharyngeal mucosal lining. Prevents poliovirus in the nervous system; immunostimulant | Diphtheria, whooping cough, tuberculosis, tetanus, polio and bladder cancer                                         | 2019 17               |
| 2                       | Grippol plus                               | Quadrivalent AbbVie                                                                 | Polymer supported inactivated influenza vaccine 3-valent, egg derived adjuvanted influenza vaccine, polyoxidoonium as a distinct adjuvant | Vaccine primes advanced meticulous anti-influenza immunity formation; immunostimulant                                                                                                                                                                                                                                                                   | To treat influenza virus infection (A/H1N1 + A/H3N2)                                                                | 2018 18               |
| 3                       | Ys-On-001                                  | Yivyka polyinosinic-polycystidylic acid/ inactivated rabies virus Yisheng Biopharma | Cancer vaccines anti-neoplastics                                                                                                          | B-cell stimulant, cytokines stimulant, dendritic cell stimulant, immune-modulator, macrophage modulator, natural cell stimulant, regulatory T lymphocyte inhibitor, Th1 cell stimulant                                                                                                                                                                  | Cancer vaccines pancreatic cancer                                                                                   | 2017 19               |
| 4                       | Adimmune's quadrivalent flu vaccine (4142) | Adimflu-S (QIS)BioB2B Taiwan                                                        | Monovalent vaccine, egg-based inactivated split virus, influenza H1N1 vaccine                                                             | Immunostimulants; inactivated virus from chick embryo culture split virus                                                                                                                                                                                                                                                                               | Influenza A (H1N1) vaccine during pregnancy                                                                         | 2017 20               |
| 5                       | Quadrivalent influenza vaccine (J0&BB02)   | Vaxigrip Tetra QIVSanofi Pasteur                                                    | Split influenza virus vaccine                                                                                                             | Induces the humeral antibodies against hemagglutinins inhibition (HAI) within 2–3 weeks; antibodies neutralize the influenza viruses                                                                                                                                                                                                                    | Potent immunization against the four influenza virus strains (A and B types have two each)                          | 2017 21               |
| 6                       | NBP-608                                    | SKY Zoster                                                                          | Attenuated zoster-, varicella-type vaccine                                                                                                | Inducing both humoral and cellular immune response, which creates an IgG humoral immune response; varicella-zoster specifically activate CD4 <sup>+</sup> T-helper and CD8 <sup>+</sup> T-lymphocyte cells                                                                                                                                              | Varicella-zoster virus (VZV), a human neurotropic alpha-herpes-virus. Major infection causes varicella (chickenpox) | 2017 22               |
| 7                       | HBV-ISS (Dynavax)                          | Heplisav-B                                                                          | Hepatitis B virus used for vaccine development                                                                                            | Immune-stimulatory DNA sequence (ISS) ISS-1018, adjuvant proceeds TLR-9 agonist, which is used for potential prevention and treatment of HIV infection                                                                                                                                                                                                  | Treatment of HBV-, HIV viruses                                                                                      | 2017 23               |



## Review

Table 2 (Contd.)

| Characteristic vaccines |                                                            |                                                |                                                                                                      |                                                                                                                                                                                                                    |                                                                           |                       |
|-------------------------|------------------------------------------------------------|------------------------------------------------|------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------|-----------------------|
| S. no.                  | Generic name                                               | Trade name                                     | Type                                                                                                 | Mode of action                                                                                                                                                                                                     | Treatment for other diseases                                              | Year of approval Ref. |
| 8                       | GSK-1437173A                                               | Shingrix GSK                                   | Non-live, recombinant subunit vaccine; varicella vaccine herpes zoster (Hz/su) 2q4vaccine candidates | Antigen IgE and adjuvant system AS01B develop VZV-specific immune response in a weakening immune system. Generates a long-term immune response, Shingrix helps to address age-related decline in shingles immunity | To prevent herpes zoster (HZ) infection                                   | 2017 24               |
| 9                       | DTPa-hepB-IPV-Hib                                          | EasySix, Hexaxim Panacea Biotech               | 6 in 1 combination Infanrix hexa whole cell pertussis antigen                                        | The body produces its own antibodies to protect against bacteria and viruses causing different infections                                                                                                          | Diphtheria, DTP, Hepatitis B, Hib, IPV, Pertussis, Polio, Tetanus vaccine | 2017 25               |
| 10                      | Ad5-EBOVJ07BX02                                            | ErveboBIB, CanSinoBIO                          | Glycoprotein recombinant adenovirus type 5 Ebola virus vaccine                                       | Recombinant replication deficient human Ad5 vector stimulating immune responses and defending against the Ebola virus                                                                                              | To treat Ebola virus                                                      | 2017 26               |
| 11                      | Tetavalent inactivated influenza vaccine (TIV)             | Vaxiflu-4 Zydus Cadila                         | Influenza vaccine                                                                                    | To increase immunity by generating antibody proteins, which defend against infection caused by the virus present in the vaccine                                                                                    | Treating influenza viruses-H1N1, H3N2, Type B (Brisbane and Phuket)       | 2017 27               |
| 12                      | Flu Vac Qs 2019–20 (4 Yr Up) CD,BL-125408                  | Flucelvax Quadrivalent Seqirus, Inc            | Cell based flu vaccines                                                                              | Vaccine motivates the body to produce its own immunity (antibodies) against the virus                                                                                                                              | To prevent influenza A and B viruses                                      | 2016 28               |
| 13                      | Quadrivalent influenza vaccine (QIV) (GQM-10)              | Vaxigrip TetraSanofi                           | Inactivated influenza vaccine (split version)                                                        | To enhance the protection from circulating influenza B viruses                                                                                                                                                     | To protect against influenza (flu) viruses                                | 2016 29               |
| 14                      | Live attenuated influenza vaccine (LAIV) J07BB03           | FluMist Quadrivalent Influenza Tetra-Medimmune | Attenuated virus                                                                                     | LAIVs induce T cell antibody-reactions against the surface protein of HA and NA; LAIVs provide hetero-subtype protection in humans                                                                                 | To prevent common flu; influenza A-(H1N1), (H3N2), and B-viruses          | 2016 30               |
| 15                      | Enterovirus type 71 vaccine (Vero cells)                   | Inlive SINOVAC                                 | Inactivated EV 71 virus antigen vaccine                                                              | Generate immune reactions against EV71 virus                                                                                                                                                                       | To prevent hand-foot-mouth disease (HFMD) caused by EV-71                 | 2016 31               |
| 16                      | Inactivated influenza vaccine                              | Cadiflu-S CBL Biologicals Pvt. Ltd             | Inactivated vaccine                                                                                  | To develop immunity against the disease by forming antibodies                                                                                                                                                      | To prevent influenza and protect against its effects                      | 2016 32               |
| 17                      | Inactivated quadrivalent influenza vaccine (split version) | Afluria Quad 2020 Seqiruspty Ltd               | Influenza quadrivalent vaccine 2020                                                                  | Against influenza-A and influenza-B type viruses                                                                                                                                                                   | To prevent influenza A and B viruses                                      | 2016 33               |
| 18                      | Riv-4 (RIV4)                                               | Flublok-QSonofi Pasteur                        | Quadrivalent recombinant influenza vaccine                                                           | Humoral immune response measured by hemagglutination inhibition antibodies                                                                                                                                         | To prevent influenza A (H1N1) and B (H3N2)                                | 2016 34               |
| 19                      | Gam Evac Combi                                             | Combined Vector Based Vaccine                  | Heterologous VSV and adenovirus type-5 vectored Ebola virus                                          | Heterologous prime-boost vaccine humoral immune response, cell facilitated immune response to Ebola                                                                                                                | To prevent Ebola viral disease (EVD)                                      | 2015 35               |



Table 2 (Contd.)

| Characteristic vaccines |                                         |                                               |                                                                  |                                                                                                                                                                                                                  |                                                                                                                                   |                  |      |
|-------------------------|-----------------------------------------|-----------------------------------------------|------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------|------------------|------|
| S. no.                  | Generic name                            | Trade name                                    | Type                                                             | Mode of action                                                                                                                                                                                                   | Treatment for other diseases                                                                                                      | Year of approval | Ref. |
| 20                      | In32 Activated Sabin polio vaccine      | Ai Bi Wei IMB China                           | Inactivated Poliomyelitis (IPV)                                  | Inactivated vaccines offer immunity by delivering an inactivated antigen. This vaccine cannot cause disease, thus, it may be administered to an immuno-compromised host                                          | To prevent polio in new born babies                                                                                               | 2015             | 36   |
| 21                      | Rotavirus Orv-116E                      | ROTAVAC 5D Vero cells Derived Bharath Biotech | Rotavirus vaccine live attenuated monovalent vaccine             | Rotarix (vaccine prevents rotavirus infectivity) reproduces in the small intestine and induces immunity; the particular mechanistic action of immunology by rotarix against rotavirus gastroenteritis is unknown | To prevent rotavirus gastroenteritis                                                                                              | 2015             | 37   |
| 22                      | NBP-607-QIV                             | SKY cell flu Quadrivalent SK Bioscience       | Cell cultured quadrivalent inactivated subunit influenza vaccine | Immunity for influenza viruses A and B subtypes. Sero-protection is generally obtained within three weeks                                                                                                        | To treat influenza virus infections                                                                                               | 2015             | 38   |
| 23                      | GC-3110A, GC-3110B                      | GCflu Quad GC Pharma                          | Quadrivalent influenza virus vaccine                             | Hemagglutination inhibition antibody response                                                                                                                                                                    | To treat influenza virus infections                                                                                               | 2015             | 39   |
| 24                      | Chimerivax-dengue (CYD-TDV)             | Dengvaxia Sanofi Pasteur                      | Live attenuated tetravalent chimeric vaccine                     | Antibody dependent improvement                                                                                                                                                                                   | To prevent Dengue virus-1, 2, 3 and 4 fever in humans                                                                             | 2015             | 40   |
| 25                      | H5N1 influenza (avian flu) vaccine (Rx) | Audenz Vn-101 Sonali Pasteur GSK              | Egg based H5N1 vaccines, inactivated influenza virus vaccine     | Induces immunity (antibodies), which act against viral HA in the vaccine, after interrupting viral attachment to human respiratory cells, provides immunity to influenza A virus subtype H5N1                    | To treat influenza virus                                                                                                          | 2014             | 41   |
| 26                      | 9-Valent HPV vaccine [9v HPV]           | GARDASIL 9                                    | Recombinant virus                                                | Prevent HPV through humoral immune responses induced by the vaccine                                                                                                                                              | Prevention of cervical, vulvar, vaginal, anal, or pharyngeal, head and neck cancer; by preventing papilloma virus (HPV) infection | 2014             | 42   |
| 27                      | Cell cultured-H5N1 vaccine KD-295       | GSK                                           | Emulsion cell culture influenza HA vaccine (Prototype)           | Antibody titer calculated through hemagglutination inhibition (HI): high-titer virus generation led to suspension of growth of MDCK (Madin–Darby Canine Kidney) and Vero cells in a serum-free distribution      | To prevent influenza A virus H5N1                                                                                                 | 2014             | 43   |



## Review

Table 2 (Contd.)

| Characteristic vaccines |                                                                                                                    |                                                                |                                                                                                              |                                                                                                                           |                                                                      |                  |      |
|-------------------------|--------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------|------------------|------|
| S. no.                  | Generic name                                                                                                       | Trade name                                                     | Type                                                                                                         | Mode of action                                                                                                            | Treatment for other diseases                                         | Year of approval | Ref. |
| 28                      | Live modified vaccine virus Ankara Ankara-Bavarian Nordic (MVA-BN)-J07BX                                           | IMVANEX/IMVAMUNE JYNNEOS                                       | Non-replicating small pox vaccine                                                                            | Non-replicating and capable of generating humoral and cellular immune response to orthopox-viruses                        | To prevent smallpox                                                  | 2013             | 44   |
| 29                      | Japanese encephalitis vaccine BBIL/JEV                                                                             | JENVAC Bharath Biotech                                         | Inactivated Vero cell-derived viral vaccine                                                                  | JENVAC is sufficient to elicit an immune response                                                                         | To protect against Japanese encephalitis virus (JEV)                 | 2013             | 45   |
| 30                      | Fluzone quadrivalent BL 103914/J07B B                                                                              | FLUZONE Quadrivalent Sanofi Pasteur Inc.                       | Inactivated quadrivalent influenza virus vaccine type A and type B (split version)                           | Stimulates the production of specific antibodies                                                                          | Prevents influenza diseases, type A and B                            | 2013             | 46   |
| 31                      | FLU-Q-QIV flu laval quadrivalent GSK-2282511A                                                                      | FluLaval™ Quadrivalent Glaxo Smith Kline (GSK)                 | Influenza virus vaccine                                                                                      | Vaccines which improve immunity against the viral pathogen leading to influenza: they induce the generation of antibodies | Prevents influenza diseases, type A and B                            | 2013             | 47   |
| 32                      | DTa-IPV-HePB-HibHexavalent vaccine6-in-1 vaccine J07CA09                                                           | Hexyon, Infanrix, Vaxelis                                      | Hexavalent vaccine                                                                                           | Booster vaccination                                                                                                       | To treat DTaP, hepatitis B, polio, haemophilus influenza diseases    | 2013             | 48   |
| 33                      | GSK-2282512a BL 125127                                                                                             | Fluarix Quadrivalent (FLU Q-QIV) Glaxo Smith Kline Biologicals | Inactivated influenza vaccine, quadrivalent, seasonal                                                        | Increases immunity for treatment of disease originating from influenza-A subtype and type B viruses                       | Prevents disease triggered by influenza A subtype and type B viruses | 2012             | 49   |
| 34                      | ChimeriVax™-JE                                                                                                     | IMOJEV Sanofi                                                  | Live attenuated Japanese encephalitis vaccine (JEV), monovalent                                              | IMOJEV is highly immunogenic and able to induce continuing immunity through both preclinical and clinical trials          | To prevent yellow fever virus                                        | 2012             | 50   |
| 35                      | Prepandemic influenza vaccine (H5N1) J07BB01                                                                       | Vepacel                                                        | Inactivated flu strain known as A/Vietnam/1203/2004 (H5N1) whole virion, derived from inactivated Vero cells | Overall the vaccine primes the immune system                                                                              | Protect against influenza H5N1 (bird flu)                            | 2012             | 51   |
| 36                      | Medi-3250STN: 125020                                                                                               | FLUMIST Quadrivalent Med-Immune                                | Influenza vaccine, live attenuated influenza vaccine (LAIV)                                                  | To provide immunity against influenza virus caused by subtypes A and B                                                    | Protects against influenza                                           | 2012             | 52   |
| 37                      | Hepatitis E hecolin (HEV-239)                                                                                      | Hecolin Xiamen Innovax Biotech                                 | Non enveloped virus with positive sense HEV vaccine, a recombinant vaccine                                   | HEV 239 treatment induces a strong anti-HEG IgG response, through early antibody mobilisation                             | Fights against hepatitis-E virus                                     | 2012             | 53   |
| 38                      | Measles/rubella vaccine                                                                                            | German measles                                                 | Live attenuated (weakened) viruses                                                                           | Immunostimulant, produces antibodies (associated proteins fight and kill measles, mumps, and rubella (MMR) viruses        | Prevent MMR viruses                                                  | 2011             | 54   |
| 39                      | Human inactivated influenza vaccine (H1N1) 2009 vaccine. Pandemic influenza strain A/California/7/2009/nyMC X-179A | HNVAC (Bharath Biotech)                                        | Inactivated Influenza A virus vaccine (H1N1). Cell culture derived vaccine                                   | Active immunization agent, which acts against the influenza A (H1N1) 2009 virus                                           | Activity against influenza A                                         | 2010             | 55   |



Table 2 (Contd.)

| Characteristic vaccines |                                                                                                                                        |                                               |                                                                             |                                                                                                                                                                                                          |                                                                                 |                  |      |
|-------------------------|----------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------------------|-----------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------|------------------|------|
| S. no.                  | Generic name                                                                                                                           | Trade name                                    | Type                                                                        | Mode of action                                                                                                                                                                                           | Treatment for other diseases                                                    | Year of approval | Ref. |
| 40                      | Influenza A virus (H1N1), monovalent vaccine                                                                                           | 2009 Influenza A (H1N1), Sonali Pasteur       | Monovalent vaccine, adjuvant                                                | Active immunization for preventing disease caused by influenza virus A (H5N1)                                                                                                                            | To prevent influenza A viral disease                                            | 2010             | 56   |
| 41                      | Quadrivalent flu vaccine                                                                                                               | FLUCELVAX                                     | Cellular influenza vaccine                                                  | Active immunization for preventing influenza subtypes A and B causative diseases                                                                                                                         | To defend against four different strains of influenza for both subtypes A and B | 2010             | 57   |
| 42                      | Influenza vaccine (whole virion). Inactivated combination                                                                              | Vaxiflu-S Fluzone Zydus Cadila Healthcare Ltd | Inactivated influenza vaccine (NZ)                                          | Active immunization for preventing Vaxiflu-X                                                                                                                                                             | To assist in protection against influenza                                       | 2010             | 58   |
| 43                      | H1N1 pandemic influenza vaccine H5N1 strain of the flu virus A/Vietnam/1203/2004Flu strain/California/07/2009 (H1N1) virus             | Celvapan Baxter International                 | Whole virion in Vero cell-based influenza vaccine, inactivated              | Motivates the immune system to produce antibodies when exposed to the virus                                                                                                                              | To protect against the influenza strains of H5N1 virus                          | 2009             | 59   |
| 44                      | Influenza A virus vaccine H5N1                                                                                                         | Fluval P-H5N1 Omninvest                       | Fluart innovative vaccine                                                   | Fluval affords active immunization against four virus strains; two for influenza A subtype, and two for B type                                                                                           | To prevent influenza a (H5N1)                                                   | 2009             | 60   |
| 45                      | Non adjuvant influenza A (H1N1) 2009 monovalent vaccine. Influenza hemagglutinin (HA) A/California/07/2009 (H1N1) V-like virus         | Panenza Sanofi-Pasteur                        | Non-adjuvanted pandemic vaccine                                             | Induces a high immune (antibody) response, three weeks post-vaccination                                                                                                                                  | To prevent influenza A (H5N1)                                                   | 2009             | 61   |
| 46                      | Monovalent, cell culture-derived, inactivated subunit influenza vaccine. Produced from A/California/07/2009 (H1N1) with adjuvant MF-59 | Celtura                                       | MF-59 adjuvanted cell cultured derivative A/H1N1 pandemic influenza vaccine | MF-59 induces a strong response in adults and substantially develops the response with growth of HA-specific Tfh (CD4 <sup>+</sup> , ICOS <sup>+</sup> , CXCR5 <sup>+</sup> , IL-21 <sup>+</sup> ) cells | Induces an immune response for protection against influenza virus               | 2009             | 62   |
| 47                      | Pandemic (H1N1) ASO3 adjuvanted influenza vaccine                                                                                      | Pandremix GSK                                 | Combination of H1N1 virus antigen and adjuvant system of H1N1               | Enhances the natural immunity of the body                                                                                                                                                                | To prevent influenza A (H1N1), swine flu viral infections                       | 2009             | 63   |
| 48                      | Influenza vaccine (H1N1) flu strain from A/California/7/2009 (H1N1) derived strain NYMC-181 J07BB02                                    | Focetria                                      | Surface antigen (hemagglutinin and neuraminidase) inactivated adjuvant      | Vaccine acts by priming the immune system                                                                                                                                                                | To protect against influenza type A (H1N1) 2009 virus                           | 2009             | 64   |
| 49                      | Cell culture-derived adjuvanted influenza virus vaccine (Grippol TC                                                                    | Grippol NeoSolvay pharmaceutical AbbVie       | Cell based adjuvanted influenza vaccine                                     | Activates the endosomal receptor, which leads non-specific activation of the surface TLRs, which induce the intracellular signals contributing to the antiviral mechanism                                | To prevent influenza virus infections                                           | 2009             | 65   |
| 50                      | Inactivated H5N1 influenza (avian flu) vaccine A/Vietnam/1194/2003/(H5N1) RG                                                           | Pan-flu (Sinovac Biotech)                     | Single shot vaccine against H1N1 influenza                                  | Body reacts by creating antibodies                                                                                                                                                                       | To prevent H5N1 pandemic influenza                                              | 2008             | 66   |



## Review

Table 2 (Contd.)

| Characteristic vaccines |                                                                                                                                                                                                       |                                        |                                                                                    |                                                                                                                                                                                                                                    |                                                |                  |      |
|-------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------|------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------|------------------|------|
| S. no.                  | Generic name                                                                                                                                                                                          | Trade name                             | Type                                                                               | Mode of action                                                                                                                                                                                                                     | Treatment for other diseases                   | Year of approval | Ref. |
| 51                      | Pandemic influenza vaccine                                                                                                                                                                            | Panvax H1N1 vaccine                    | Split virion inactivated vaccine                                                   | Panvax, is an inactive viral part of H1N1, the immune system responds by developing antibodies to the virus particle                                                                                                               | Influenza virus (H1N1), swine flu infections   | 2008             | 67   |
| 52                      | Pre-pandemic influenza vaccine H5N1 (split virion, adjuvanted, inactivated) AS03-H5N1 vaccine A/Vietnam/1194/2004; or A/Indonesia/5/2005 GSK-1562902a                                                 | Prepandrix GSK                         | Split virion, inactivated, adjuvanted; hemagglutinin, antigen, adjuvanted          | Potential immunization against influenza A subtype H5N1 virus                                                                                                                                                                      | Influenza (H5N1) virus, swine flu infection    | 2008             | 68   |
| 53                      | Influenza vaccine (surface antigen, inactivated, prepared in cell cultures) J07BB02                                                                                                                   | Optaflu Flucelvax TETRA Novartis       | Influenza virus surface antigens, hemagglutinin and neuraminidase subunit vaccines | Optaflu comprising different strain surface regions of the flu virus. Vaccine causes immune system to recognize the unknown viral components and produce antibodies against them                                                   | Prevent influenza viral infection              | 2008             | 69   |
| 54                      | Smallpox (vaccinia) vaccine ACAM2000                                                                                                                                                                  | Acam-2000 Sanofi Pasteur Biologics Co. | The vaccine is made from a live virus, vaccinia                                    | Possessing potent immunization against smallpox                                                                                                                                                                                    | Prevention of smallpox                         | 2007             | 70   |
| 55                      | Inactivated quadrivalent influenza vaccine (split virion)                                                                                                                                             | Afluria Seqirus Pty. Ltd.              | Quadrivalent split virion, influenza virus hemagglutinin as the active ingredient. | Immunity against 3 (type A (2) and type B (1)) or 4 (type A (2) and type B (2)) microbial strains. Suitable for the annual flu season                                                                                              | To stop infection caused by influenza virus    | 2007             | 71   |
| 56                      | H5N1 avian flu vaccine, avian influenza or bird flu, vaccine is derived from A/Vietnam/1203/2004 influenza virus                                                                                      | Fluzone Sanofi Pasteur                 | Inactivated influenza virus vaccine                                                | Provides protection against the H5N1 influenza virus stimulating the immune response                                                                                                                                               | To prevent infection caused by influenza virus | 2007             | 72   |
| 57                      | Adjuvanted H5N1 pre-pandemic vaccine                                                                                                                                                                  | Daronix, GSK                           | Second generation pandemic vaccine                                                 | Prepare the body's immune system to prevent a flu epidemic                                                                                                                                                                         | Prevent common flu disease                     | 2007             | 73   |
| 58                      | Influenza vaccine surface antigen, inactivated, prepared in cell culture: A/California/7/2009 (H1N1) pdm09-like strain, A/Switzerland/9715293/2013 (H3N2)-like strain, B/Phuket/3073/2013-like strain | Optaflu                                | Vaccine containing flu surface antigen                                             | Enhances the body's defence system, priming the immune system to make antibodies against the flu virus                                                                                                                             | To prevent infection caused by influenza virus | 2007             | 74   |
| 59                      | Birch pollen allergy vaccine                                                                                                                                                                          | Oralgen Birch Pollen ALK-Abello        | Peptide based vaccine                                                              | Suppression of allergic reactions after immunization with fusion protein, which is caused by releasing the immune messenger interleukin-10 (IL-10), an autologous cytokine, functioning to decrease the overacting immune response | To prevent allergy                             | 2007             | 75   |



Table 2 (Contd.)

| Characteristic vaccines |                                                                                                                                                            |                                                |                                                  |                                                                                                                          |                                                                                                      |                  |      |
|-------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------|--------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------|------------------|------|
| S. no.                  | Generic name                                                                                                                                               | Trade name                                     | Type                                             | Mode of action                                                                                                           | Treatment for other diseases                                                                         | Year of approval | Ref. |
| 60                      | Influenza vaccine, split virion, inactivated hemagglutinin of A/California/7/2009 Hemagglutinin of A/Victoria/210/2009 Hemagglutinin of B/Brisbane/60/2008 | Anflu Sinovac                                  | Split virion, inactivated                        | Body generates immuno-reactions against influenza virus                                                                  | To prevent infection caused by influenza virus                                                       | 2006             | 76   |
| 61                      | Rec hepatitis B vaccine                                                                                                                                    | Supervax                                       | Hepatitis B vaccine, immunoglobulin (HBIG)       | Active immunization of hepatitis B vaccine induces the immune system to create anti-HBs without an active infection risk | To prevent infection against hepatitis B virus                                                       | 2006             | 77   |
| 62                      | Antirabies vaccine                                                                                                                                         | RABIRIX Bharath Biotech                        | Vero cell based rabies vaccine (PVRV)            | Inactivated virus vaccine boosts immunization against rabies                                                             | To prevent infection against rabies virus                                                            | 2006             | 78   |
| 63                      | Live herpes zoster vaccine                                                                                                                                 | Zostavax Merck                                 | Live attenuated virus vaccine                    | Enhancing VZV – specific immunity against zoster                                                                         | Treatment for herpes zoster/postherpetic neuralgia (PHN), a neuropathic pain                         | 2006             | 79   |
| 64                      | Rotavirus vaccine, live rotavirus 116E strain derived in Vero cells                                                                                        | RotaTeq Bharath Biotech                        | Live attenuated, oral, monovalent                | Active immunization for infants from the age of 6 weeks                                                                  | To protect against rotavirus gastro-enteritis disease in infants and young children                  | 2006             | 80   |
| 65                      | Human papillomavirus quadrivalent (type 6, 11, 16, and 18) vaccine (HPV vaccine) J07BM01                                                                   | Gardasil, Cervarix Merck                       | Protein subunit quadrivalent, recombinant        | Body's immune system identifies the viral proteins in Gardasil, develops antibodies against them                         | To protect against either two or four or nine types of HPV (cervical, vaginal and vulvar in females) | 2006             | 81   |
| 66                      | Rotavirus vaccine with five strains of rotavirus, from both human and animal sources                                                                       | ROTARIX                                        | Live, attenuated, oral                           | Usually to develop immunity against rotavirus-based disease                                                              | Rotarix vaccine assists to prevent this disease in children                                          | 2005             | 82   |
| 67                      | Hepatitis B (r DNZ) vaccine, adjuvanted, absorbed J07BC01                                                                                                  | Fendrix GSK                                    | Adjuvanted, absorbed, recombinant DNA technology | Vaccine works by priming the immune system                                                                               | Prevents hepatitis B virus infection                                                                 | 2005             | 83   |
| 68                      | MR vaccine freeze-dried, live, attenuated, measles-rubella combined vaccine                                                                                | Mearubik, Mitsubishi Tanabe Pharma Corporation | Live vaccine for measles, rubella                | Immune system induced to produce antibodies (proteins that fight and also kills the rubella virus)                       | To prevent rubella virus                                                                             | 2005             | 11   |
| 69                      | Inactivated hepatitis A and hepatitis B (rDNA) – HAB adsorbed vaccine                                                                                      | Bilive Sinovac                                 | Recombinant DNA technology                       | Suboptimal immune response to the vaccine                                                                                | Prevents hepatitis B viral infection                                                                 | 2005             | 84   |
| 70                      | Virosomal influenza vaccine, Invivac influenza vaccine                                                                                                     | Invivac Solvay Pharmaceuticals                 | Adjuvant                                         | The virosome mechanism remains complex; it is the transporter as well as an immune stimulant                             | To treat influenza                                                                                   | 2004             | 85   |
| 71                      | Influenza virus (live) (LAIV), inactivated influenza vaccine (IIV)                                                                                         | Flumist                                        | Wild-type                                        | Disease-causing viruses has been attenuated and inactivated, using the influence of heat/chemicals such as formaldehyde  | Protects against infection from influenza viruses                                                    | 2003             | 86   |
| 72                      | Hepatitis A and B Vac DB10989, DB11627                                                                                                                     | Ambirix                                        | Vaccine based                                    | B-lymphocytes anti-HBs antibodies                                                                                        | Immunization against hepatitis A and B viral infectivity                                             | 2003             | 87   |



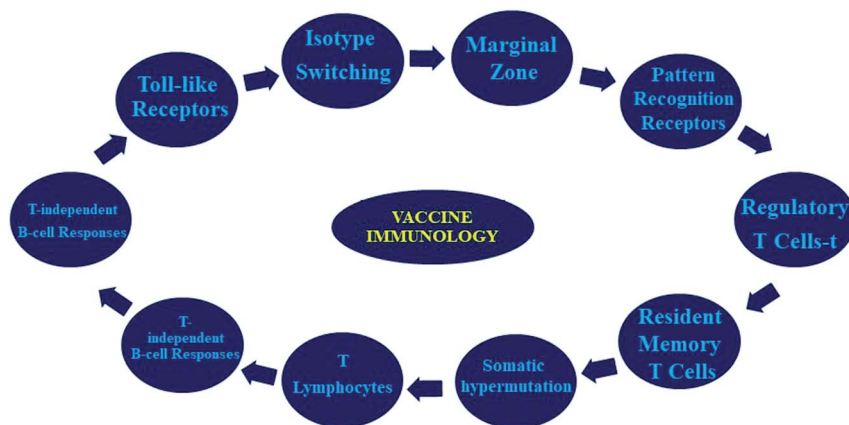


Fig. 2 General features of vaccine immunology.

immunoglobulin) at a particularly high rate throughout the proliferation of B-cells. This mechanistic process occurs as an effect of the cytidine deaminase enzyme and results in antibody diversification.

**T lymphocytes.** Cells that mature in the thymus, and become stimulated in the spleen or nodes if their T-cell receptors interact with an antigen marked with MHC molecules, and they obtain supplementary co-stimulation indications motivating them to kill the infected cell (mainly CD8<sup>+</sup> T cells)/secondary (mainly CD4<sup>+</sup> T cells) roles.

**T-Independent B-cell responses.** This B cell differentiation pathway is mostly triggered by polysaccharides, in the marginal zone and extra-follicular regions of the spleen or nodes. It provides a fast response (days), and generates transient (over months) low affinity antibodies without impacting immune memory.

**T-Dependent B-cell responses.** This B cell differentiation occurs due to protein antigens which recruit T and B cells into germinal centers of the spleen/nodes. It is slower (weeks), but provides enduring stimulation (years) with high affinity antibodies formation and immune memory.

**Toll-like receptors.** A cluster of 10 receptors (TLR1 to TLR10) existing at immune cell external regions, which identify pathogens and trigger characteristic immunity.

The main target of immunization through vaccination is to inhibit specific infections and their unavoidable difficulties. The best vaccine is the one which concomitantly accomplishes the following criteria such as:

- Actively inhibit the infective disease or else minimize the adverse effects of the disease;
- Offer a strong and continuing defence against a specific disease;
- Improves immunity through a minimum quantity of administrations;
- Deliver abundant antigens to afford wide-ranging safety against infection;
- Never results in side effects, or keeps them to a minimum;
- Remains stable under storage conditions, preferably mild storage conditions, for its shelf-life;

- Can be produced on a huge scale;
- Should be economical and easily available.

Herein we consider the COVID-19 pandemic, the main resolution of vaccination against SARS-CoV-2 are:

- Inhibition of characteristic clinical symptoms so hospitalization is avoided, and reduces severe infectivity;
- Prevention of disease spreading before the corresponding antibodies are produced (sero conversion)
- Producing a strong neutralizing immune response able to link with the viral protein spike (S) that must prevent it from attaching to human cells.

From this perspective, the various immunological response effects which neutralize antibodies and CD8<sup>+</sup> T cells are most significant.

The antibodies of anti-SARS-CoV-2 alert the host organism's immune response to the presence of the virus; such antibodies are immunoglobulins, which are appropriately split into IgA, IgM, IgG, and less frequently IgD. Prior serological antibody model responses to viral infections have usually proven the subsequent sequence of these antibodies resulting from virus infection: the antibodies of IgA are primary, which are followed by IgM, IgG-type continues at high levels for a longer time than the preceding ones (IgA and IgM). For certain viruses, sometimes the antigen (the virus itself) co-occurs with antibodies, particularly with the antibodies of primary IgA and IgM. Further, viruses have a "serological window," *i.e.*, a period between the initial arrival of the antigen (in the blood) and the antibody response, thus phase intervals of infection occur. Eventually, IgG-type antibodies are specific to the novel coronavirus (SARS-CoV-2) and can be examined through chemiluminescence immunological routes, which is an automated laboratory process using enzyme-linked immunosorbent assays with higher arrangement. The examination mostly identifies the body's immune response to SARS CoV-2 infection.

### Significant characteristics of (corona) virus

Before 2019, there were two pandemics caused in the past two decades by coronavirus; namely SARS during 2002–2003 and Middle East Respiratory Syndrome (MERS) in 2011. According



to the International Committee on Taxonomy of Viruses (ICTV), the coronaviruses (CoVs) are sub-classified as Othocoronavirinae, which in turn consist of four categories – alpha-CoV, beta-CoV, gamma-CoV, and delta-CoV. The alpha- and beta-CoVs transmit disease in mammals such as bats, pigs, cats and mice. Gamma- and delta-CoVs usually infect birds. In addition, the seven different types of human CoVs – which include HCoV-229E, HCoV-NL63 – belong to alpha-CoVs. The HCoV-OC43, HCoV-HKU1, SARS-, MERS- CoVs, and the current pandemic SARS-CoV-2 belong to beta-CoVs, and the genus of such beta coronavirus occurrences are zoonotic infections. The December 2019 outbreak coronavirus, which leads to the respiratory-associated syndrome, originated from Wuhan, China, and is called the novel corona virus disease 2019 or nCOVID-19, and its genome is fully sequenced.<sup>88</sup> The genetic sequential arrangement of SARS-CoV-2 has an identical genomic array of SARS-/MERS-CoV.<sup>89,90</sup>

Initially, the taxonomy of coronaviruses are split in to three groups, on account of genetic and serological interactions, the first set (Group-1) comprised several viruses including the porcine epidemic diarrhoea virus (PEDV), porcine transmissible gastroenteritis virus (TGEV), canine coronavirus (CCoV), feline infectious peritonitis virus (FIPV), the previously identified coronavirus of HCoV-229E, and HCoV-NL63. Whilst the second combination (Group-2) consists of murine hepatitis virus (MHV), bovine coronavirus (BCoV), human coronavirus OC43 (HCoV-OC43), rat sialodacryoadenitis virus (SDAV), porcine hemagglutinating encephalomyelitis virus (PHEV), canine respiratory coronavirus (CRCoV), and equine coronavirus (ECoV). In a similar pattern, the third group (Group-3) contains the avian infectious bronchitis virus (IBV) and Turkey coronavirus (TCoV). Now, the SARS coronaviruses (SARS-CoV) cannot be associated with any of these representative groups, however they possess some similarities along with the second group coronaviruses.<sup>91a</sup>

The battle between scientists and viral infections is a perpetual process and thus the identification of specific potential drugs with high efficiency and low toxicity is a continuous aim. Globally, this is the first time, that scientific researchers, diplomats, politicians, and capitalists have convened to work towards a common objective. The FDA approved drugs chloroquine and hydroxychloroquine to be utilized in critical illness cases, but clinical practices were still becoming overwhelmed by CoV cases. The suggestion to employ extensive use of these antiviral drugs was not sufficient. Certain polymerase nucleoside/nucleotide inhibitors are promising agents. Favipiravir a selective viral RdRp inhibitor, has been tested in clinical trials against COVID-19. Furthermore, the antiviral drugs lopinavir, ritonavir, remdesivir, nelfinavir, serine based protease inhibitors of nafamostat, camostat, efficient lipid reducing statin, rosuvastatin, TNF alpha inhibitors, interleukin 1 receptor antagonists; Janus associated kinases (JAK) as well as monoclonal antibodies, tocilizumab, baricitinib and ruxolitinib *etc.*, along with their combinations and other antiviral components, are also under investigation in clinical studies to combat COVID-19. In the future, researchers are requested to accumulate their results to provide more

knowledge to repurpose significant drugs appropriately, and provide cheap drugs with the minimum toxicity profile.<sup>91b,c</sup>

### The manufacture of SARS-CoV-2 vaccines through various approaches

Vaccines have to be approved based on sufficient evidence. Many scientific researchers have reviewed the current literature and worked towards the development of a specific treatment for COVID-19. Current studies have exposed several beneficial opportunities, a few of them are more established and are in preclinical trials in addition to some in clinical trials.<sup>92,93</sup> Vaccines endeavor to represent the antibody to the antigen, and they should help the immune system (innate and adaptive immune responses, Fig. 3).

### Virus vaccines

A number of vaccines are being developed for SARS-CoV-2, including inactivated and weakened, replicating and non-replicating viral vectors, and the tentative use of nucleic acid in the form of DNA and RNA. Now, we can distinguish each vaccine characteristic as follows:

**Weakened virus.** Viruses cause disease by reproducing in a rapid manner. A weakened virus reproduces inside the human or animal host very poorly thereby decreasing its virulence and disease causing ability. One pharmaceutical company in the USA (Codagenix, New York) is collaborating with an Indian research team (Serum Institute at Pune) for the manufacture of SARS-CoV-2 with changed genetic sequence – making viral proteins with lower potency.

**Inactivated virus.** Chemicals such as formaldehyde, or thermal heating, are applied to inactivate a virus which is then used as a vaccine.

**Viral-vector vaccines.** In this technique other virus types are modified to render them safe and then used as a vaccine. They combine the qualities of DNA vaccines with those of live attenuated vaccines. The viral vector vaccine consists of a live attenuated virus which is genetically modified to carry the DNA encoding protein antigens from a different organism. For example, the virus belonging to measles/adenovirus is genetically modified and thus made safe. Currently there are products for veterinary use but not for human use. There are two types of viral vectors,<sup>94</sup> one is still able to replicate inside the cells, while the other is non-replicating, since the key genes have been rendered inoperative.

*Replicating viral vector (weakened measles).* One of the approved Ebola vaccines is a typical paradigm for a viral-vector vaccine. Even if replication occurs inside cells, this vaccine is safe and will intensify the immune response.

*Non-replicating viral vector (adenovirus variety).* In general, viral vectors are genetically transformed to generate defective replications which are termed non-replicating vectors. Numerous viruses like adenovirus, adeno-linked, measles, and human parainfluenza viruses has been extensively utilized as viral vectors. Ultimately, the virus attains an attenuated state in which they can activate the anticipated human immune



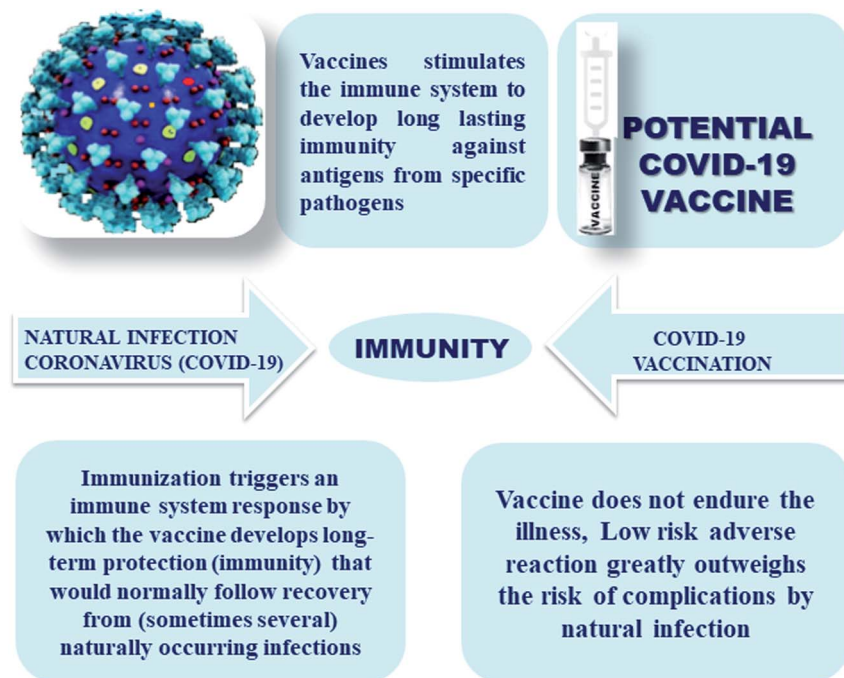


Fig. 3 Pictorial representation of characteristic immunity.

responses, however, they are unable to reproduce in human cells.

**Nucleic-acid vaccines.** A number of groups are aspiring to utilize gene sequences (based on DNA or RNA) for a coronavirus protein to instantaneously provide an immune response. The majority of these vaccines focus on the spike protein present in the virus.<sup>95</sup>

**Protein-based vaccines.** Several researchers have focused on injecting proteins present in coronavirus into the body. Also, protein fragments or shells of protein as copies of the coronavirus' surface could also be utilized.

**Protein subunits.** Nearly 28t groups are working on vaccines using subunits of viral proteins, predominantly on the spike protein at the receptor binding arena. Vaccines for protecting monkeys against SARS virus infection are available which have not been examined for humans. In fact, to employ such vaccines, adjuvant-immune-stimulation drug molecules should be administered together with the vaccine.<sup>96</sup>

**Virus-like particles.** Shells (outer regions) of an empty virus resemble the CoV structure, however, it is non-transmittable due to the absence of the genetic factors. Several research groups are engaged in vaccines based on virus-like particles that can activate a good immune response, however it is quite complex to synthesize such particles, and 70% of the teams directing their research in this way are from industry or private companies rather than academic laboratories.<sup>97,98</sup>

Table 3 and 4 provide a comprehensive overview of the current development of vaccines for SARS-CoV-2, their mode of action, their utility and the name of the manufacturer.<sup>99</sup> More than 10 vaccines have almost reached the approval stage, while many are in the preclinical stage. Fig. 4 portrays the number of vaccine aspirants which are under clinical (~63 drug aspirants) and preclinical (~172 drug aspirants) stages against nCOVID-19.<sup>99a</sup> Protein-based techniques seem to be the most popular among the various mechanisms (see <https://www.who.int/publications/m/item/draft-landscape-of-covid-19-candidate-vaccines>).

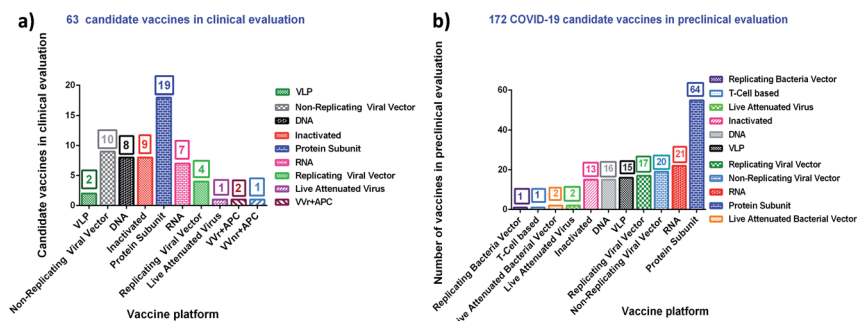


Fig. 4 nCOVID-19 vaccine aspirants at the clinical (63 drug aspirants) and preclinical (172 drug aspirants) stage.

Table 3 COVID-19 aspirant developers and their significant details<sup>99a</sup>

| S. no. | Vaccine aspirant developer                                                                                             | Vaccine aspirant category                                                                                                                          | Clinical trials <sup>a</sup> |            |          |              |           |
|--------|------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------|------------|----------|--------------|-----------|
|        |                                                                                                                        |                                                                                                                                                    | Phase I                      | Phase I/II | Phase II | Phase II/III | Phase III |
| 1      | Sinovac Research and Development Co., Ltd.                                                                             | SARS-CoV-2 vaccine (Inactivated)                                                                                                                   |                              |            |          |              | ■         |
| 2      | Sinopharm + China National Biotech Group Co + Wuhan Institute of Biological Products                                   | Inactivated SARS-CoV-2 vaccine (Vero cell)                                                                                                         |                              |            |          |              | ■         |
| 3      | Sinopharm + China National Biotech Group Co + Beijing Institute of Biological Products                                 | Inactivated SARS-CoV-2 vaccine (Vero cell)                                                                                                         |                              |            |          |              | ■         |
| 4      | AstraZeneca + University of Oxford                                                                                     | ChAdOx1-S (AZD1222) (Covishield)                                                                                                                   |                              |            |          |              | ■         |
| 5      | CanSino Biological Inc./Beijing Institute of Biotechnology                                                             | Recombinant novel coronavirus vaccine (Adenovirus type 5 vector)                                                                                   |                              |            |          |              | ■         |
| 6      | Gamaleya Research Institute/Health Ministry of the Russian Federation                                                  | Gam-COVID-Vac Adeno-based (rAd5-S/Ad5-S)                                                                                                           |                              |            |          |              | ■         |
| 7      | Janssen Pharmaceutical                                                                                                 | Ad26.COV2.S                                                                                                                                        |                              |            |          |              | ■         |
| 8      | Novartis                                                                                                               | SARS-CoV-2 rMMatrix-MJ-Adjuvant (full length recombinant SARS-CoV-2 glycoprotein nanoparticle vaccine adjuvanted with Matrix M)                    |                              |            |          |              | ■         |
| 9      | Moderna + National Institute of Allergy and Infectious Diseases (NIAID)                                                | mRNA-1273                                                                                                                                          |                              |            |          |              | ■         |
| 10     | Pfizer/BioNTech + Fosun Pharma                                                                                         | BNT162 (3 LNP-mRNA)                                                                                                                                |                              |            |          |              | ■         |
| 11     | Akzo/ Zhihi Longmen Biopharmaceutical + Institute of Microbiology, Chinese Academy of Sciences                         | Recombinant SARS-CoV-2 vaccine (CHO Cell)                                                                                                          |                              |            |          |              | ■         |
| 12     | CureVac AG                                                                                                             | CVnCoV vaccine                                                                                                                                     |                              |            |          |              | ■         |
| 13     | Institute of Medical Biology + Chinese Academy of Medical Sciences                                                     | SARS-CoV-2 vaccine (Vero cell)                                                                                                                     |                              |            |          |              | ■         |
| 14     | Research Institute for Biological Safety Problems, Rop of Kazakhstan                                                   | QazCovid-in® COVID-19 inactivated vaccine                                                                                                          |                              |            |          |              | ■         |
| 15     | Inovio Pharmaceuticals + International Vaccine Institute + Advaccine (Barbados) Biopharmaceutical Co., Ltd.            | INO-4800 + electroporation                                                                                                                         |                              |            |          |              | ■         |
| 16     | AsGen + Takara Bio + Osaka University                                                                                  | AG001-COVID-19                                                                                                                                     |                              |            |          |              | ■         |
| 17     | Cadila Healthcare Ltd.                                                                                                 | nCoV vaccine                                                                                                                                       |                              |            |          |              | ■         |
| 18     | Genesee Consortium                                                                                                     | GX-19                                                                                                                                              |                              |            |          |              | ■         |
| 19     | Bharat Biotech International Limited                                                                                   | Whole-Viron Inactivated SARS-CoV-2 Vaccine (BBV152)                                                                                                |                              |            |          |              | ■         |
| 20     | Kentucky Bioprocessing Inc.                                                                                            | KBP-COVID-19 (RBD-based)                                                                                                                           |                              |            |          |              | ■         |
| 21     | Sandoz Partner + GSK                                                                                                   | SARS-CoV-2 vaccine formulation 1 with adjuvant 1 (S protein (BioSolvation production))                                                             |                              |            |          |              | ■         |
| 22     | Astara Therapeutics                                                                                                    | ARCT-021                                                                                                                                           |                              |            |          |              | ■         |
| 23     | Serum Institute of India + Accugen Pty                                                                                 | RBD SARS-CoV-2 H10Aq VLP vaccine                                                                                                                   |                              |            |          |              | ■         |
| 24     | Shenzhen Kangxin Biological Products Co., Ltd.                                                                         | Inactivated SARS-CoV-2 vaccine (Vero cell)                                                                                                         |                              |            |          |              | ■         |
| 25     | BuTera + Leucocyte + Universells                                                                                       | GRACOV2 (replication defective Simian Adenovirus (GRAC) encoding S)                                                                                |                              |            |          |              | ■         |
| 26     | Vaxart                                                                                                                 | VXA-CoV2-1: Ad5 adjuvanted oral vaccine platform                                                                                                   |                              |            |          |              | ■         |
| 27     | University of Munich (Ludwig-Maximilians)                                                                              | MVA-SARS-2-S                                                                                                                                       |                              |            |          |              | ■         |
| 28     | Choror Biopharmaceuticals Inc./GSK/Dynavax                                                                             | SCB-2019 + A003 or CpG 1018 adjuvant plus adjuvant (active like intranasal adjuvant spike protein vaccine)                                         |                              |            |          |              | ■         |
| 29     | Vaccine Pty Ltd + Medigen                                                                                              | COVID-19 vaccine                                                                                                                                   |                              |            |          |              | ■         |
| 30     | CSL Ltd + Seqirus + University of Queensland                                                                           | MF59 adjuvanted SARS-CoV-2 Schimp vaccine                                                                                                          |                              |            |          |              | ■         |
| 31     | Moderna Vaccine Biologics + Dynavax + National Institute of Allergy and Infectious Diseases (NIAID)                    | MVC-COV190 (S-P protein + CpG 1018)                                                                                                                |                              |            |          |              | ■         |
| 32     | Instituto Finlay de Vacunas                                                                                            | FINLAY-FR anti-SARS-CoV-2 Vaccine (RBD + adjuvant)                                                                                                 |                              |            |          |              | ■         |
| 33     | Federal Budgetary Research Institution State Research Center of Virology and Biotechnology "Vector"                    | EpVacCorona (EpVacCorona vaccine based on peptide antigens for the prevention of COVID-19)                                                         |                              |            |          |              | ■         |
| 34     | West China Hospital + Sichuan University                                                                               | RBD (BioSolvation production) expressed in SP cells recombinant SARS-CoV-2 vaccine (SP cell)                                                       |                              |            |          |              | ■         |
| 35     | University Hospital Tuebingen                                                                                          | BMP-GVacc1 (SARS-CoV-2 HLA-DR peptide)                                                                                                             |                              |            |          |              | ■         |
| 36     | COVAXX + United Biomedical Inc.                                                                                        | UB-612 (multipeptide based SI-RBD-protein based vaccine)                                                                                           |                              |            |          |              | ■         |
| 37     | March & Co. + Thermo + Sharp & Duane + Institute Pasteur + University of Pittsburgh                                    | V591-601 – Membrane-anchored (TMV-c91)                                                                                                             |                              |            |          |              | ■         |
| 38     | Jiangsu Provincial Center for Disease Prevention and Control                                                           | DN2S1-2019-CoV-RBD-OP1 (Intranasal RBD-based RBD)                                                                                                  |                              |            |          |              | ■         |
| 39     | Synovate Corporation                                                                                                   | hucTL-Spike                                                                                                                                        |                              |            |          |              | ■         |
| 40     | ImmunityBio, Inc.                                                                                                      | BAE5-S-Fusion + N-TSD vaccine                                                                                                                      |                              |            |          |              | ■         |
| 41     | City of Hope Medical Center + National Cancer Institute                                                                | COB401 (MVA-SARS-2-S)                                                                                                                              |                              |            |          |              | ■         |
| 42     | Israel Institute for Biological Research                                                                               | AVIV-SARS-CoV-2-S vaccine                                                                                                                          |                              |            |          |              | ■         |
| 43     | Akzo Biomedical, Inc., National Institute of Health Research and Development, Ministry of Health Republic of Indonesia | Dendritic cell vaccine AV-COVID-19-A vaccine consisting of autologous dendritic cells loaded with antigens from SARS-CoV-2, with or without GM-CSF |                              |            |          |              | ■         |
| 44     | Codagen/Seum Institute of India                                                                                        | COVI-VAC                                                                                                                                           |                              |            |          |              | ■         |
| 45     | Center for Genetic Engineering and Biotechnology (CEGB)                                                                | CIGB-609 (RBD + AgalIB)                                                                                                                            |                              |            |          |              | ■         |
| 46     | Center for Genetic Engineering and Biotechnology (CEGB)                                                                | CIGB-66 (RBD + aluminum hydroxide)                                                                                                                 |                              |            |          |              | ■         |
| 47     | Valvira, National Institute for Health Research, United Kingdom                                                        | VLA2001                                                                                                                                            |                              |            |          |              | ■         |
| 48     | Biological E Limited                                                                                                   | BECOV2                                                                                                                                             |                              |            |          |              | ■         |
| 49     | Celld Co., Ltd.                                                                                                        | ARCLD-CoV19                                                                                                                                        |                              |            |          |              | ■         |
| 50     | GeneBio Life Sciences, Inc.                                                                                            | GLS-510                                                                                                                                            |                              |            |          |              | ■         |
| 51     | Nanogen Pharmaceutical Biotechnology                                                                                   | Recombinant Sars-CoV-2 spike protein, aluminum adjuvanted                                                                                          |                              |            |          |              | ■         |
| 52     | Shangri                                                                                                                | Recombinant protein vaccine S-200019 (using BioSolvation expression vector system)                                                                 |                              |            |          |              | ■         |
| 53     | Altimmune, Inc.                                                                                                        | ARCoVIR – Adenovirus-based “platform” expresses the receptor-binding domain (RBD) of the SARS-CoV-2 spike protein                                  |                              |            |          |              | ■         |
| 54     | University Medical Center Groningen + Altimmune Biotech Inc.                                                           | SARS-CoV-2-RBD-Fc fusion protein                                                                                                                   |                              |            |          |              | ■         |
| 55     | Enceus University                                                                                                      | ERICOV-VAC, inactivated virus                                                                                                                      |                              |            |          |              | ■         |

<sup>a</sup> Vaccine candidate list and relevant information shown according to the World Health Organization (WHO) statistical data for COVID-19 during 2019–2020. Clinical research data (last updated on January 08, 2021) may be variable.

## Comparison between previous vaccines and COVID-19 vaccines

Comparing COVID-19 vaccine novelty may seem challenging, but it is normal for queries to arise on a vaccine's overall potential, safety, toxicity and its side effects *etc.*,<sup>100a</sup> and some of the efficacy, efficiency and safety of the COVID-19 frontrunner vaccines can be determined by comparison with other vaccines, like flu vaccines. For example, Pfizer/BioNTech released their COVID-19 vaccine initially in the UK and USA, and the rest of the world could see their effectiveness. However, prior to this demonstration the three prominent COVID-19 vaccines – Pfizer/BioNTech claims an efficacy of 95%, the Oxford/AstraZeneca provides 70%, and the Moderna is stated as 94.1% efficacy – were compared with other available vaccines such as the flu, polio, and measles, which helped provide their effectiveness and efficiency predictions.

It is important to discuss here that “effectiveness” and “efficacy” are different.<sup>100b</sup> Efficacy states how a vaccine performs in ideal lab circumstances, like those present in clinical trials. Whereas, effectiveness means how the vaccine works in normal, non-controlled conditions. For example, in a clinical trial, 90% efficacy refers to 90% lower disease rates in the group getting the vaccine compared with the sample group. But, the members in a group selected for a clinical trial have to be in good health and young and they usually have no underlying health conditions. Besides, medical researchers will not generally consider some demographics in these clinical research studies like children/pregnant women. Thus, once a vaccine is able to prevent disease in clinical trials, we may observe the effectiveness drop when directed to different demographics.

Vaccines do not necessarily need high effectiveness to protect several thousands of lives from disease. For instance, the vaccine for flu<sup>100c</sup> has 40–60% effectiveness according to CDC data. During 2018–19, this vaccine prevented millions of influenza cases and its associated illness, but determining the exact effectiveness rate is challenging. Dosages can also increase effectiveness for some vaccines. The two doses of a vaccine can give a protection boost, nevertheless this advantage is sometimes limited to only certain groups like children/organ transplanted people. The booster dosage may not provide an advantage in people aged 65 years.

Through comparing vaccines, like the ones for polio and measles, we see heavy dosages are needed to realize effectiveness.

Polio vaccines<sup>100d</sup> should be up to 100% effective. According to the CDC, “Two inactivated polio vaccine (IPV) dosages have 90% effectiveness; three dosages are 99–100% effective.” The IPV vaccine prevents poliomyelitis (poliovirus), which can activate infection in the brain and spinal cord leading to paralysis.

The MMR vaccine<sup>100e</sup> defends against measles, mumps, and rubella, which tends to have up to 97% effectiveness at inhibiting measles once directed in two dosages. A single dose is around 93% effective, as reported by the CDC. They suggest to give the initial dose at “12–15 months of age, followed by the second dose at 4–6 years.”

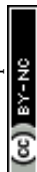


Table 4 Range of SARS-CoV-2 vaccine aspirants and their representative data<sup>16a,b,99c</sup>

| Vaccine ID | Vaccine division acronym | Vaccine sector description     | Vaccine category and composition                                                                            | Characteristic nature of vaccination                                                                                                                                                                 | Producers                                                           | Clinical phases                                                                                | Vaccine utility for other viruses                        | Ref. |
|------------|--------------------------|--------------------------------|-------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------|------------------------------------------------------------------------------------------------|----------------------------------------------------------|------|
| 1          | IV                       | Inactivated virus              | Vaccines of CoronaVac, PiCoVaccine, based on vaccine cultured in Vero cells, inactivated pathogen           | Based on an inactivated pathogen, body generates a varied immune response against several viral antigens, producing neutralizing antibodies                                                          | Sinovac R&D Co. Ltd                                                 | Phase III NCT04456595, Phase 1/2 NCT04383574, NCT04352608                                      | Vaccine being developed for treatment of SARS-CoV-2      | 101  |
| 2          | IV                       | Inactivated virus              | Inactivated vaccine, Vero cell based                                                                        | Vaccine from non-living viral particles, bacteria, and other pathogens which are developed in a culture medium. No potential for infection, but induces an immune system response                    | Sinopharm + WIBP                                                    | Phase 3, Phase-I/II/III ChiCTR2000031809                                                       | Vaccine being developed for treatment of SARS-CoV-2      | 102  |
| 3          | IV                       | Inactivated virus              | Inactivated and similar to virus vaccine based on Vero cells                                                | The foremost vaccine, does not exhibit any adverse side effects with favored immunogenicity and safety; also an inactivated new crown vaccine which completely neutralizes the antibodies in 28 days | Sinopharm + BIBP                                                    | Phase 3 ChiCTR2000032459                                                                       | Vaccine being developed for treatment of similar viruses | 92d  |
| 4          | VVnr                     | Viral vector (non-replicating) | Covishield ChAdOx1-S-(AZD1222)                                                                              | Adenovirus vector based on chimpanzee adenovirus                                                                                                                                                     | AstraZeneca + Oxford University                                     | Phase 3 ISRCTN89951424, Phase2b/3 2020-001228-32, Phase 1/2 PACTR2020069221651322020-001072-15 | MERS, influenza, TB, Chikungunya, Zika, MenB, plague     | 103  |
| 5          | VVnr                     | Viral vector (non-replicating) | Ad5-nCoV recombinant vaccine for CoV                                                                        | Recombinant adenovirus type 5 vector based vaccine aspirant, which is genetically modified with replication-deficient groups, mimics SARS-CoV-2 spike protein                                        | Beijing Biotechnology Institute/CanSino Biological Inc.             | Phase 2 ChiCTR2000031781, Phase 1 ChiCTR2000030906                                             | EBOV (Ebola virus)                                       | 104  |
| 6          | VVnr                     | Viral vector (non-replicating) | Gam-COVID-Vac adeno-based (rad26-S+rAd5-S), Russian COVID-19 vaccine, adenovirus based, and non-replicating | Develop immunity against the coronavirus, and strengthens the immune system                                                                                                                          | Gamaleya Research Institute; Ministry of Health, Russian Federation | Phase-I/III NCT04436471, NCT04437875                                                           | Vaccine being developed for treatment of SARS-CoV-2      | 105  |
| 7          | VVnr                     | Viral vector (non-replicating) | Ad26.COVS2 recombinant serotype 26, adenovirus vectors, multivalent vaccine                                 | Activates specific acquired immunity for Ebola virus                                                                                                                                                 | Janssen Pharmaceutical                                              | Phase 3 NCT04505722, NCT04614948                                                               | Vaccine being developed for treatment of similar viruses | 106  |



Table 4 (Contd.)

| Vaccine division ID | Vaccine sector acronym | Vaccine category and composition | Characteristic nature of vaccination                                                                                       | Producers                                                | Clinical phases                                               | Vaccine utility for other viruses                            | Ref. |
|---------------------|------------------------|----------------------------------|----------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------|---------------------------------------------------------------|--------------------------------------------------------------|------|
| 8                   | PS                     | Protein subunit                  | SARS-CoV-2 rS/Matrix M1-adjuvant (full size recombinant SARS CoV-2 GP nanoparticle based vaccine with Matrix Mas adjuvant) | Novavax                                                  | Phase 3 NCT04611802, NCT04583995                              | Ebola, Lassa, MERS, Nipah, Rift Valley Fever and Chikungunya | 107  |
| 9                   | RNA                    | RNA based vaccine                | Moderna mRNA-1273 vaccine, LNP encapsulated cell bank; mRNA, VAX (non-replicating viral vector)                            | Moderna + NIAID                                          | Phase 3 NCT04470427, Phase 2 NCT04405076, Phase 1 NCT04283461 | Multiple agents                                              | 108  |
| 10                  | RNA                    | mRNA based vaccine               | BNT162 (3 LNP-mRNAs)                                                                                                       | BioNTech + Fosun Pharma; Jiangsu Provincial CDC + Pfizer | Phase 2/3 NCT04368728                                         | To be implemented                                            | 109  |
| 11                  | PS                     | Protein subunit                  | Recombinant COVID-19 vaccine, CHO (Chinese hamster ovary) cell system, prevents CoVs                                       | Anhui Zhifei Longcom Biopharmaceutical + IMCAS           | Phase-I NCT04453852, NCT04445194, Phase 3 NCT04646590         | Vaccine being developed for treatment of SARS-CoV-2          | 110  |
| 12                  | RNA                    | mRNA based vaccine               | CVnCoV vaccine                                                                                                             | CureVac AG                                               | NCT04449276 Phase 1, NCT04515147 Phase 2, NCT04674189 Phase 3 | CureVac's vaccine candidate against SARS-CoV-2               | 111  |
| 13                  | IV                     | Inactivated virus                | Vaccine for SARS-CoV-2 based on Vero cells                                                                                 | IMB + CAMS                                               | Phase 3 NCT04659239, Phase 1/2 NCT04470609, NCT04412538       | To be implemented                                            | 112  |
| 14                  | IV                     | Inactivated virus                | QazCovid-in@-COVID-19 inactivated vaccine                                                                                  | RIBSP, Republic of Kazakhstan                            | Phase 1/2 NCT04530357                                         | Vaccine being developed for treatment of SARS-CoV-2          | 113  |



Table 4 (Contd.)

| Vaccine ID | Vaccine division acronym | Vaccine sector description | Vaccine category and composition                                                                     | Characteristic nature of vaccination                                                                                                                                                                                                                                                                                        | Producers                             | Clinical phases                                                                   | Vaccine utility for other viruses                           | Ref. |
|------------|--------------------------|----------------------------|------------------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------|-----------------------------------------------------------------------------------|-------------------------------------------------------------|------|
| 15         | DNA                      | DNA based vaccine          | INO-4800 + electroporation vaccine with DNA-plasmid based connecting electroporation device          | DNA plasmids distribute by electro-permeabilization (electro-transfer), a computationally sequenced design to produce a specific immune response                                                                                                                                                                            | Inovio Pharmaceuticals + IIVI         | Phase 1 NCT04336410, NCT04447781, Phase 2 ChiCTR2000040146, Phase 2/3 NCT04642638 | Multiple agents, Cancer, HBV, HIV, HPV, Lassa, Nipah, Zika, | 114  |
| 16         | DNA                      | DNA based vaccine          | AG0301-COVID19                                                                                       | Plasmid DNA vaccine developed through an intradermal gene transfer targeting the S protein, which increases the efficiency of gene expression and antibody production                                                                                                                                                       | AnGes + Takara Bio + Osaka University | Phase 2/3 NCT04655625, Phase 1/2 NCT04463472, NCT04527081                         | To be implemented                                           | 115  |
| 17         | DNA                      | DNA based vaccine          | nCov vaccine viral vector, membrane protein based vaccine                                            | Progression of DNA vaccine with the viral membrane protein liable for CoV cell entry; plasmid DNA incorporated into the host cell changes the viral protein, leading to a strong immune response intervened by cellular and humoral immunity. Also used to produce live attenuated recombinant measles viral vector vaccine | Cadila Healthcare Ltd.                | Phase 1/2 CTRI/2020/07/026352                                                     | To be implemented                                           | 116  |
| 18         | DNA                      | DNA based vaccine          | GX-19                                                                                                | Designed to make antigens by adding nucleic acids into the body, the antigens then produce an immune response                                                                                                                                                                                                               | Genexine Consortium                   | Phase-1/2 NCT04445389                                                             | To be implemented                                           | 117  |
| 19         | IV                       | Inactivated virus          | Whole-virion inactivated vaccine (BBV152), Covaxin for SARS-CoV-2, India's trials on nCoV-19 vaccine | Isolated from asymptomatic COVID-19 patient at NIV Pune, India                                                                                                                                                                                                                                                              | Bharat Biotech International Limited  | Phase 1/2 NCT04471519, Phase 3 NCT04641481; CTRI/2020/11/028976                   | Vaccine being developed for treatment of SARS-CoV-2         | 118  |
| 20         | PS                       | Protein subunit            | KBP-COVID-19 (RBD-based), plant based technology                                                     | A distinctive plant-based vaccine technology for the generation of antigens to various diseases                                                                                                                                                                                                                             | KBP., Inc.                            | Phase 1/2 NCT04473690                                                             | To be implemented                                           | 119  |



Table 4 (Contd.)

| Vaccine ID | Vaccine division acronym | Vaccine sector description                                | Vaccine category and composition                                                                             | Characteristic nature of vaccination                                                                                                                                                                                              | Producers                              | Clinical phases               | Vaccine utility for other viruses                   | Ref. |
|------------|--------------------------|-----------------------------------------------------------|--------------------------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------|-------------------------------|-----------------------------------------------------|------|
| 21         | PS                       | Protein subunit                                           | Adjuvanted vaccine of SARS-CoV-2 formulation spike protein (baculovirus production)                          | ARCT-021                                                                                                                                                                                                                          | Sanofi Pasteur + Glaxo SmithKline plc. | Phase 1/2 NCT04537208         |                                                     | 120  |
| 22         | RNA                      | RNA based vaccine                                         | RBD SARS-CoV-2 HBsAg VLP vaccine                                                                             | Powerful single dose, vaccine is based upon self-replicating mRNA, self-transcribing and replicating RNA along with its liquid-enabling and resolving nucleomonomer facilitator altered RNA                                       | Arcturus Therapeutics                  | Phase 1/2 NCT0448095          | Vaccine being developed for treatment of SARS-CoV-2 | 121  |
| 23         | VLP                      | Virus like particle                                       | SARS-CoV-2 vaccine                                                                                           | Neutralizing antibodies response in pigs, treated together with pseudo type lentivirus and live SARS-CoV-2 viruses. Immunoglobulin (IgG) response observed after booster immunization                                             | SII + Accelagen Pty                    | Phase 1/2 ACTRN12620000817943 | Vaccine being developed for treatment of SARS-CoV-2 | 122  |
| 24         | VVnr + APC               | Viral vector (non-replicating) + APC IV inactivated virus | SARS-CoV-2 vaccine (inactivated)                                                                             |                                                                                                                                                                                                                                   | Shenzhen Geno-Immune Medical Institute | Phase 1/2 NCT04276896         | Vaccine for SARS-CoV-2                              | 123  |
| 25         | VVnr                     | Viral vector (non-replicating)                            | GRAd-COV2 replication using defective simian adenoviral vector (GRAd), encodes full length CoV spike protein | Able to generate immune response (antibodies and T cells)                                                                                                                                                                         | ReiThera + Leukocare + Univercells     | Phase 2 ChiCTR2000039462      | Ebola and RSV (respiratory syncytial virus)         | 124  |
| 26         | VVnr                     | Viral vector (non-replicating)                            | VXA-CoV2-1 Ad5 adjuvanted oral vaccine platform                                                              | Vaccine mainly provides mucosal immunity, this vital factor targets mucosal pathogens, including the current coronavirus. Stimulates antigen-specific CD4 <sup>+</sup> and CD8 <sup>+</sup> T cells at low and high dosage levels | Vaxart                                 | Phase 1 NCT04563702           | Influenza infection H1N1                            | 125  |



Table 4 (Contd.)

| Vaccine division ID | Vaccine sector acronym | Vaccine category description   | Vaccine category and composition                                                                                | Characteristic nature of vaccination                                                                                                                                                                                                                                   | Producers                                            | Clinical phases                                                                                    | Vaccine utility for other viruses                     | Ref. |
|---------------------|------------------------|--------------------------------|-----------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------|----------------------------------------------------------------------------------------------------|-------------------------------------------------------|------|
| 27                  | VVnr                   | Viral vector (non-replicating) | MVA-SARS-2-S                                                                                                    | Immune responses and neutralizing antibodies directing the S antigen used to defend against infectivity. The double recombinant sMVA-CoV-2 vectors give S-specific CD8 <sup>+</sup> and CD4 <sup>+</sup> T cells to both S and N antigens in Balb/c mice, respectively | University of Munich (Ludwig-Maximilians)            | Phase 1 NCT04569383                                                                                | Infectious diseases and cancer                        | 126  |
| 28                  | PS                     | Protein subunit                | SCB-2019 + AS03/CpG 1018 adjuvant plus alum adjuvant (similar to native trimeric subunit spike protein vaccine) | S-Trimer, antigen used to target the antibodies of novel CoV spike protein and is in ACE2-viable human convalescent sera samples, obtained from recovered COVID-19 patients                                                                                            | Clover Biopharmaceuticals Inc./GSK/Dynavax           | Phase 1 NCT04405908                                                                                | Vaccine being developed for treatment of SARS-CoV-2   | 127  |
| 29                  | PS                     | Protein subunit                | COVID19 vaccine                                                                                                 | Effective T cells and neutralizing antibody response                                                                                                                                                                                                                   | Vaxine Pty Ltd. + Medytox                            | Phase 1 NCT04453852                                                                                | Vaccine being developed for treatment of SARS-CoV-2   | 128  |
|                     |                        | Protein subunit                | MF59 adjuvanted SARS-CoV-2 S clamp vaccine                                                                      | Advanced antibodies are able to neutralize infectivity through live virus in the cell culture                                                                                                                                                                          | CSL Ltd. + Seqirus + University of Queensland        | Phase 1, development was suspended and the candidate vaccine was removed from the summary analysis | Vaccine production against similar viruses is on hold |      |
| 30                  | PS                     | Protein subunit                | MVC-COV1901 (S-2P protein + CpG 1018)                                                                           | High titer of neutralizing antibodies are prompted against pseudo type novel CoV in sera of immunized mice                                                                                                                                                             | Medigen Vaccine Biologics Corp + NIAID + Dynavax     | Phase 1 NCT04487210                                                                                | Vaccine being developed for treatment of SARS-CoV-2   | 129  |
| 31                  | PS                     | Protein subunit                | FINLAY-FR anti-SARS-CoV-2 vaccine (RBD + adjuvant)                                                              | It has the capability to generate a substantial immune reaction                                                                                                                                                                                                        | Instituto Finlay de Vacunas                          | Phase 2 RPCEC00000347, Phase 1/2 RPCEC00000332, Phase 1 RPCEC00000338                              | Vaccine being developed for treatment of SARS-CoV-2   | 130  |
| 32                  | PS                     | Protein subunit                | EpiVacCorona, based on peptide antigens                                                                         | This antigens-based preparation stimulates the immune reaction                                                                                                                                                                                                         | Federal Budgetary Research Institution SRC VB VECTOR | Phase 1/2 NCT04527575                                                                              | Vaccine being developed for treatment of SARS-CoV-2   | 131  |
| 33                  | PS                     | Protein subunit                | RBD, baculovirus production expressed in Sf9 cells, recombinant SARS-CoV-2 vaccine (Sf9 cell)                   | Specific immune blotting RBD, S-WT and S-2P, motivates high neutralization titers                                                                                                                                                                                      | WCH, Sichuan University                              | Phase 2 NCT04640402, ChiCTR2000039994, Phase 1 NCT04530656                                         | Vaccine being developed for treatment of SARS-CoV-2   | 132  |
| 34                  | PS                     | Protein subunit                | IMP CoVac-1 (SARS-CoV-2 HLA-DR peptides)                                                                        |                                                                                                                                                                                                                                                                        | University Hospital Tuebingen (UKT)                  | Phase 1                                                                                            | Vaccine being developed for treatment of SARS-CoV-2   | 133  |



Table 4 (Contd.)

| Vaccine division ID acronym | Vaccine sector description       | Vaccine category and composition                                                                                            | Characteristic nature of vaccination                                                                                                                                                                                                                                        | Producers                                                                       | Clinical phases                                    | Vaccine utility for other viruses                        | Ref. |
|-----------------------------|----------------------------------|-----------------------------------------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------|----------------------------------------------------|----------------------------------------------------------|------|
| 35 PS                       | Protein subunit                  | UB-612, multipeptide based S1-RBD-protein based vaccine                                                                     | This vaccine has the immunity potential to reduce future pandemic rates                                                                                                                                                                                                     | COVAXX + United Biomedical Inc                                                  | Phase 1 NCT04545749                                | Vaccine being developed for treatment of SARS-CoV-2      | 134  |
| 36 VVr                      | Viral vector (replicating)       | V591-001-measles-vector based (TMV-o38)                                                                                     | Attains the target immune response in humans                                                                                                                                                                                                                                | Merck & Co., Themis + Sharp & Dohme + Pasteur Institute + Pittsburgh University | Phase 1 CT04498247, Phase/2 NCT04497298NCT04569786 | Vaccine being developed for treatment of SARS-CoV-2      | 135  |
| 37 VVr                      | Viral vector (replicating)       | DelNS1-2019-nCoV-RBD-OPT1, intranasal flu-based-RBD                                                                         | SARS-CoV-2 RBD protein based vaccine used to motivate cross-reactivity or cross-neutralizing antibodies; moreover it blocks previous CoV pseudovirus and the advanced novel CoV pseudovirus into hACE2 expressing 293 T cells ( $IC_{50} = 4.1$ and $11.63 \mu g ml^{-1}$ ) | Jiangsu Provincial CDC                                                          | Phase 2 ChiCTR2000039715, Phase 2 ChiCTR2000037782 | Vaccine being developed for treatment of SARS-CoV-2      | 136  |
| 38 RNA                      | RNA based vaccine                | LNP-nCoVsaRNA                                                                                                               | The saRNA vaccine can activate a strong immune response; it is suggested that the IM vaccination antigen is expressed in muscle cells, then moves to antigen presenting cells (APC), representing a cross priming mode of potentiality to prominent $CD8^+$ T cells         | Imperial College London                                                         | Phase 1 ISRCTN17072692                             | Vaccine being developed for treatment of SARS-CoV-2      | 137  |
| 39 RNA                      | RNA based vaccine                | SARS-CoV-2 mRNA vaccine                                                                                                     | —                                                                                                                                                                                                                                                                           | Shulan (Hangzhou) Hospital and CDC at Guangxi Zhuang Autonomous Region          | Phase 1                                            | Vaccine being developed for treatment of SARS-CoV-2      | 138  |
| 40 VLP                      | Virus-like particle              | Coronavirus-like particle COVID-19 (CoVLP)                                                                                  | Provides immunity by generating a harmless spike protein member, reduces severe infection effects                                                                                                                                                                           | Medicago Inc.                                                                   | Phase 2/3                                          | Vaccine being developed for treatment of similar viruses | 139  |
| 41 VVr + APC                | Viral vector (replicating) + APC | COVID-19/aAPC vaccine produced from a variation of lentivirus with immune modulatory genes and viral minigenes to the aAPCs | Artificial antigen-pathogen-specific vaccine, which uses the spike protein binding to the ACE2 receptor; the vaccine utilizes the modified minigenes to direct viral proteins, vary aAPC, and trigger T-cells                                                               | Shenzhen Genoimmune Medical Institute                                           | Phase 1 NCT04299724                                | Vaccine being developed for treatment of SARS-CoV-2      | 140  |



Table 4 (Contd.)

| Vaccine ID acronym | Vaccine division | Vaccine sector description           | Vaccine category and composition                                                                                                                                                                                                       | Characteristic nature of vaccination                                                                                                                                                                                                               | Producers                    | Clinical phases       | Vaccine utility for other viruses                        | Ref. |
|--------------------|------------------|--------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------|-----------------------|----------------------------------------------------------|------|
| 42                 | VVnr + APC       | Viral vector (non-replicating) + APC | LV-SMENP-DC vaccine. Dendritic cells are modified with lentivirus vectors transmitting COVID-19 minigene SMENP and immune modulatory genes. Cytotoxic T lymphocytes (CTLs) are activated by LV-DC offering COVID-19 specific antigens. | Vaccine directed with antigen-specific CTLs                                                                                                                                                                                                        | Same medical institute       | Phase 1/2 NCT04276896 | Vaccine being developed for treatment of similar viruses | 141  |
| 43                 | PS               | Protein subunit                      | AdimrSC-2f (recombinant RBD $\pm$ aluminium)                                                                                                                                                                                           | —                                                                                                                                                                                                                                                  | Adimmune Corporation         | Phase 1 NCT04522089   | Vaccine being developed for treatment of SARS-CoV-2      | 142  |
| 44                 | DNA              | DNA based vaccine                    | Covigenix VAX-001                                                                                                                                                                                                                      | Induces neutralizing antibody levels and stable T helper cell immunity                                                                                                                                                                             | Entos Pharmaceuticals Inc.   | Phase 1 NCT04591184   | Vaccine being developed for treatment of SARS-CoV-2      | 143  |
| 45                 | DNA              | DNA based vaccine                    | CORVax                                                                                                                                                                                                                                 | CORVax12 initiates a coordinated vaccine response, able to expose the innate adaptive humoral and cellular arms. These cellular immune responses have the potential to generate a strong antiviral response                                        | Providence Health & Services | Phase 1 NCT04627675   | Vaccine being developed for treatment of similar viruses | 144  |
| 46                 | RNA              | RNA based vaccine                    | ChulaCov19 mRNA vaccine                                                                                                                                                                                                                | The mRNA based vaccine encodes a protein antigen, while RNA is considered to be unstable; the design and development of this novel vaccine is improving its constancy and protein translation efficacy, so it efficiently enhances immune response | Chulalongkorn University     | Phase 1 NCT04566276   | Vaccine being developed for treatment of similar viruses | 145  |



Table 4 (Contd.)

| Vaccine ID | Vaccine division acronym | Vaccine sector description       | Vaccine category and composition                                                                                                | Characteristic nature of vaccination                                                                                                                                                                                                    | Producers                            | Clinical phases                               | Vaccine utility for other viruses                        | Ref. |
|------------|--------------------------|----------------------------------|---------------------------------------------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------|-----------------------------------------------|----------------------------------------------------------|------|
| 47         | DNA                      | DNA based vaccine                | bacTRL-Spike                                                                                                                    | Symvivo's bacTRL gene therapy platform associates an advanced gene-expression plasmid with a probiotic bacterium to resolve restrictions and distribute the DNA vaccine directly to the gut. The bacterium triggers the immune response | Symvivo Corporation                  | Phase 1 NCT04334980                           | Vaccine being developed for treatment of similar viruses | 146  |
| 48         | VVnr                     | Viral vector (non-replicating)   | hAd5-S-Fusion + N-ETSD vaccine                                                                                                  | A viral spike protein. Virus enters the host cells through the ACE2 receptor with a fusion linker, S is antigenic, and generates an efficient immune response                                                                           | ImmunityBio, Inc.                    | Phase 1 NCT04591717, NCT04710303, NCT04732468 | Vaccine being developed for treatment of similar viruses | 147  |
| 49         | VVnr                     | Viral vector (non-replicating)   | COH04S1 (MVA-SARS-2-S)                                                                                                          | Vaccine holds the SARS-CoV-2 spike and nucleocapsid proteins inserted into the MVA platform that can replicate DNA within cells. Thus it generates novel CoV protein expression to trigger host immunity against the virus              | City of Hope National Medical Center | Phase 1 NCT04639466                           | Vaccine in development                                   | 148  |
| 50         | VVr                      | Viral vector (replicating)       | rVSV-SARS-CoV-2-S vaccine                                                                                                       | rVSV-ΔG-spike stimulated a safe, efficient and adequate neutralizing antibody. Vaccination leads to lower morbidity, protects lungs, and provides fast viral clearance                                                                  | IIBR, Israel                         | Phase 1/2 NCT04608305                         | Vaccine in development                                   | 149  |
| 51         | VVr + APC                | Viral vector (replicating) + APC | Dendritic cell vaccine AV-COVID-19: contains autologous dendritic cells load with antigens from SARS-CoV-2, with/without GM-CSF | Produced from isolated peripheral blood monocytes from patients. Monocytes are then distinguished into dendritic cells with GM-CSF and IL=4                                                                                             | Aivita Biomedical, Inc.              | Phase 1/2 NCT04690387 NCT04386252             | Vaccine in development                                   | 150  |



Table 4 (Contd.)

| Vaccine division ID acronym | Vaccine sector description | Vaccine category and composition    | Characteristic nature of vaccination                                                                                                                                                                                                                                                      | Producers                          | Clinical phases               | Vaccine utility for other viruses | Ref. |
|-----------------------------|----------------------------|-------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------|-------------------------------|-----------------------------------|------|
| 52 LAV                      | Live attenuated virus      | COVI-VAC                            | Codagenix's synthetic attenuated virus engineering (SAVE) platform utilizes synthetic biology to re-code the virus genes into the vaccine. Since this design involves distributing a benign, live attenuated version of SARS-CoV-2 it may stimulate a strong and enduring immune response | Codagenix/Serum Institute of India | Phase 1 NCT04619628           | Vaccine in development            | 151  |
| 53 PS                       | Protein subunit            | CIGB-669 (RBD + AgnHB)              | Stimulates the generation of antibodies that enable a strong immune response to the pathogen                                                                                                                                                                                              | CIGB                               | Phase 1/2 RPCEC00000345       | Vaccine in development            | 130  |
| 54 PS                       | Protein subunit            | CIGB-66 (RBD + aluminium hydroxide) | Stimulates the generation of antibodies that boost the targeted immune response to the virus                                                                                                                                                                                              | CIGB                               | Phase 1/2 RPCEC00000346       | Vaccine in development            | 130  |
| 55 IV                       | Inactivated virus          | VLA2001                             | Vero-cell supported the refined inactivated candidate, based on Valneva's JE vaccine, uses the spike protein normal structural array with CpG 1018 and can stimulate an immune response with a high titer of neutralizing antibodies                                                      | Valneva, NIHR, United Kingdom      | Phase 1/2 NCT04671017         | Vaccine in development            | 130  |
| 56 PS                       | Protein subunit            | BECOV2                              | Candidate based on inactivated SARS-CoV-2 virus components                                                                                                                                                                                                                                | Biological E Limited               | Phase 1/2 CTRI/2020/11/029032 | Vaccine in development            | 130  |
| 57 VVr                      | Viral vector (replicating) | AdCLD-CoV19                         | Immunotherapeutic vaccine based on cells                                                                                                                                                                                                                                                  | Cellid Co., Ltd.                   | Phase 1/2 NCT04666012         | Vaccine in development            | 152  |
| 58 DNA                      | DNA based vaccine          | GLS-5310                            | Nucleic acid-based vaccine platform                                                                                                                                                                                                                                                       | GeneOne Life Science, Inc.         | Phase 1/2 NCT04673149         | Vaccine in development            | 152  |



Table 4 (Contd.)

| Vaccine division ID acronym | Vaccine sector description | Vaccine category and composition                                                     | Characteristic nature of vaccination                                                                                                                                                                                                                                                                                                                         | Producers                            | Clinical phases          | Vaccine utility for other viruses | Ref. |
|-----------------------------|----------------------------|--------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------|--------------------------|-----------------------------------|------|
| 59 PS                       | Protein subunit            | Recombinant SARS-CoV-2 spike protein, aluminum adjuvanted                            | Protein subunit containing the recombinant SARS-CoV-2 S1 domain of the spike protein, also integrates either CoVaccine HT™ or alhydrogel. CoVaccine HT™, a single dose stimulated high titers of antigen-binding IgG. This accelerates the affinity maturation and switching of class to higher values, enhancing cell-induced immunity and virus antibodies | Nanogen Pharmaceutical Biotechnology | Phase 1/2 NCT04683484    | Vaccine in development            | 153  |
| 60 PS                       | Protein subunit            | Recombinant protein vaccine S-268019 (based on Baculovirus expression vector system) | Vaccine is based on recombinant protein units together with GSK; development of mRNA vaccine by Sanofi, in collaboration with Translate Bio. Preclinical data revealed that the two immunizations of the mRNA based vaccine stimulated high neutralizing antibody levels equal to those produced in infected humans                                          | Shionogi                             | Phase 1/2 jRCT2051200092 | Vaccine in development            | 130  |

<sup>a</sup> Globally, vaccine producers have the ability to achieve rapid development of highly efficient, safe – minimum toxicity – vaccines.

In order to determine the similarities and differences between previously available vaccines and COVID-19 vaccines, the Bacillus Calmette–Guérin (BCG) vaccine against tuberculosis is provided as one more example; it confers a wide range of immunity against other infections, and it also may minimize the intensity of COVID-19. Moreover one epidemiological analyses provided universal connections between the vaccinations of BCG and COVID-19 mortality: the suggestion of BCG vaccination results on COVID-19 fatality are dominated by socio-economical and demographical variations between countries. In the wake of reducing the manifold distracting factors, many substantial connections between the BCG vaccination and decreased COVID-19 fatalities were perceived. Obviously this investigation emphasizes the necessity for an intrinsic mechanism of studies supporting BCG vaccination

effects on COVID-19, and also for clinical evaluation to control the COVID-19 pandemic.<sup>99b</sup>

COVID-19 is a serious respiratory related disease, so the scientific and medicinal community are working hard across the globe to develop a vaccine. Presently, around sixty vaccine candidates are on trial in many countries. Now, nearing twenty candidate vaccines are in phase 3 clinical trials. Gratifyingly, seven vaccines have been approved in many countries.

## Conclusions

Globally, as of January 2021, there have been around 88 million COVID-19 cases, including nearly 1.9 million fatalities, WHO have registered (see WHO Covid-19 case report).<sup>99d</sup> These numbers are expected to increase further, so there is an



emergency requirement to produce vaccines to protect people. Several candidates vaccines are being developed which are in pre-clinical and clinical trials. The mechanism of action of these candidates varies significantly, as better knowledge becomes available<sup>154a</sup> about this virus, researchers can adapt their design so if one candidate shows low efficacy, another one may be more active.

Similarly, different demographics may necessitate the need for designing vaccines with different mechanisms of action. The time required for the various stages of clinical trials needs to be shortened (without compromising on the ethics and safety) to achieve the desired goal in a short period of time.<sup>154b</sup> Finally, manufacturing such large quantities of the vaccine or vaccines, quickly (without compromising on the quality, purity and efficacy) and distributing them to all parts of the world is another problem which the present planners have not faced before. This requires sufficient manufacturing capacity, availability of raw materials, logistics and several other factors. Also, as all resources are diverted towards SARS-CoV-2, epidemiologists and public health organizations should not lose track of their fight against other viruses.<sup>155</sup>

The current assemblage of vaccine developers<sup>156a</sup> may reward researchers with increased capability, as numerous basic, transformational and preclinical statistical data have become available during coronavirus exploration. These factors combine together as a substantial promising source for rapid vaccine development.<sup>156b</sup>

Since, December 2020, some new vaccines have emerged and also have been approved by certain national regulatory authorities for use against COVID-19. Amongst these, as per the universal expectations of the WHO EUL/PQ assessment, the Pfizer vaccine and some other candidates have been approved. More studies on vaccine aspirant efficacy and safety results,<sup>157</sup> including on the Moderna and AstraZeneca vaccine, therein have been widely reported, and AstraZeneca have published their results in well reputed journals. Thus we are eagerly expecting more potential COVID-19 vaccine candidates will be offered to governing authorities for approval in the coming years. Gratifyingly, the growth of many efficient COVID-19 vaccine aspirants under clinical trials is fascinating. When the vaccine candidates are proven to be benign and efficient, they need to be acknowledged by the governing authorities, produced to the necessary standard, and distributed. WHO is collaborating all over the world to assist with the roles in this process, which includes facilitating reasonable access to safe and effective COVID-19 vaccines for everyone.

## Author contributions

KD and SA equally contributed to the collection of data and developed the entire manuscript. MD corrected the manuscript. SG and ST assisted in drafting.

## Conflicts of interest

We have no conflicts of interest.

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