Vaccine Innovation for Pandemic Preparedness: Patent Landscape, Global Sustainability, and Circular Bioeconomy in Post-COVID-19 era

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Abstract
In this article, we present breakthroughs and challenges in vaccine development for COVID-19 pandemic, discussing issues related to pandemic preparedness and their implications for circular bioeconomy and sustainability. Notwithstanding the unprecedented accelerated speed of COVID-19 vaccine development, just 9 months after the emergence of the pandemic in Wuhan, China, benefiting from previous developments in SARS and MERS vaccines, significant gaps persist in global vaccine preparedness. These gaps include issues related to immunity and protection, particularly to the limited vaccine protection against recent emergence of concerning new viral variants in the UK, South Africa, and Brazil and the consequent need for vaccine redesign. We examine these gaps and discuss the main issues that could impact on global vaccine availability in the current pandemic scenario: (1) breakthroughs and constraints in development and production of leading global COVID-19 vaccines; (2) innovation and technological development advances and gaps, providing information on global patent assignees for COVID-19, SARS, and MERS vaccine patents; (3) local capacity for development and production of COVID-19, SARS, and MERS vaccines in three emerging agro-based countries (India, Brazil, and South Africa); and (4) future scenarios, examining how these issues and vaccines redesign for new SARS-CoV-2 variants could impact on global access to vaccines and implications for circular bioeconomy and sustainability in the post-COVID era.

Keywords Vaccine innovation · COVID-19 · SARS · MERS · Pandemic preparedness · Circular economy

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Introduction

Vaccines, due to their relatively low costs when compared with pharmaceutical drugs, high cost-effectiveness, and huge impact on economy and society, have emerged in the COVID-19 pandemic as a key bioeconomic product and a crucial component of circular economy, in the direction of a more sustainable and equal world. Vaccines are widely recognized as the most successful and low-cost public health interventions, saving millions of lives, and considered one of the greatest medical achievements of the contemporary world. Significant global scientific and technological progress has been made in the last two decades in this area, with extraordinary breakthroughs in human and veterinary vaccines. These advances, resulting from new paradigms in Vaccinology 4.0, such as genomic vaccines, mRNA technology, and plant expression of vaccine antigens, are evidenced by the unprecedented speed in vaccine development initiatives and international collaborations during COVID-19 pandemic, with several clinical trials just 9 months since the emergence of the outbreak in Wuhan, China. Notwithstanding this accelerated pace of vaccine development, many gaps and challenges persist in global vaccine development and STI governance for COVID-19, including the need to protect against recent concerning new viral variants in the UK, South Africa, and Brazil. In this article, we examine four major issues that could impact on global vaccine availability and sustainability in the post-COVID-19 scenario: (I) leading SARS-CoV-2 vaccines; (II) patent protection, global patent assignees for vaccines against pandemic diseases (COVID-19) or potentially pandemic diseases (SARS, MERS); (III) patent assignees for SARS, MERS, and COVID-19, clinical trials, local capacity for production, and for development of COVID-19 vaccines patents for COVID-19, SARS, and MERS for three agro-based emerging countries (India, South Africa, and Brazil) are described for the period 2010–2020; and (IV) the role of vaccines in the circular bioeconomy model, as a key global resource-saving strategy, preventing expenditures with COVID-19 hospitalizations and life-saving medical devices and equipments, reducing high-risk biological hospital waste, and contributing to meet sustainability and circular economy goals. Finally, in the “Future Scenarios” section, we provide considerations on post-COVID vaccine scenarios. These future scenarios, supported by vaccine patent information, indicate that vaccines are a crucial component of global pandemic preparedness, and thus an exponential increase in funding vaccine research, development, innovation, and production should be an utmost priority.

Conceptual Framework: Vaccines in Circular Bioeconomy Model

Vaccines have emerged in the Vaccinology 4.0 scenario as the future of medicine, expected to increasingly substitute high-cost and resource-consuming curative interventions and consequently, to drastically reduce the high-risk biological waste from these interventions in hospitals and healthcare units, a major environmental and social concern, recently severely aggravated in COVID-19 pandemic. For these reasons, vaccines are key for pandemic preparedness in the direction of a circular bioeconomy model. The urgent need for a transition in the health sector from a linear bioeconomy model (“take, make, dispose”) to a circular bioeconomy model (“renew, remake, share”) is becoming increasingly evident. Notwithstanding, although this economic paradigm change is clear, with circular economy replacing the linear economy, in the direction of a resource-saving and sustainable world, the huge contribution that vaccines can provide to the circular bioeconomy model has not been discussed in
the so far scarce literature on circular economy and health [1], focused on the potential health impacts of circular economy and not on the extraordinary contributions that innovative health products, such as vaccines or preventive antiviral drugs, could provide in a circular bioeconomy model. Vaccines are definitely a key resource-efficient strategy for circular economy, as indicated in Fig. 1, and have already contributed to a sharp reduction in interventions and related direct and indirect expenditures from COVID-19 hospitalizations [2], minimizing the environmental and social impacts of high-risk biological hospital waste, contributing to global sustainability and emerging as a crucial circular economy component.

Methodology

Patent information for COVID-19 vaccines and vaccines for other potentially pandemic diseases (SARS and MERS) is presented for the period 2010–2020, in a search from Derwent Innovation database and from Espacenet database. Information is also provided for the different phases of vaccine clinical trials, from the registry ClinicalTrials.gov. We also provide information for vaccine patents and clinical trials in three emerging agro-based countries (India, Brazil, and South Africa) and an overview of their relative participation in BRICS countries, in a search from abovementioned databases.

COVID-19 Vaccines: State-of-the Art

Vaccines usually require decades of research and testing before reaching clinical trials, but in 2020, in an unprecedented speed and extraordinary international collaboration to respond to the pandemic, scientists succeeded to develop and produce safe and effective COVID-19 vaccines in just 9 months after the emergence of the pandemic in Wuhan, China. Researchers are currently testing 67 vaccines in clinical trials on humans, and 20 have reached the final

Fig. 1 Vaccines in circular bioeconomy model: post- COVID-19 era. Elaborated by the authors
stages of testing [3, 4]. In Table 1, we present the leading SARS vaccines according to technology, efficacy, phase, and regulatory status.

The innovative COVID-19 vaccines presented in Table 1 are in fact the result of a cumulative process starting two decades ago with novel vaccine platforms and new technologies, such as mRNA, modRNA, engineered viral vectors, and other previous novel vaccine strategies against other coronavirus vaccines such as SARS and MERS. The rationale is that, from a cumulative knowledge perspective, these previous breakthroughs were crucial for the accelerated speed of just 9 months from the emergence of the pandemic in Wuhan, China, to produce safe and effective COVID-19 vaccines. Concerning SARS and MERS vaccines, this cumulative process is explained by the structural similarity among the viral proteins responsible to the viral activity (entry and replication into host cells) [9].

In addition, it should be noted that in spite of these breakthroughs in COVID-19 vaccine development, significant knowledge gaps persist, related to immunity and protection. It is not clear yet the extent of duration of immunity and protective immune responses. There are evidence suggesting that adaptive immune responses elicited by SARS-CoV-2 infection might protect against reinfection [10]. Results from seasonal coronaviruses infection [11] and current experience with SARS-CoV-2 suggest that immunity to natural infection might wane over time (“waning immunity”) and reinfection has been reported [12]. There is the possibility that

| Enterprise                   | Technology                | Efficacy (%) | Phase | Status                                           |
|------------------------------|---------------------------|--------------|-------|-------------------------------------------------|
| Moderna                      | mRNA                      | 94.1         | 3     | Approved in Switzerland Emergence use in US, UK, EU, others |
| Pfizer-BioNTech              | mRNA                      | 95           | 2-3   | Approved in Several Countries Emergence use in US, EU, others |
| Oxford/AstraZeneca           | Viral vector ChAdOx1 (engineered) | 62-90*      | 2-3   | Emergence use in UK, EU, others |
| Sinovac                      | Inactivated               | 50.4**       | 3     | Approved in China Emergency use in Brazil, others |
| Johnson&Johnson              | Ad26                      | 72           | 3     | Emergency use in U.S., E.U., other countries. |
| Sinopharm                    | Inactivated               | 79           | 3     | Approved in China U.A.E., Bahrain Emergency use in Egypt, others |
| Gamaleya                     | rAd26, rAd5               | 92           | 3     | Early use in Russia Emergency use in other countries |
| Bharat Biotech               | Inactivated               | Not available: no publication | 2-3 | Restricted emergency use in India |

*62% at first half dose, 90% at second dose

**Different efficacy results from trials in different countries: Brazil 50.4%, Turkey 91.2%, and Indonesia 65.3%. For Turkey and Indonesia provided interim results from late stage trials. Publications and additional data needed to clarify efficacy rates

(1)As of February 20, 2021

Sources: Baden et al. [5], Polack et al. [6], Logunov et al. [7], WHO [3], Voysey et al. [8]

Elaborated by the authors
additional booster doses might be necessary to extend the duration of protection. It is also unclear whether previously infected persons would benefit from vaccination, although vaccination against SARS-CoV-2 is recommended irrespective of infection status [13].

In the “Leading Global Patent Assignees: Vaccines for COVID-19, SARS, and MERS” section, a search in patent documents is provided, describing these preceding breakthroughs and novel vaccine platforms.

Leading Global Patent Assignees: Vaccines for COVID-19, SARS, and MERS

In this section, we examine, for the period 2010–2020, patent data from leading global patent assignees for vaccines against pandemic (COVID-19) or potentially pandemic (SARS and MERS). To define the universe of research, our first search was directed to identify, from a global perspective, where the vaccines and related technologies for COVID-19, SARS, and MERS disease had been developed1. A total of 1235 individual records were recovered, with the USA and China as top assignees, as indicated in Fig. 2.

The top assignees are shown in Fig. 3. From the graphs’ analysis, in spite of CureVac leading position, no company stood out for numbers of applications.

A further novel finding scenario was observed for the BRICS’ countries (Brazil, Russia Federation, India, China, and South Africa) (Fig. 4). Here the top assignee is China. India and Russia Federation have a small representation in the rank. Brazil and South Africa had not applied any priority document for vaccines for SARS, MERS, or COVID-19 according to the strategy search2(A).

India, South Africa, and Brazil: Patent Assignees for SARS, MERS, and COVID-19

In order to describe the specific role of the three agro-based emerging countries, India, South Africa, and Brazil, in vaccine development scenario, a new strategy search was applied2(B). For all countries, a few vaccine patent documents for COVID-19 were observed (considering both criteria, country’s priority or application of global assignees in the country, which was expected due to the patent secrecy’s period). The results are described below (Table 2).

EPO European Patent Office

(*) 1 document: Canada and Netherlands

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1 The search was done in Derwent Innovation Database (Clarivate Analytics) using terms for Respiratory Syndromes associated to coronavirus (SARS OR “SEVERE ACUTE RESPIRATORY SYNDROME” OR MERS OR “MIDDLE EAST RESPIRATORY SYNDROME” OR COVID* OR “SARS-CoV-2” OR “WU-HAN VIRUS” OR “2019 nCoV” OR “SARS-2”) associated to the word vaccine or vaccination in the search field “Title/abstracts/claims.” The strategy search used the following International Patent Classification (IPC): A61k0039 (IPC group: for medical preparations containing antigens or antibodies) or A61k0039215 (IPC subgroup: vaccine against coronavirus (European Patent Office–EPO).

2 The search was done in Derwent Innovation Database (Clarivate Analytics) using terms for Respiratory Syndromes associated to coronavirus (SARS OR “SEVERE ACUTE RESPIRATORY SYNDROME” OR MERS OR “MIDDLE EAST RESPIRATORY SYNDROME” OR COVID* OR “SARS-CoV-2” OR “WU-HAN VIRUS” OR “2019 nCoV” OR “SARS-2”) associated to the word vaccine or vaccination in the search field “Title/abstracts/claims.” The strategy search used the following International Patent Classification (IPC): A61k0039 (IPC group: for medical preparations containing antigens or antibodies) or A61k0039215 (IPC subgroup: vaccine against coronavirus (European Patent Office–EPO). A. The “priority country/region earliest” are from BRICS (Brazil, Russia, India, China, and South Africa (total: 111 DWPI families); B. The priority countries were IN (India), ZA (South Africa), and BR (Brazil) separately.
India: Vaccine Patent Assignees for SARS, MERS, and COVID-19

The search recovered only 6 Indian patent documents (Indian companies), most of them from 2020. There was not a leading Indian vaccine applicant among them. The Patent Document\(^3\) PN: IN202011016574A (Mangalmay Institute of Engineering and Technology, India) [14] discloses about a novelty: an isolated human coronavirus which causes SARS and comprises a nucleotide sequence/acid molecule which under stringent condition can hybridize. This invention is “related to Computer Engineering and newly isolated human coronavirus” with scientific research and clinical applications including the development of vaccines.

The invention from Sinha Kanishk (PN: IN202031014832A (individual’s application) [15] “Vaccine for Novel Coronavirus” is related to advanced immunotherapy to treat novel coronavirus patients. Its composition comprises “cytokine-expressing, proliferation incompetent, whole novel coronavirus cells, anti-programmed death 1 (PD-1) antibody, and toll like receptor (TLR) agonist.” Another invention related to composition of infectious mRNA, RNA, and RNA of SARS coronavirus, SARS coronavirus replicon particles, recombinant ones, and methods of making and using the compositions. It is related to Computer Engineering and Medical Science. The composition expresses heterologous nucleic acids and can be used as vaccines and/or immunogens (PN: IN202041016724A-individual’s application) [16].

The document PN: IN202011018851A (individual’s application) [17] and PN: IN202011024811A (individual’s application) [18] described a diagnostic kit reagent related to proteins and nucleic acids from COVID-19 and to vaccine formulations which comprises respiratory virus antigens (one or more) and COVID-19 virus antigens (one or more).

\(^3\) In the present research, the term “Patent Number or PN” refers to the Publication Number of the applied patent.
The results highlighted that little was recovered about the COVID-19’s vaccine. A complementary search was done to give clear better results about the subject matter.

How it was expected (due to patent document’s secrecy period), only one document from Indian individual applicants was recovered: PN: IN202021024459A [19], which is about a high immunogenic and stable vaccine with the inactivated virus from partially attenuated and/or normal strain.

Foreign Assignees in India We identified 18 applicants/assignees (companies/institutions/countries) which applied in India (8 from the USA, 8 from Europe, and 2 from Asia). The main applicant was CureVac GMBH from Germany (with 4 documents).

North America The patent document from Novavax Inc. - US (PN: IN201617013371A) [20] discloses about nanotechnology: Immunogenic compositions which comprises nanoparticles containing MERS virus proteins. These particles are in polymer compositions and structures. These compositions can be used as vaccines to induce neutralizing antibodies against the virus. One intriguing finding is that an immunogenic composition for inducing immune response against MERS-CoV in subjects is also cost-effective, with an excellent biological stability (PN: IN201617021769A) (the Trustees of the University of Pennsylvania, USA) [21]. Another promising finding is that the Regeneron Pharmaceuticals, Inc., USA applied (PN: IN201647039008A) [22] an invention which provides monoclonal antibodies to bind MERS-CoV spike protein (Regeneron Pharmaceuticals, Inc., USA). Moreover, a novelty in polymeric nanoparticles was applied by President and Fellows of Harvard College/Massachusetts Institute of Technology/The Brigham and Women’s Hospital, Inc., USA (PN: IN201503426P2) [23] which stimulates mucosal immune response by the administration of an inactivated pathogen (preferably bacteria) or virus like SARS coronavirus or common cold influenza (President and Fellows of Harvard College; Massachusetts Institute of Technology; The Brigham and Women’s Hospital, Inc.). Our results casts a new light on immunogenic compositions: The use of a silicified virus/silicified virus, including SARS, MERS, and

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4 Espacenet and Derwent Innovation Index Databases. Strategy Search: Terms: COVID-19 or COVID or “Coronavirus Disease-19” or “SARS-COV-2” or nCoV or “novel coronavirus” or “SARS2” or “Wuhan coronavirus”) and vaccine, without a specific period. The research was conducted on January 25, 2021.
influenza virus to induce (PN: IN201506559P1) [24] a virus-specific immune response which are useful as vaccines (Portland State University; Providence Health & Services Oregon, USA). Moreover, an applied biotechnology method (PN: IN201510078P1-University of Iowa Research Foundation; The University of Kansas, USA): a chimeric virus was isolated and it comprises “bocavirus capsid protein and recombinant adeno-associated viral (AAV) genome.” Virus gene product proteins from SARS or influenza were claimed [25].

The document recovered PN: IN201305009P4 [26] (Medicago Inc., Canada) was aligned with applied biotechnology: it is a method to introduce the nucleic acid, expressed in the plant. Some virus fragments or virus trimeric surface protein was chosen. Among them, SARS virus,

Table 2 Initial results of applicant documents for the priority countries: India, South Africa, and Brazil according to strategy search

| Countries | Origin: priority country | Number of applicant documents for SARS, MERS, and COVID-19 |
|-----------|--------------------------|----------------------------------------------------------|
|           | Origin: countries with applications in India, South Africa, and Brazil (priority countries) |
| Groups    | North America | Europe | Asia |
|-----------|--------------|--------|------|
| India     | 6            | 8      | 8    | 2    |
| South Africa | 0          | 3 (*)  | 1    | -    |
| Brazil    | 0            | 6      | 4    | 2    |

Elaborated by the authors

Source: Derwent Innovation (Clarivate Analytics)
influenza virus, measles virus, etc. That is a production of a virus like particle (VLP) in a plant to prevent or treat (vaccine) these viral infections. The document from Folia Biotech Inc., Canada (PN: IN201310534P1) [27] is also an example of VLPs in vitro. It comprises “combining recombinant Potexvirus coat protein and single stranded RNA, and treating the VLPs with nuclease to remove any RNA protruding from the particles” to be used in combination with antigens (from virus like SARS) as a vaccine.

**Europe** From Novartis AG from Switzerland (PN: IN201203534P2) [28] was applied an invention of methods to prepare squalene which can be used in a vaccine composition for diseases like SARS coronavirus and others (HIV, varicela, human papillomavirus, etc.).

The document (PN: IN291696B-CureVac AG from Germany) [29] is an immunostimulatory composition that may be used like a vaccine or a pharmaceutical composition from CureVac AG. It can be used to treat SARS, influenza, autoimmune diseases, tumor diseases, and others.

The document (PN: IN201404812P1) [30] (CureVac GMBH from Germany) reinforces the general belief that the use of the nucleic acid to increase the expression of a pathogenic antigen which was encoded. About this invention from CureVac GMBH, the nucleic acid “comprises coding for a histone stem-loop and a poly(A) sequence or a polyadenylation signal” for genetic vaccination for prophylactic treatment or therapeutic use of infectious diseases, like SARS or influenza, for example. A further novel finding is about vaccine formulations. A polyple formulation which can be used to deliver RNA to a target organ/cell after intramuscular administration particularly (PN: IN202047028961A-BioNTech RNA Pharmaceuticals GmbH and TRON-Translationale Onkologie an der Universitätsmedizin der Johannes Gutenberg-Universität Mainz GmbH. Both of them from Germany) is used against the antigen or epitope of infectious diseases like SARS, influenza, etc. [31]. A further novel finding is the new carbohydrate-glycolipid conjugates (PN: IN201408326P1-Max-Planck-Gesellschaft zur Förderung der Wissenschaften e. V. from Germany and Universitätsspital Basel from Switzerland) That is used to prepare vaccines against infectious diseases [32]. The saccharide antigen can be from a viral glycoprotein of SARS, influenza viral, etc. And a RNA vaccine (PN: IN201506225P1-CureVac GMBH, Germany) with an “open reading frame coding for antigen and composition with a programmed death-1 pathway inhibitor” for treatment and prevention of cancer or infectious diseases, like SARS and influenza [33]. An intranasal immunization (compositions and methods) with recombinant Modified Vaccinia virus Ankara was claimed using the nucleic acid which encodes the antigen for the induction of an immune response. The antigens can be a tumor-associated antigen, bacteria, parasite, fungus, or virus (e.g., SARS, influenza, or parainfluenza virus) and flagellin (PN: IN201717009706A-Bavarian Nordic A/S, Denmark) [34]. A RNA composition (PN: IN201817022402A) was applied by CureVac AG (Germany) used as vaccine, gene-therapeutic agent, or immunotherapeutic agent. The selected pathogen can be a virus like SARS [35].

**Asia** The document applied by the University of Tokushima, Japan (PN: IN201207596P1) [36] is a mucosal vaccine which comprises the drug and antigen vehicle, carboxyvinyl polymer, and the antigenic protein for induction of immune response against many virus, including SARS virus, influenza virus, yellow fever virus, etc. Indeed, the method, the mutated influenza virus, attenuated, and the live vaccine from the document PN: IN201817042073A-Emerging Viral Vaccine Limited and Versitech Limited. Both from China) comprise “an insertion of gene encoding for an exogenous antigen” which can be an SARS/MERS virus or influenza virus, for example [37].
South Africa: Vaccine Patent Assignees for SARS, MERS, and COVID-19

According to our strategy search, no document was recovered for South Africa as priority country.

Foreign Assignees in South Africa Our results demonstrated that there are a few number of patent documents (four) from other country. The main applicants were Medicago Inc. (from Canada), Novartis AG (Switzerland), Irdeto BV (Netherlands), Univ Iowa Res Found, and Univ Kansas (both of them from US). Three documents were applied in 2010 and only one in 2013.

North America A “new isolated chimeric virus comprises bocavirus capsid protein and recombinant adeno-associated viral genome” (PN: ZA201507946B-Univ Iowa Res Found; Univ Kansas. Both from US). The gene product includes a viral antigen (from SARS virus protein, e.g.) which is a prophylactic gene product [38].

In addition, production of virus like particle to treat and prevent viral infections, including SARS, influenza, rabies, etc. (PN: ZA201305407B-Medicago Inc., Canada; ZA201400166B (Medicago Inc., Canada; Irdeto BV - Netherlands) [39, 40].

Europe A vaccine comprising squalene as adjuvant (PN: ZA201208381B-Novartis AG, Switzerland) against SARS and other virus like influenza virus, HIV, etc [41].

Brazil: Vaccines Patent Assignees for SARS, MERS, and COVID-19

No Brazilian documents were recovered according to our search strategy.

Foreign Assignees in Brazil Only a few documents have shown the applications in Brazil: 12 documents. None of them was recovered to Brazil as priority country. Most documents were applied in 2013 (four) and the main assignee was CureVac GMBH (Germany).

North America A new live and attenuated coronavirus (SARS, MERS, etc.) which has excellent stability to prevent diseases was claimed by Loyola University Chicago from the USA. The coronavirus (mutation of wild-type) has amino acid changes (PN: BR112019018251A2) [42].

From Portland State University, USA (PN: BR112015017966A2) [43]: An immunogenic composition to induce a virus-specific immune response and to raise the “virus specific cell-mediated immune response” against SARS, MERS, etc.

BR112015021970A2 [44] from President and Fellows of Harvard College, USA, is about a composition which includes an adjuvant loaded polymeric nanoparticles - attached to the inactivated antigen to promote the stimulation of mucosal response against SARS, common cold-influenza, dengue fever, etc. Moreover, a MERS-CoV nanoparticle where it is possible the administration of “the purified high affinity antibody to a human subject” from Novavax Inc., US (PN: BR112016006122A2) [45].

The technology to produce human monoclonal antibodies (recombinant antibodies) from Regeneron Pharmaceuticals, Inc., USA (PN: BR112016025009A2) to prevent MERS infection. Moreover, MERS-CoV spike proteins bind to “isolated recombinant antibody or its antigen-binding fragment” [46].
Production of VLP in a plant was also claimed, by the introduction of the nucleic acid in a plant. The composition can be used to treat/prevent viral infection, including SARS, smallbox (PN: BR112013016185A2–Medicago Inc., Canada) [47].

**Europe** A method to prepare a squalene composition for vaccines preparations (adjuvant) is new. The composition, which comprises parvalbumin and proteins, uses antigens for the development of the vaccine against SARS, Influenza, HIV, etc. (PN: BR112012028711A2–Novartis AG, Switzerland) [48].

A protein/peptide which increases expression to treat infectious diseases from the nucleic acid sequence (pathogenic antigen for SARS, HIV, dengue, etc) from CureVac GMBH, Germany (PN: BR112014016361A2) [49].

From Universitätsspital Basel, Switzerland (PN: BR112014023193A2) [50], an active agent raises the immune response against infectious diseases (e.g., SARS and influenza). Bacterial capsular saccharide antigen or other antigens (e.g., viral glycoproteins are used).

A RNA vaccine which comprises a códon-optimized region and (at least) one open reading frame (ORF) to treat/prevent disease like cancer and infectious ones: SARS, influenza, etc. (PN: BR112015018989A2–CureVac GMBH, Germany) [51].

**Asia** From the University of Tokushima, Japan, a new vaccine of mucous membrane was applied. A mucosal vaccine (antigen and drug vehicle) can be used to induce immune response against antigens as SARS, influenza virus, dengue virus, etc. (PN: BR112012022059A2) [52].

The mutant virus (e.g., SARS and I was applied by Peking University from China can be used as an attenuated vaccine. The mutated virus is prepared by (e.g., “expressing mutated nucleic acid expression vector” (PN: BR112018013168A2) [53].

**India, South Africa, and Brazil: Clinical Trials for COVID-19 Vaccines**

Brazil, India, and South Africa play an important role as sites for collaboration in international clinical trials for COVID-19 vaccines. The significant investment by pharmaceutical companies in clinical trials in these three countries contrasts with limited investments in technological transfer agreements and strengthening local capacity for technological development in these countries.

Clinical Trials for COVID-19 vaccines are illustrated in Tables 3 (India), 4 (South Africa), 5 (Brazil), and 6 (multi-country). These tables present the clinical trial phase, status, and sponsor’s descriptions. The search was conducted from the base clinicaltrials.gov and a strategy search. We also searched the COVID-19, living NMA initiative provided by WHO that collected 164 RCTs and 48 non-randomized studies of vaccines for COVID-19 from the ICTRP. Our results cast a new light on the participation of the referred countries in vaccine’s development (Tables 4, 5, and 6).

**Development and Production of COVID-19 Vaccines: India, South Africa, and Brazil**

Concerning the development and production of COVID-19 vaccines, Table 7 presents the partnerships with leading enterprises made by institutions in India, South Africa, and Brazil, for

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5 Database: Clinical Trials. The search used the terms “COVID-19, COVID, SARS-CoV-2, Severe Acute Respiratory Syndrome Coronavirus 2” (January 22, 2021)
vaccine development based on data recovered from the Chemical Business NewsBase (CBNB). Table 8 presents local capacity for production in these countries. These Tables evidence the leading role of India in technological capacity for development and production of COVID-19 vaccines.

### Table 3: India clinical trials for COVID-19 vaccines—country-based trials as of February 18, 2021

| Title                                                                 | Phases | Status | Sponsor                                                                 |
|----------------------------------------------------------------------|--------|--------|-------------------------------------------------------------------------|
| An Efficacy and Safety Clinical Trial of an Investigational COVID-19 Vaccine (BBV152) in Adult Volunteers | 3      | R      | Bharat Biotech International Limited|Indian Council of Medical Research|iqvia Pty Ltd |
| BCG Vaccine in Reducing Morbidity and Mortality in Elderly Individuals in COVID-19 Hotspots | 3      | R      | Tuberculosis Research Centre, India|ICMR-National Institute for Research in Tuberculosis, Chennai, Tamil Nadu|All India Institute of Medical Sciences, New Delhi|National Institute for Research in Environmental Health, Bhopal, Madhya Pradesh|National Institute of Occupational Health, Ahmedabad, Gujarat|King Edward Memorial Hospital|National Institute for Implementation Research on Non-Communicable Disease |
| Clinical Trial to Assess Safety and Immunogenicity of Gam-COVID-Vac Combined Vector Vaccine for Severe Acute Respiratory Syndrome Coronavirus 2 (SARS-D(OV-2) Infection – Gamaleya-Sputnik Vaccine | 2 / 3  | R      | Dr. Reddy's Laboratories Limited|Gamaleya Research Institute of Epidemiology and Microbiology, Health Ministry of the Russian Federation|RDIF (Russian Direct Investment Fund)|CRO: JSS Medical Research India Pvt. Ltd. |
| Whole-Virion Inactivated SARS-CoV-2 Vaccine (BBV152) for COVID-19 in Healthy Volunteers | 1 / 2  | A      | Bharat Biotech International Limited|Indian Council of Medical Research |
| Safety and Immunogenicity of an Intranasal SARS-CoV-2 Vaccine (BBV154) for COVID-19 | 1      | N      | Bharat Biotech International Limited |
| Novel Corona Virus-2019-nCov vaccine by intradermal route in healthy subjects. | 1 / 2  | A      | Cadila Healthcare Ltd |
| A Phase 2/3, Observer-Blind, Randomized, Controlled Study to Determine the Safety and Immunogenicity of Covishield (COVID-19 Vaccine) in Healthy Indian Adults | 2 / 3  | A      | Serum Institute of India Private Limited|Indian Council of Medical Research ICMR |
| Whole-Virion Inactivated SARS-CoV-2 Vaccine (BBV152) in Healthy Volunteers | 1 / 2  | A      | Bharat Biotech International Limited |
| A Phase 1, Followed by a Phase 2, vaccine given Randomly in subjects in different sites to Evaluate the Safety, side effects, and resistance of the Virus Vaccine, BBV152D Administered between the layers of the skin (intradermal) in Healthy Volunteers | 1 / 2  | A      | Bharat Biotech International limited |
| Biological E’s novel Covid-19 vaccine of SARS-CoV-2 for protection against Covid-19 disease. | 1 / 2  | A      | Biological ELimited |
| Novel Corona Virus-2019-nCov vaccine by intradermal route in healthy subjects. | 3      | R      | Cadila Healthcare Ltd |

*R* recruiting, *N* not yet recruiting, *A* active, not recruiting

Source: clinicaltrials.gov Database and COVID-19-living NMA initiative

Elaborated by the authors
vaccines, led by the Serum Institute of India. In second position, Brazil with two technological transfer agreements from Oxford/AstraZeneca with Oswaldo Cruz Foundation/Bio-Manguinhos and from SINOVAC with Butantan Institute and local capacity for production in both institutions.

### Table 4 South Africa clinical trials for COVID-19 vaccines—country-based trials as of February 18, 2021

| Title                                                                 | Phases | Status | Sponsor                                                                 |
|----------------------------------------------------------------------|--------|--------|-------------------------------------------------------------------------|
| A Study Looking at the Effectiveness and Safety of a COVID-19 Vaccine in South African Adults | 2      | R      | Novavax|Bill and Melinda Gates Foundation                                         |
| COVID-19 Vaccine (ChAdOx1 nCoV-19) Trial in South African Adults With and Without HIV-infection | 1 / 2   | A      | University of Oxford|Medical Research Council, South Africa|Bill and Melinda Gates Foundation|Wits Health Consortium (Pty) Ltd|University of Witwatersrand, South Africa |

R recruiting, N not yet recruiting, A active, not recruiting
Source: [clinicaltrials.gov](https://clinicaltrials.gov) Database and COVID-19-living NMA initiative
Elaborated by the authors

### Table 5 Brazil clinical trials for COVID-19 vaccines—country-based trials as of February 18, 2021

| Title                                                                 | Phases | Status | Sponsor                                                                 |
|----------------------------------------------------------------------|--------|--------|-------------------------------------------------------------------------|
| A Study of a Candidate COVID-19 Vaccine (COV003)                      | 3      | R      | University of Oxford                                                   |
| Clinical Trial of Efficacy and Safety of Sinovac's Adsorbed COVID-19 (Inactivated) Vaccine in Healthcare Professionals | 3      | R      | Butantan Institute|Sinovac Life Sciences Co., Ltd.                                      |
| COVID-19: BCG As Therapeutic Vaccine, Transmission Limitation, and Immunoglobulin Enhancement | 4      | R      | University of Campinas, Brazil|Conselho Nacional de Desenvolvimento Científico e Tecnológico|Instituto de Infectologia Emilio Ribas|Pontificia Universidade Católica de Campinas, PUC-Campinas|Faculty of Medicine of Ribeirão Preto (FMRP-USP)|Faculdade de Medicina de Botucatu, UNESP, Botucatu, Brasil|Federal University of São Paulo|State Hospital Dr. Leandro Franceschini, Sumaré, Unicamp|Paulinia Municipal Hospital Universidade Federal do Rio de Janeiro|Ministry of Science and Technology, Brazil |
| Use of BCG Vaccine as a Preventive Measure for COVID-19 in Health Care Workers | 2      | R      | University of Oxford                                                   |
| A phase III study to investigate a vaccine against COVID-19           | 3      | A      | University of Oxford                                                   |
| Clinical Trial of Efficacy and Safety of Sinovac's Adsorbed COVID-19 (Inactivated) Vaccine in Healthcare Professionals (PROFISCOV) | 3      | A      | Butantan Institute; Sinovac Life Sciences Co., Ltd.                     |

R recruiting, N not yet recruiting, A active, not recruiting
Source: [clinicaltrials.gov](https://clinicaltrials.gov) Database and COVID-19-living NMA initiative
Elaborated by the authors
Our results indicate that a concerning geopolitical scenario is emerging, with developed countries and three BRICS’ countries (China, India, and Russia) dominating the development and production of patented vaccines, active principles (APIs) and adjuvants, leaving developing countries, in greatest need, with very limited vaccine stocks.

Table 6  Multi-country clinical trials for COVID-19 vaccines–country-based trials as of January 22, 2021

| Title                                                                 | Phases | Status | Sponsor                                                                                       | Countries                                      |
|----------------------------------------------------------------------|--------|--------|------------------------------------------------------------------------------------------------|------------------------------------------------|
| The Efficacy, Safety and Immunogenicity Study of Inactivated SARS-CoV-2 Vaccine for Preventing Against COVID-19 | 3      | N      | Chinese Academy of Medical Sciences                                                             | Brazil and Malaysia                            |
| A Controlled Phase 2/3 Study of Adjuvanted Recombinant SARS-CoV-2 Trimeric S-protein Vaccine (SCB-2019) for the Prevention of COVID-19 | 2 / 3  | N      | Clover Biopharmaceuticals AUS Pty Ltd/The Coalition for Epidemic Preparedness Innovations/International Vaccine Institute | Belgium, Brazil, Colombia, Dominican Republic, Germany, Nepal, Panama, Philippines, Poland, South Africa |
| Study to Describe the Safety, Tolerability, Immunogenicity, and Efficacy of RNA Vaccine Candidates Against COVID-19 in Healthy Individuals | 2 / 3  | A      | BioNTech SE/Pfizer                                                                             | USA, Argentina, Brazil, Germany, South Africa, Turkey |
| Phase III Double-blind, Placebo-controlled Study of AZD1222 for the Prevention of COVID-19 in Adults | 3      | A      | AstraZeneca | Iqvia Pty Ltd                                                                 | Argentina; Chile; Colombia; Czechia; France; Germany; India; Italy; Netherlands; Peru; Spain; USA |
| A Study of Ad26.COV2.S for the Prevention of SARS-CoV-2-Mediated COVID-19 in Adult Participants (ENSEMBLE) | 3      | A      | Janssen Vaccines & Prevention B.V.                                                             | USA; Chile; Colombia; Mexico; Peru; Brazil; South Africa |
| A Study of Ad26.COV2.S for the Prevention of SARS-CoV-2-mediated COVID-19 in Adults (ENSEMBLE 2) | 3      | R      | Janssen Vaccines & Prevention B.V.                                                             | Belgium, Brazil, Colombia, France, Germany, Philippines, South Africa, Spain, UK, USA |
| COVID-19: a Phase 2a, partially observer-blind, multicenter, controlled, dose-confirmation clinical trial to evaluate the safety, reactogenicity and immunogenicity of the investigational SARS-CoV-2 MRNA vaccine cvncov in adults ≥60 years of age and 18 to 60 years of age | 2      | A      | CureVac Ag                                                                                     | Brazil; Panama; Peru                            |

R recruiting, N not yet recruiting, A active, not recruiting

Source: clinicaltrials.gov Database and COVID-19-living NMA initiative
Elaborated by the authors

Discussion

Our results indicate that a concerning geopolitical scenario is emerging, with developed countries and three BRICS’ countries (China, India, and Russia) dominating the development and production of patented vaccines, active principles (APIs) and adjuvants, leaving developing countries, in greatest need, with very limited vaccine stocks.
| Institution (country)/vaccine or vaccine candidate | COVID-19 vaccine: development partnerships | Note: vaccine/vaccine candidate |
|--------------------------------------------------|---------------------------------------------|---------------------------------|
| Fiocruz/Instituto De Tecnologia De Imunobiológicos (Brazil) | - | - Technology Transfer from AstraZeneca |
| Akers Biosciences Inc. (USA) and Premas biotech (India) | Development Partnership: vaccine prototype completed | - |
| Beximco Pharmaceuticals Ltd. (Bangladesh) | Investment in Serum Institute of India–development of COVID-19 vaccine | - |
| Bharat Biotech (India)/University Of Wisconsin-Madison and FluGen (USA) | Development and testing of a intranasal vaccine | - |
| Biological E Limited (India) and Ohio State Innovation Foundation (USA) | License to the “novel live attenuated measles virus vectored vaccine candidates” | - |
| Instituto Butantan (Brazil) | - | - Technology Transfer from Sinovac Life Sciences CoronaVac |
| Codagenix Inc. (US)/CDX 005 | Collaboration for vaccine development with Serum Institute of India | - |
| Etna Biotech (the European research arm of Zydus Cadila (India) | Working in measles vaccines for COVID-19 | - |
| Gennova Biopharmaceuticals (India)/HGCO19 | “First vaccine based on messenger ribonucleic acid (mRNA) platform” | - |
| Indian Institute of Technology and Hester Biosciences (India) | Development of a vaccine versus COVID-19 | - |
| Mynvax, Bharat Biotech, Serum Institute of India, and Cadila Healthcare (Zydus Cadila) (India) | Development of a vaccine | - |
| Novavax (USA)/Covovax | Collaboration for the development: Serum Institute of India; Indian Council of Medical Research (ICMR) | - |
| PFIZER (USA) | India as arsenal to the vaccine development (announced by the company) | - |

Table 7 COVID-19 vaccines: development partnerships in India, Brazil, and South Africa
Consequently, as recently confirmed by the WHO Director-General in the Parliamentary Assembly of the Council of Europe (PACE), 75% of global vaccine doses are concentrated in just 10 countries [4]. Our results also indicate the dynamism and proactivity of India in the production and clinical trials of different vaccines (Tables 7 and 8). Among the three agro-based countries in our study (India, South Africa, and Brazil), India has by far a leading protagonist role in the vaccine preparedness. Our study provides evidence that this post-COVID scenario tends to be dramatic for developing countries, particularly in other two BRICS’s and agro-dependent countries examined here (South Africa and Brazil). It will be necessary otherwise to conceive novel governance strategies to strengthen local manufacturing capacity in these countries, supporting vaccine RD&I institutes and manufacturers to incorporate new technologies for production of innovative products. Multinational and startup companies have the intellectual property of new vaccine technologies, adjuvants, and novel vaccine platforms, but often many of these companies do not have sufficient production capacity to meet the global demand for these products and rely on technological transfer to public and private manufacturers both in developed and developing countries to expand their production. Other important issues are the increasingly strict international vaccine regulatory requirements, which are pressuring developing countries’ manufacturers for exponential investments in compliance, modernization, and

### Table 7 (continued)

| Institution (country)/vaccine or vaccine candidate | COVID-19 vaccine: development partnerships | Note: vaccine/vaccine candidate |
|---------------------------------------------------|------------------------------------------|---------------------------------|
| **India** | | |
| Pune-Based Serum Institute of India (India); Max Planck Institute for Infection Biology and Vaccine Projekt Management (Germany) | Evaluation of tuberculosis vaccine for COVID-19 vaccine | - - - |
| União Química (Brazil) | - | Pilot Scale |
| Zydus Cadila (India) | Development of vaccine (*) : two methods | - - |
| Zydus Cadila (India) | Development: Two candidate vaccines | - - |

(*): Including this country

Source: Chemical Business New Base (CBNB) in STN International (January, 2021), INPI.gov.br; Butantan.gov.br; UniãoQuímica.com.br

Elaborated by the authors
capacity-building to meet these requirements, although resources in these countries are often insufficient to comply to them [54].

**Future Scenarios**

Our study evidences that a breakthrough scenario has emerged, in a new era of vaccine development, the Vaccinology 4.0 revolution, with accelerated development of disruptive technologies contributing to a new paradigm in vaccine development, such as for instance, the new mRNA, engineered viral vector and peptide-based vaccine platforms. These novel vaccines platforms radically differ from the previous traditional ones, allowing rapid and readily scalable production, with high productivity and flexibility, in which the same process can be used to produce vaccines against different indications. We have now several COVID-19 vaccines authorized in different countries and policy makers often question if it would possible to predict which vaccines will perform the best in real-world against the recent concerning variants (United Kingdom B.1.1.7; South-Africa B.1.351; Brazil P1). We consider that it is not possible to compare these vaccines and predict their performance, at this point it

| Institution (country) | COVID-19 vaccine: production | Note: vaccine/vaccine candidate |
|-----------------------|-----------------------------|---------------------------------|
| Aurobindo Pharma (India) | Vaccine production plant | The company builds a production facility for viral vaccines |
| Fiocruz (Brazil) | - | Vaccine’s production AstraZeneca’s vaccine |
| Hetero Biopharma (India) | Production of Sputnik (COVID-19 vaccine) | - |
| Instituto Butantan | - | Vaccine’s production Sinovac Biotech (CoronaVac Vaccine) |
| Panacea Biotec (India) and Refana Inc. (USA) | Development, production, and commercialization | - |
| Serum Institute of India (India) | To export vaccines and collaboration with GAVI (The Vaccine Alliance) | Production of 2 billions doses (approximately) around June 2021. The company teamed up with many countries |
| Serum Institute of India and Indian Council of Medical Research (India)/Novavax (USA) | Manufacturing capacity expanded | Based on a Matrix-M adjuvant and protein |

(*) Including this country

Source: Chemical Business New Base (CBNB) in STN International (January, 2021), Butantan.gov.br, Portal. Fiocruz.br
Elaborated by the authors
would certainly be premature. Although there a few preliminary results and pre-print publications indicating that some vaccines might be effective against these three more worrying COVID-19 variants, many unknown issues related to nature and duration of immunity and protection remain in the scientific agenda to be clarified. Current SARS-CoV-2 vaccines were designed for earlier coronavirus versions. Some scientists, based on these preliminary results and pre-print publications, believe they might also protect against these new variants, but the extent of this protection is not clear yet. We list below the vaccines that might offer some protection against these COVID-19 variants:

1. Pfizer/BioNTech—preliminary results suggest this vaccine protects against these new variants, but there are some indications that the vaccine could be slightly less effective for these variants.
2. Moderna—preliminary results indicate that this vaccine is effective against the South Africa variant, but it is not clear how strong and long-lasting could be the immune response.
3. Oxford/AstraZeneca—preliminary results suggest it protects against the new UK variant. It offers less protection against the South Africa variant, although there are indications it could protect against severe illness.
4. Novavax and Janssen vaccines—there are preliminary results indicating that these two vaccines might offer some protection against these variants.

The emergence of more severe and lethal variants of COVID-19, exponentially aggravating the current pandemic scenario is a real possibility with potentially huge global economic impact and should be a topic of major global concern, requiring urgent redesign of vaccines. The development of an universal SARS-CoV-2 that would protect against all variants ("pan-CoV vaccine"), as supported now by NIAID/NIH, should be a global priority, receiving urgently massive funding and incentives, in an unprecedented effort to intensify international and national collaborations on innovative universal vaccine development, led by WHO, COVAX-Facility, GAVI, CEPI and financially supported by G-7, G-20, The World Economic Forum, the World Bank, the World Trade Organization (WTO), Gates Foundation and other organizations. Developing countries’ scientists and manufacturers should be active players in these global initiatives supporting technological transfer agreements for local production, to accelerate vaccine development against these variants.

Finally, for future scenarios, the issue of access should also be considered. It should be recognized that the novel mRNA vaccines (Pfizer/BioNTech and Moderna) are extraordinary breakthroughs, a revolution in vaccine development and a new era in Vaccinology. Their efficacy is greater, with few adverse events, as stated by CDC in a recent document. Nevertheless, there are some constraints, particularly in developing countries, for their immediate incorporation into National Immunizations Programs-NIPs. Besides the very high prices of these multipatented vaccines [55] there are other infrastructure and logistic constraints, particularly with the Pfizer/BioNTech vaccine, such as the requirement of refrigeration under minus 75 C and once taken out kept refrigerated and used within five days. These constraints are particularly severe in countries of continental dimensions such as Brazil and India, with limited vaccine purchasing capacity and the need to transport these vaccines by road or by river to indigenous and rural communities in distant locations. For the above mentioned
reasons, the development of universal and more accessible vaccines against all variants (“pan-CoV vaccine”), which is also an issue for future Influenza vaccines and other vaccines for potentially pandemic diseases, should be of utmost priority. If approved, these universal vaccines could completely change the future scenario for pandemics.

Conclusion

The COVID-19 pandemic came at a time that the global community was preparing for a circular economy transition by 2050, as illustrated by the recent European Union Circular Economy Action Plan [56]. Our results indicate that an extensive range of breakthrough SARS-CoV-2 vaccines (mRNA, engineered viral vector and peptide-based) emerged in this scenario just nine months after the onset of the pandemic, inaugurating a new era in vaccine development, in the context of Vaccinology 4.0. These novel vaccines are, for their potentially huge economic impact, a crucial circular bioeconomy component and a key strategy for the transition towards a prevention-oriented, resource-saving and sustainable world. Our results also indicate a great concentration of multipatented vaccines in developed countries, with two BRICS’ countries leading the development and production of patented active pharmaceutical ingredients-API and adjuvants (India and China) and a pro-active dynamism of India in vaccine production and clinical trials. This concentration, aggravated by a complex global geopolitical scenario in which just 10 countries account for 75% of global COVID-19 vaccine doses, is dramatically restricting access to vaccines by populations in the poorest countries. Without universal access to SARS-CoV-2 vaccines and other vaccines for potentially pandemic diseases, the world will continue to submerge in a long-standing pandemic scenario, permeated by economic depression, social exclusion and extreme poverty, compromising equity, human rights and environmental sustainability. Although it should be recognized that the poorest countries, such as Brazil and South Africa, are now gradually incorporating the new SARS-CoV-2 vaccines into their National Immunization Programs (NIP), with financial support from the COVAX-Facility, GAVI Alliance, WHO, Gates Foundation and other international organizations, this inclusive donor strategy, in spite of its highly positive results, has limitations in supporting local manufacturing and innovation capacity, constraining local vaccine self-sufficiency and achievement of Sustainable Development Goals (SDG). Finally, as indicated in our future scenarios, the emergence of concerning new SARS-CoV-2 variants, such as the ones in UK, Brazil and South Africa, which are now exponentially aggravating the pandemic, will require an urgent redesign of COVID-19 vaccines, with new funding mechanisms in support to an effective global pandemic preparedness system. In this new system, an innovative global vaccine preparedness strategy is urgently needed, in the direction of a sustainable and resource-saving circular bioeconomy. Finally, our results also indicate, that in order to support this new global vaccine preparedness strategy, developing countries should be active players not only in clinical trials of multinational companies but in sustainable local vaccine RD&I and production.
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Declarations

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