Advances in Vaccine Development: Novel Platforms, Efficacy, and Global Distribution Challenges

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ABSTRACT

Recent advances in vaccine development have revolutionized the prevention and control of infectious diseases, leveraging novel platforms such as mRNA, viral vectors, and nanoparticle-based vaccines. This paper reviews current innovations in vaccine technologies, assessing their mechanisms, immunogenicity, and clinical efficacy against emerging and re-emerging pathogens. Additionally, it explores challenges related to large-scale manufacturing, equitable global distribution, cold-chain logistics, and vaccine hesitancy, which impact the effectiveness of vaccination programs worldwide. Comparative analyses of different vaccine platforms highlight their respective advantages, limitations, and suitability for rapid deployment in response to pandemics. The study emphasizes the need for integrated strategies combining scientific innovation, policy frameworks, and international collaboration to enhance vaccine accessibility and uptake. By synthesizing evidence on novel vaccine technologies and global distribution challenges, this paper provides critical insights for researchers, policymakers, and public health authorities seeking to optimize immunization strategies and strengthen global health resilience.

Keywords: Vaccine Development, mRNA Vaccines, Viral Vector Vaccines, Global Distribution, Immunogenicity

INTRODUCTION

Vaccination remains one of the most effective public health interventions for preventing infectious diseases and reducing global morbidity and mortality. Over the past decade, advances in biotechnology have revolutionized vaccine development, enabling the rapid creation of novel platforms such as mRNA vaccines, viral vectors, protein subunits, and nanoparticle-based vaccines. These innovations have proven particularly crucial in responding to emerging infectious diseases, including COVID-19, Ebola, and influenza variants, demonstrating unprecedented speed in design, production, and deployment. Despite significant scientific progress, several challenges continue to limit the global impact of vaccines. Issues such as cold-chain logistics, manufacturing scalability, regulatory approval processes, and equitable access remain critical barriers, particularly in low- and middle-income countries. Additionally, vaccine hesitancy, misinformation, and sociocultural factors can reduce immunization coverage, undermining herd immunity and public health outcomes. This paper aims to explore the latest advancements in vaccine technologies, evaluate their efficacy and safety profiles, and examine the challenges associated with global distribution and uptake. By synthesizing evidence from clinical trials, technological innovations, and public health strategies, the study provides insights into optimizing vaccine development and ensuring equitable access, ultimately strengthening global health resilience.

THEORETICAL FRAMEWORK

The theoretical framework for this study integrates **immunological principles**, **vaccine platform technologies**, **and global public health policy frameworks** to analyze the development, efficacy, and distribution of vaccines. It provides a structured lens to evaluate both scientific innovation and implementation strategies.

1. Immunological Principles

• Adaptive Immune Response: Effective vaccines elicit a strong and durable adaptive immune response, generating both humoral (antibody-mediated) and cellular immunity.

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- Memory Formation: Vaccines aim to establish immunological memory, ensuring rapid and robust responses upon subsequent exposure to pathogens.
- **Herd Immunity:** Widespread vaccination reduces pathogen transmission, protecting populations and vulnerable groups.

2. Vaccine Platform Frameworks

- **mRNA Vaccines:** Utilize messenger RNA to encode antigens, prompting in vivo protein expression and immune activation. Benefits include rapid development, high efficacy, and adaptability to emerging variants.
- **Viral Vector Vaccines:** Employ engineered viruses to deliver genetic material encoding target antigens. They are effective for inducing strong cellular immunity but may face pre-existing vector immunity challenges.
- **Protein Subunit and Nanoparticle Vaccines:** Deliver purified antigens or antigen-presenting nanoparticles to stimulate immune responses while minimizing safety concerns.
- Comparative Platform Analysis: Each platform differs in terms of scalability, stability, immunogenicity, and manufacturing complexity, which influences global deployment strategies.

3. Public Health and Policy Frameworks

- Global Vaccine Access Models: Policies such as COVAX aim to ensure equitable distribution of vaccines, especially in low-resource settings.
- **Regulatory Oversight:** National and international regulatory bodies (e.g., FDA, EMA, WHO) establish standards for vaccine safety, efficacy, and emergency use authorization.
- **Distribution and Cold-Chain Logistics:** Effective vaccine deployment relies on robust supply chains, storage infrastructure, and monitoring systems to maintain efficacy.
- Vaccine Hesitancy Frameworks: Sociobehavioral theories help understand public acceptance, guiding strategies to increase uptake and trust.

Synthesis

This framework combines **scientific, technological, and policy perspectives** to analyze vaccine development and distribution comprehensively. By integrating immunological mechanisms, platform-specific characteristics, and global public health strategies, the framework provides a basis for evaluating vaccine efficacy, scalability, and accessibility challenges.

PROPOSED MODELS AND METHODOLOGIES

This study employs a combination of analytical models, comparative frameworks, and empirical methodologies to evaluate novel vaccine platforms, clinical efficacy, and global distribution challenges. The approach integrates scientific, public health, and policy perspectives.

1. Proposed Models

a. Vaccine Platform Comparative Model

- Compares mRNA, viral vector, protein subunit, and nanoparticle vaccines across key parameters: immunogenicity, safety, scalability, cost, and adaptability to emerging pathogens.
- Evaluates advantages and limitations of each platform for rapid pandemic response and long-term vaccination programs.

b. Global Distribution and Accessibility Model

• Examines factors affecting equitable vaccine distribution, including manufacturing capacity, supply chain logistics, cold-chain requirements, and regulatory policies.

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Uses a multi-layered framework combining national strategies, international initiatives (e.g., COVAX), and socio-economic determinants.

c. Vaccine Uptake and Hesitancy Framework

- Integrates behavioral, social, and cultural variables influencing public acceptance.
- Analyzes strategies to enhance vaccine coverage, combat misinformation, and foster trust in immunization programs.

2. Research Methodologies

a. Literature Review and Systematic Analysis

- Comprehensive review of peer-reviewed studies, clinical trial data, and policy reports related to vaccine development, efficacy, and distribution.
- Thematic analysis identifies trends, innovations, and persistent challenges across vaccine platforms.

b. Comparative Platform Evaluation

- Analysis of clinical trial outcomes to assess vaccine efficacy, immunogenicity, adverse events, and durability of protection.
- Cross-platform comparison highlights technological advantages, limitations, and suitability for rapid deployment.

c. Case Study and Policy Analysis

- Evaluation of global vaccination campaigns, distribution strategies, and regulatory frameworks during major outbreaks (e.g., COVID-19, Ebola).
- Focus on lessons learned for optimizing supply chains, equitable access, and vaccination coverage.

d. Stakeholder Analysis

- Surveys and reviews of perspectives from public health authorities, vaccine manufacturers, policymakers, and community groups.
- Identifies barriers to adoption, logistical challenges, and public perceptions affecting vaccine uptake.

3. Evaluation Metrics

- Clinical Efficacy: Vaccine effectiveness in preventing infection, severe disease, and mortality.
- Safety Profile: Incidence of adverse events and long-term effects.
- Scalability and Logistics: Manufacturing capacity, cold-chain requirements, and distribution efficiency.
- Equitable Access: Coverage rates across high-, middle-, and low-income regions.
- Public Acceptance: Vaccine hesitancy levels and uptake rates influenced by socio-cultural factors.

Synthesis

By integrating comparative platform analysis, distribution frameworks, and stakeholder perspectives, this methodology allows a **comprehensive assessment of vaccine innovations, efficacy, and global deployment challenges**. It provides actionable insights for researchers, public health authorities, and policymakers to optimize vaccination strategies and improve global health outcomes.

EXPERIMENTAL STUDY

This study evaluates vaccine development and deployment through the analysis of clinical trial outcomes, distribution strategies, and real-world implementation data. It focuses on assessing efficacy, safety, and the challenges associated with global accessibility of novel vaccine platforms.

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1. Study Design

- Type: Multi-method evaluation combining clinical trial data analysis, case studies, and policy assessment.
- **Scope:** Includes mRNA vaccines (e.g., Pfizer-BioNTech, Moderna), viral vector vaccines (e.g., AstraZeneca, Johnson & Johnson), and protein subunit/nanoparticle vaccines (e.g., Novavax).
- **Data Sources:** Peer-reviewed literature, regulatory filings, vaccine surveillance reports, WHO and CDC datasets, and national immunization program records.

2. Intervention / Context

- Clinical Trials: Analysis of Phase II and III trial results for immunogenicity, efficacy, and safety.
- **Distribution Strategies:** Case studies of global vaccination campaigns, focusing on supply chain logistics, cold-chain management, and equity in access.
- **Public Health Programs:** Assessment of vaccination uptake, hesitancy, and public acceptance across different populations.

3. Data Collection and Metrics

- Efficacy Metrics: Reduction in infection rates, hospitalization, and mortality.
- Safety Metrics: Adverse event reporting, severity grading, and long-term monitoring.
- **Distribution Metrics:** Coverage rates, logistic efficiency, cold-chain compliance, and timeliness of delivery.
- Equity Metrics: Vaccine availability in high-, middle-, and low-income regions.
- Public Perception Metrics: Vaccine hesitancy levels, acceptance rates, and engagement with public health campaigns.

4. Key Findings

- **Efficacy:** mRNA vaccines showed >90% efficacy in preventing symptomatic COVID-19 in clinical trials, while viral vector and protein-based vaccines ranged between 60–85%, depending on population and variant exposure.
- **Safety:** All platforms demonstrated acceptable safety profiles; most adverse events were mild or moderate. Rare severe events were identified and monitored post-deployment.
- **Global Distribution:** High-income countries achieved rapid vaccination coverage, whereas low-income regions faced supply, cold-chain, and regulatory challenges.
- Vaccine Uptake: Public hesitancy, misinformation, and logistical barriers influenced overall vaccination rates, highlighting the importance of education and outreach.

5. Limitations of the Experimental Study

- Reliance on reported trial data and government statistics may omit underreported adverse events or informal distribution efforts.
- Regional variability in data quality may limit generalizability of findings.
- Rapidly evolving viral variants and emerging vaccine platforms may require continuous updating of the analysis.

Synthesis

The experimental study demonstrates that **novel vaccine platforms are effective and generally safe**, but challenges in global distribution, equity, and public acceptance remain significant barriers. Effective vaccination programs require integration of scientific innovation, logistical planning, and policy measures to ensure broad, equitable immunization coverage.

RESULTS & ANALYSIS

The analysis of clinical trial outcomes, distribution strategies, and public health data provides insights into the efficacy, safety, and accessibility of novel vaccine platforms. Key findings from mRNA, viral vector, and protein-based vaccines are summarized below.

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1. Vaccine Efficacy

- mRNA Vaccines: Demonstrated the highest efficacy (>90%) against symptomatic infection and severe disease in clinical trials. Efficacy remained high across various demographic groups and with emerging viral variants, though booster doses improved protection against new strains.
- **Viral Vector Vaccines:** Moderate efficacy (60–80%) observed, with stronger T-cell responses noted. Some efficacy variability occurred due to pre-existing immunity to the viral vector and regional variant prevalence.
- **Protein Subunit/Nanoparticle Vaccines:** Efficacy ranged from 70–85% with favorable safety profiles. They offer advantages in storage stability and are suitable for low-resource settings.

2. Safety and Adverse Events

- Common Reactions: Mild-to-moderate local (pain, redness) and systemic (fever, fatigue) reactions were observed across all platforms.
- **Severe Events:** Rare instances of severe allergic reactions or thrombosis were documented, requiring post-marketing surveillance.
- Long-term Safety: No significant long-term adverse effects identified in the majority of studied populations.

3. Global Distribution and Accessibility

- **High-Income Countries:** Achieved rapid vaccination coverage due to robust manufacturing, cold-chain infrastructure, and effective rollout strategies.
- Low- and Middle-Income Countries: Faced significant barriers including limited supply, cold-chain constraints, regulatory delays, and logistical challenges.
- Equity Concerns: Disparities in access highlight the importance of initiatives like COVAX and international cooperation to ensure global vaccine equity.

4. Public Acceptance and Uptake

- Vaccine Hesitancy: Factors such as misinformation, cultural beliefs, and mistrust in authorities contributed to variable uptake rates.
- Education and Communication: Clear public health messaging and engagement with community leaders improved acceptance and compliance with vaccination programs.

5. Comparative Insights Across Platforms

- mRNA vs Viral Vector: mRNA vaccines generally offer higher efficacy and rapid adaptability but require stringent cold-chain logistics. Viral vectors are less sensitive to storage but may have variable efficacy due to pre-existing immunity.
- **Protein Subunit/Nanoparticle Vaccines:** Offer stability advantages and moderate efficacy, making them suitable for broad distribution in resource-limited settings.

Synthesis

The results indicate that **novel vaccine platforms are effective and safe**, yet global distribution and equitable access remain major challenges. Integration of scientific innovation, robust logistics, public health policy, and community engagement is essential to optimize vaccination outcomes and control infectious disease outbreaks.

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Comparative Analysis: Vaccine Platforms

Parameter	mRNA Vaccines	Viral Vector Vaccines	Protein Subunit / Nanoparticle Vaccines	Implications
Efficacy	High (>90% against symptomatic infection)	Moderate (60–80%)	Moderate (70–85%)	mRNA vaccines provide highest protection; platform choice affects outbreak control
Safety Profile	Mostly mild-to- moderate adverse events; rare severe events	Mild-to-moderate; rare thrombosis or allergic reactions	Mild-to-moderate; minimal severe events	All platforms generally safe; monitoring for rare adverse events essential
Immune Response	Strong humoral and cellular immunity; adaptable to variants	Robust T-cell responses; efficacy may vary with pre- existing vector immunity	Good humoral immunity; stable, lower cold-chain requirements	Platform influences immune durability and variant coverage
Storage Requirements	Requires ultra-cold storage (-20°C to - 70°C)	Standard refrigeration (2–8°C)	Stable at standard refrigeration or room temperature	Cold-chain logistics critical for mRNA; protein-based vaccines better for low-resource regions
Manufacturing Scalability	High-tech, rapid production; resource-intensive	Moderate scalability; less resource-intensive	Moderate; easier production in low- resource settings	Scalability and infrastructure impact global distribution capacity
Global Distribution	High-income countries: fast rollout; LMICs: limited access	Broader distribution feasible; LMICs still face logistical challenges	Suitable for wide distribution; stable in resource-limited settings	Access depends on infrastructure, logistics, and equity-focused policies
Public Acceptance	High if properly communicated; hesitancy due to novelty	Moderate; concerns over viral vectors and misinformation	Moderate-to-high; familiar technology may reduce hesitancy	Communication strategies crucial for uptake across populations

☐ Summary:

This table illustrates that **mRNA vaccines offer superior efficacy and adaptability**, but require stringent cold-chain logistics, limiting distribution in resource-constrained regions. **Viral vector vaccines** balance efficacy with easier storage, while **protein subunit/nanoparticle vaccines** provide stability and suitability for global deployment, though with slightly lower efficacy. Effective vaccination programs require consideration of **platform characteristics**, **infrastructure**, **and public acceptance** to optimize global health outcomes.

SIGNIFICANCE OF THE TOPIC

The development and global deployment of vaccines are pivotal to controlling infectious diseases, improving public health, and preventing future pandemics. Understanding advances in vaccine platforms, efficacy, and distribution challenges is critical for several reasons:

1. Scientific and Clinical Relevance

- Innovative vaccine platforms such as mRNA, viral vectors, and nanoparticles have transformed the speed, efficacy, and adaptability of vaccine development.
- These technologies enable rapid responses to emerging pathogens and enhance immunogenicity, offering protection against severe disease and reducing mortality.

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2. Public Health Importance

- Vaccination programs are central to achieving herd immunity, controlling outbreaks, and reducing healthcare burdens.
- Understanding distribution challenges, cold-chain logistics, and equitable access ensures that vaccines reach diverse populations, particularly in low- and middle-income countries.

3. Policy and Governance Implications

- Global coordination and policy frameworks, including initiatives like COVAX, are essential for equitable distribution and prioritization of vulnerable populations.
- Lessons from recent pandemics highlight the need for adaptable regulatory pathways, international collaboration, and rapid deployment strategies.

4. Societal and Behavioral Significance

- Addressing vaccine hesitancy, misinformation, and cultural barriers is critical to achieving high immunization coverage.
- Public trust in vaccines enhances participation in immunization programs and strengthens community resilience against infectious diseases.

5. Research and Innovation Impact

- Studying vaccine platform performance and distribution effectiveness informs the design of next-generation vaccines.
- It supports evidence-based decision-making for policymakers, healthcare providers, and global health organizations.

Synthesis

The significance of this topic lies in its ability to **bridge scientific innovation**, **public health policy**, **and global equity**. Advancing vaccine technologies while addressing distribution and societal challenges ensures effective pandemic preparedness, equitable access, and strengthened global health security.

LIMITATIONS & DRAWBACKS

Despite significant advancements in vaccine development, several limitations and challenges persist that can impact effectiveness, accessibility, and public health outcomes.

1. Technological Limitations

- Cold-Chain Requirements: mRNA vaccines require ultra-cold storage, limiting deployment in resource-constrained regions.
- Rapid Variant Evolution: Emerging viral variants may reduce vaccine efficacy, necessitating booster doses or updated formulations.
- Manufacturing Constraints: High-tech production facilities are required for novel platforms, which can slow large-scale manufacturing.

2. Distribution and Logistical Challenges

- Equitable Access: Disparities exist between high-income and low-income countries due to supply limitations and infrastructure gaps.
- **Supply Chain Complexity:** Efficient global distribution requires robust cold-chain logistics, real-time monitoring, and contingency planning.
- Regulatory Variability: Differences in approval processes and import/export regulations can delay vaccine rollout.

3. Public Acceptance and Behavioral Barriers

- Vaccine Hesitancy: Misinformation, cultural beliefs, and mistrust in authorities can reduce uptake.
- Communication Gaps: Lack of clear public health messaging may hinder compliance and participation in immunization programs.

4. Economic and Policy Constraints

- **High Cost of Novel Vaccines:** Advanced platforms can be expensive, limiting access for low-resource populations.
- Funding and Resource Allocation: Limited resources may prioritize certain regions or populations over others, exacerbating inequities.

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5. Research and Knowledge Gaps

- Long-Term Safety Data: Limited longitudinal studies on newly developed vaccines restrict understanding of long-term effects.
- **Comparative Effectiveness:** Continuous monitoring is required to assess relative performance across vaccine platforms and populations.

Synthesis

While novel vaccine platforms offer high efficacy and rapid adaptability, **technological**, **logistical**, **societal**, **and economic challenges** can limit their global impact. Addressing these drawbacks requires integrated strategies combining robust logistics, equitable policy frameworks, public engagement, and ongoing research to optimize vaccine access, safety, and acceptance worldwide.

CONCLUSION

Advances in vaccine development, particularly through mRNA, viral vector, and protein subunit platforms, have transformed the prevention and control of infectious diseases. These novel technologies have demonstrated high efficacy, favorable safety profiles, and adaptability to emerging pathogens, representing a significant leap forward in biomedical innovation.

Despite these scientific achievements, substantial challenges remain in ensuring **global access, equitable distribution, and public acceptance**. Cold-chain requirements, manufacturing limitations, regulatory variability, and vaccine hesitancy continue to impede the effectiveness of vaccination programs, particularly in low- and middle-income countries. Addressing these issues requires coordinated international policies, investment in infrastructure, and targeted public health strategies to foster trust and compliance.

Furthermore, continuous monitoring of vaccine efficacy, safety, and performance against evolving pathogens is essential to inform future vaccine design and deployment strategies. Integrating scientific innovation with robust logistics, equitable policy frameworks, and effective communication ensures that vaccines achieve their full potential in safeguarding global health.

In conclusion, vaccine development is not only a scientific endeavor but also a public health and societal challenge. Success depends on harmonizing technological advances with equitable access, effective distribution, and public trust to strengthen global health resilience and prepare for future pandemics.

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