Interventions to Improve Access to Medicine in Developing Countries: Mapping WHO’s Building Blocks and Supply Chain Functions

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Abstract

Access to medicine remains poor and inequitable in many low- and middle-income countries (LMICs). This is a complex and multi-dimensional issue calling for holistic solutions. Studies in this area focus on singular disciplines, highlighting one or two main issues; this paper seeks to consider this issue from a multi-disciplinary perspective. It first enumerates the supply chain bottlenecks which lead to poor access to medicine. Since access is dependent on a host of factors, it is critical to understand each of these in the context of LMICs. Secondly, the paper proposes interventions to improve access by focusing on availability, affordability, quality and obtainability of medicine. These interventions are categorised into broader areas of focus to help stakeholders understand their role and responsibilities across the supply chain functions. Finally, the paper establishes a rationale for each intervention, matching it to a WHO Building Block and the corresponding supply chain management function. The resulting map will allow stakeholders to envision policies that will contribute to comprehensive solutions that strengthen the public health supply chains in LMICs.

Introduction

In 2015, the UN agreed that Sustainable Development Goals will include targets such as the elimination of major disease epidemics and the reduction of the burden of childhood obesity. The progress in global health is not inevitable and requires consistent efforts by countries, donors, policymakers, supply chain participants, etc. In 2017, the improvements in global mortality rates were less pronounced than in the previous decade, and non-communicable diseases (NCDs), for example, accounted for 73.4% of deaths, an increase of 22.7% since 2007 [4]. This was attributed to changing diets, urbanisation, and sedentary lifestyles. Moreover, inappropriate and overuse of medication coupled with wastage of resources...
created a gap between the demand and supply that will ultimately lead to a loss of social protection for the poor [5].

In addition, new public health challenges are putting further pressure on already strained health systems and contributing to a high volume of out-of-pocket expenses in LMICs. These challenges include supply chain impediments coupled with legislative and economic transitions taking place in different countries. The various stakeholders including the scientific research community, local governments, public health and regulatory agencies, overseas development agencies, philanthropists, multi-lateral agencies and the non-profit sector’s specific duties will contribute to the improvement of healthcare coverage.

In many respects, the cornerstones of public health growth are the pharmaceutical companies who should aim to ensure that medicine is available and accessible, irrespective of socio-economic considerations. These companies have the power to improve supply chains and save lives by providing better access to medicine at affordable prices. Through purposeful collaborations, the said companies can respond to the changing disease patterns, introduce new products and technologies, and promote sustainable, long-term access to medicine.

Emerging economies accounted for 23% of global spending on pharmaceuticals in 2015 and they are expected to account for 25% by 2020 [6]. Since exact predictions of diseases and outbreaks are not always possible, a collaborative approach between the public and private sector to develop, support and implement innovative practice is of utmost importance. The ideal solution to these challenges would be to introduce free provision of medicine to patients with low costs for governments. This would help to improve health benefits and decrease out-of-pocket expenditures. However, in spite of the attempts to support this approach, not much progress has been witnessed in LMICs [7].

Literature Review

Since the adoption of the Millennium Development Goals (MDG) in 2000 and the establishment of the Global Fund to Fight AIDS, Tuberculosis and Malaria (GFTAM) in 2002, there has been an increase in financial donations from multilateral and bilateral donors to LMICs [8,9], resulting in an exponential increase in the procurement of medical products. But the success of health programmes is not only dependent on the availability of funds to finance the procurement of medicine; if these medicines are inaccessible to the patients who require them, suffering and premature mortality will persist. Issues with access to medicine remain one of the leading causes of preventable deaths and poor health outcomes in LMICs.

Public health supply chains carry the responsibility of improving the health outcomes by addressing inaccessibility, which can put an additional strain on the already struggling complex public health supply chain. Despite increased donor funding and an array of new products (including vaccines and essential medicine), there are factors that continue to restrict access to health products. Namely, the capacity of a country’s supply chain to select, forecast, procure, and deliver health supplies is a major hindrance [10]. There is an urgent need for more research into public health supply chains in LMICs if any improvement in the access to medicine is to be made.

Supply chains and the corresponding management

Van Wassenhove [11], outlines three types of flows supported by a supply chain that require careful design and close coordination. Those include material flows, the physical product flows from suppliers to customers, including reverse flows for product returns, servicing, and recycling; information flows, such as order transmission and order tracking, which coordinate the material flows; and financial flows–credit terms, payment schedules, and consignment arrangements. Closely related to ‘supply chain’ is ‘supply chain management’ (SCM). To some, SCM is the coordination of the supply chain activities [12]. Others provide similar definitions but refer to ‘business functions’ rather than ‘activities,’ reflecting their view of the supply chain as a network of entities [13].

Health systems

The World Health Organization (WHO) defines a health system as ‘all the activities whose primary purpose is to promote, restore or maintain health’ [14]. The WHO 2007 framework for health systems is widely recognised, and its broad approach is the most suitable for identifying the range of issues that might affect the availability of medicine. It consists of six interconnected building blocks, required for an effective health system, namely, service delivery, health workforce, information, medical products, vaccines and technology, financing, and leadership and governance. According to Shakarishvili et al. [15], this approach is ‘a useful means for locating, describing, and classifying health system constraints, for identifying where and why investments are needed, what will happen as a result, and by what means the change can be monitored’ [15].

Building on the literature review, this paper examines and assesses the strategies for improving the accessibility of medicine LMICs. The paper highlights supply chain challenges throughout the process and the resultant effect on health systems. This paper recommends areas for holistic improvement. The paper will help to disseminate information and shorten the learning curve of all the stakeholders involved in trying to improve the health outcomes of any developing nation. The paper is structured as follows: Section 2 discusses the supply chain bottlenecks in achieving better medicine accessibility; Section 3 highlights interventions to increase healthcare coverage through improved access to medicine; Section 4 matches the interventions to WHO building blocks and supply chain management; and Section 5 concludes the paper.

Citation: Pamela Steele, et al. “Interventions to Improve Access to Medicine in Developing Countries: Mapping WHO’s Building Blocks and Supply Chain Functions”. Acta Scientific Pharmaceutical Sciences 3.7 (2019): 111-120.
Supply chain bottlenecks in achieving better medicine accessibility

The methods of procurement and distribution of medicine can vary across countries. However, supply chain issues and disruptions can interrupt the availability of medicine at the lower echelons of the population. Procurement delays can hamper the patients’ confidence in public health systems and defeat the basic purpose of public health supply chain. Delays can emerge due to supplier mismanagement, delayed tendering, customs issues, poor specifications and many more. Moreover, each country has its unique set of challenges which cannot be solved through generalised solutions. Therefore, understanding contextualised factors and avoiding non-customised approaches is a step towards better health outcomes.

Capacity planning and development is crucial throughout the supply chain functions [16]. LMICs often face skill-gaps which contributes to the fragmentation and inefficiency of public health systems. There continues to be a dire need for training and capacity development of existing staff [17]. The employment of procurement staff without the right qualifications and training further jeopardises the procurement system. Employees become vulnerable to audit and vigilance issues, leading to defensive office procedures. Furthermore, the distribution of medicine, in terms of last-mile delivery, can pose various challenges due to insufficient infrastructure and lack of legislative support.

The following table highlights the key supply chain challenges and the solutions required to overcome them in an LMIC context. Besides the challenges mentioned in Table 1, inherent systemic

<table>
<thead>
<tr>
<th>Supply chain functions</th>
<th>Challenges</th>
<th>Requirements</th>
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<td>Selection of medicine</td>
<td>• Poor selection of essential medicine</td>
<td>• Proper understanding of the Essential Medicine List</td>
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<tr>
<td></td>
<td>• Lack of standard practice</td>
<td>• Linking requirements to disease pattern of the country through proper data collection and feedback system</td>
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<td></td>
<td>• Lack of transparency</td>
<td>• Evidence-based decisions</td>
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<td></td>
<td>• Low human capacity to carry on this function</td>
<td>• Ability to forecast built on need and patient demand</td>
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<td>Quantification</td>
<td>• Inaccurate and less-reliable data on consumption</td>
<td>• Matching of procurement to the Essential Medicine List</td>
</tr>
<tr>
<td></td>
<td>• Poor quantification and forecasting skills</td>
<td>• Use of technology to reduce lead times and expedite procurement cycle</td>
</tr>
<tr>
<td>Procurement</td>
<td>• Unclear procedures</td>
<td>• Better supplier selection and management</td>
</tr>
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<td></td>
<td>• Procedural delays due to poor quantification</td>
<td></td>
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<tr>
<td></td>
<td>• Use of low/outdated technology</td>
<td></td>
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<tr>
<td></td>
<td>• Supplier uncertainty</td>
<td></td>
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<tr>
<td>Quality of the medicines</td>
<td>• Counterfeit drugs crowding the public health systems</td>
<td>• Strict quality checks and adherence to criteria</td>
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<td></td>
<td>• Treatment failures extending illness, adverse reactions, disability and death (IFPMA)</td>
<td>• Blacklisting of suppliers for substandard/spurious medicine</td>
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<td></td>
<td>• Detection is also difficult as the supply chain utilizes complex international trade routes, within a system where police, customs, and drug regulators are not unified (WHO)</td>
<td>• Sample testing at delivery points and along the supply chain by independent accredited laboratories.</td>
</tr>
<tr>
<td></td>
<td>• Use of modern technology</td>
<td>• Use of modern technology</td>
</tr>
<tr>
<td>Inventory management</td>
<td>• Mismatch between demand and supply</td>
<td>• Inventory decisions based on consumption data</td>
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<td></td>
<td>• Inadequate methods of inventory control leading to overstocking/understocking</td>
<td>• Real-time inventory data shared across the public health supply chains for better replenishment and distribution strategies.</td>
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<td></td>
<td>• Lack of human capacity</td>
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<td>Warehousing</td>
<td>• Poor conditions of storage facilities</td>
<td>• Adoption of a warehouse management system</td>
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<td></td>
<td>• Lack of adequate infrastructure (temperature control)</td>
<td>• Modern infrastructure for fast order processing, such as racking and mechanical handling equipment (MHE)</td>
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<td></td>
<td>• Low utilisation of technology</td>
<td>• Well-trained and skilled staff for warehouse operations; storage and security.</td>
</tr>
<tr>
<td></td>
<td>• Reduced security and human capacity</td>
<td></td>
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<tr>
<td>Distribution</td>
<td>• Low availability of good quality public transport and very expensive private fleet</td>
<td>• Using modern fleet management technology to improve distribution</td>
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<td></td>
<td>• Low frequency of distribution to lower tiers and rural areas</td>
<td>• Embracing GPS enabled last mile delivery for better health outcomes</td>
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<td></td>
<td>• Last-mile delivery failure due to inaccessible locations and transportation constraints</td>
<td>• Outsourcing, depending on the health commodities and the regions to be delivered.</td>
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<td></td>
<td>• Poor coordination between the central warehouses and health facilities</td>
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<tr>
<td>Support functions</td>
<td>• Poor funding</td>
<td>• Capacity development through workforce engagement and motivation</td>
</tr>
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<td></td>
<td>• Low human capacity</td>
<td>• Performance management tools to instil better feedback and structure</td>
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<tr>
<td></td>
<td>• Lack of clear procedures</td>
<td>• Improved relationship management supporting long-term alliances and support</td>
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<td></td>
<td>• Poor vendor management practice</td>
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</table>

Table 1: Supply chain challenges and requirements to be met to improve the availability of medicine in LMICs.

Citation: Pamela Steele., et al. “Interventions to Improve Access to Medicine in Developing Countries: Mapping WHO’s Building Blocks and Supply Chain Functions”. Acta Scientific Pharmaceutical Sciences 3.7 (2019): 111-120.
Interventions to increase healthcare coverage through improved access to medicine

The various supply chain challenges highlighted in Table 1 have a negative impact on the availability and accessibility of medicine at the last mile. Moreover, another reason for supply chain interruptions could be the diverse geographical topographies ranging from plains to difficult-to-access mountainous areas. Public health chains can become more effective through interventions to improve access to medicine focused on better availability, affordability, quality, and obtainability. The interventions will result in tools to improve healthcare coverage and achieve better health outcomes. They will also act as solutions to the various supply chain bottlenecks that cannot be solved in isolation [18,19].

Improving the availability of medicine

It is discouraging to realise that an overwhelming majority of medicine is designed for the needs of high-income countries. This results from a lack of incentives for research and development for medicine of the needs of LMICs. Pharmaceutical companies operate in a system influenced by the profits, where the consumers 'choose' from the portfolio of drugs produced through demand [20]. However, the purchasing power is very low or non-existent in LMICs. Again, lack of funds results in high out-of-pocket expenditure that represents a similarly high proportion of the population's income. The lack of health insurances aggravates the issue further. Another trend that affects the availability of medicine is the inequality in research and development of medicine. Pharmaceutical companies consider the needs of LMICs only when a specific disease pathway lead to them and forcing the stretch beyond low profit margins [21].

Some of the existing solutions embraced by countries to strengthen research and development to improve the availability of medicine are:

- **Grants:** Research grants are a common tool to boost research and innovation into this problem. Applications are usually initiated by international donors and agencies. The grants help to cover the initial costs, and they are a step towards improving health outcomes. Yet, such grants create dependency on the donors and their availability can be sporadic.

- **Treaties:** The implementation of a global research and development (R&D) treaty was recommended by the WHO Consultative Expert Working Group (CEWG) on Research and Development: Financing and Coordination. The treaty is supported by over 80 health research institutions, product development partnerships, and public interest NGOs [22]. These treaties are crucial for the growth of LMICs [23].

- **Market-based commitments:** Advance market commitments create an agreement in advance of the development of a product to purchase guaranteed amounts that meet criteria set by the donor (Commission on Intellectual Property Rights, Innovation and Public Health, 2006). Critics have opined that it is difficult to measure the success of these strategies because of the complexity and time involved [24].

  - **Priority Review Vouchers:** These provide incentives for research and development by shortening the review process of future medicine. However, investigation shows that these are not used widely [25].

  - **Product Development Partnerships:** These are initiatives to bring together public, private and funding organisations to provide an impetus for improving the availability of medicine through research and development [25]. This is an opportunity for developing nations to combine the efforts of the various stakeholders and work towards better health coverage. These partnerships try to strike a balance between public and private interests [26].

  - **Patent Pools:** These involve the cross-licensing of intellectual property by participants to increase access to essential technologies for products [26]. These pools act as a uniform platform for licensing of all patents and make the process of drug discovery more efficient. However, it should be noted that use of patent pools for pharmaceutical R&D is relatively new, and they have not been widely used [27].

However, the above-mentioned initiatives have not yielded many results in LMICs. Therefore, some of the interventions that can be adopted are

  - ** Tradable patent terms can be introduced, meaning that pharmaceutical initiatives serving a humanitarian purpose would receive an extended patent term that can be used with a different pharmaceutical product [28].**

  - **Pharmaceutical companies are urged to begin systematically planning ahead during clinical development to ensure that successful products can be made widely available more quickly in developing countries.** They can pioneer voluntary licensing and establish multi-sector capacity development partnerships [29].

  - **Refocusing the direction of the major stakeholders may be needed to boost meaningful research and development in LMICs. Pharmaceutical companies need to take more responsibility, allocate time and resources to improve healthcare in LMICs [30]. Priorities must span across diseases, conditions and pathogens and the different products needed per disease, such as medicine or diagnostic tools.**

  - **State-run programmes can ease off the pressure from private sector. This will also induce accountability into the public sector for the availability of medicine.**

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Differential pricing can be adopted allowing LMICs to pay for medicine based on their ability and per capita household income.

Improving the affordability of medicine

The affordability of medicine is the ability of the population to pay for medicine without compromising other essentials, such as housing, education and food. Affordability also depends on who is paying—the patient or another stakeholder in the local healthcare system. In LMICs nearly 70% of spending on medicine is out-of-pocket. Different countries have different income capacities and these differences should be kept in mind while deciding the prices of medicine. The cost of medicine is usually determined by the cost of manufacturing, supplying, and the research involved. The costs often do not include the opportunity cost and are inflated to reap higher profits. Studies have highlighted that efforts to make medicine more affordable are lacking in LMICs by comparison to developed countries [31], mainly due to lack of funding [32]. LMICs do not have efficient systems for clinical trials, top-level medical schools and experienced clinicians [33] which hinders access to affordable medicine (IFPMA). Again, affordability of medicine is influenced by huge mark-ups resulting in poor healthcare coverage.

Some of the existing tools embraced by countries to improve affordability by reducing the cost of medicine are:

- Differential pricing, which occurs when varying prices are charged by the seller according to the purchasing power of governments and households in different countries [34]. However, it involves a high risk of arbitrage and erosion of high-income markets [35].
- Monopsonies are market structures where the buyer has the control power. In many countries, when public procurement is done by the government, it functions as a monopsony, allowing for the negotiation of lower prices (e.g.: PHARMAC in New Zealand). This kind of a market structure can help to reduce the cost of the medicine, generating better savings and allowing affordable health treatment. On the other hand, this model can suffer from lack of transparency and knowledge, since most contracts are confidential [36].
- Patent law flexibilities can include changing the terms of patents, strengthening patent criteria and introducing competition [37].

As with availability, the previous initiatives have not yielded visible results in LMICs. Therefore, some of the interventions that can be adopted to improve affordability of medicine could be:

- Increased transparency between the pharmaceutical sector and the country government to understand contextualised needs and challenges. Pharmaceutical companies can work with national governments and partners to expand pricing schemes and donation programmes.
- LMICs must have the political will to increase affordability, and the capacity and financing to develop policy and effective systems to that effect.
- Backdoor deals and evergreening need to be stopped completely. Backdoor deals occur when a monopolist incentivises a local company to stay away from the market [38]. Evergreening refers to penetration of secondary patenting of the most successful medicine, which may further reduce the ability of generics to enter the market [39].
- Time-bound targets for new products in LMICs will help to secure market access and grow a strong market share.
- It is necessary to enforce obligations on pharmaceutical firms to ensure that life-saving medicine is provided in LMICs [30]. This can be achieved by prioritising research and development on disease gaps.
- Socio-economic factors are to be frequently considered while setting medicine prices. The factors considered the most relevant are disease burden, healthcare system and financing; and the level of economic and human development. This will help in creating patient access profiles and facilitate better affordability.
- Stakeholders should explore ways of mobilising new and innovative means of financing, developing and increasing resources available for health.

Improving the quality of medicine

Inferior quality medication and counterfeit products have a negative impact on the health systems of any country, especially of LMICs, where some disease reports indicate that half of the drugs may have few to no active ingredients [40]. Medicine may also contain toxic substances which are harmful for the community. These issues can cause treatment failure, extended illness, adverse reactions, disability and death [41]. Substandard medicine often looks very similar to the original and it is difficult to differentiate them. The lucrative nature of these products often fosters cross-border criminal activities, making detection and prosecution difficult. The counterfeits attract the various stakeholders alike and lack of proper legislation, enforcement and heavy penalties make deterrence very limited [42]. Detection is also made difficult by the spread of supply chains across different countries and continents, meaning that they are not unified in a single system [43]. Another factor that impacts on the quality of medicine is that legally produced generics legally produced may be confiscated (en route) on grounds of IP breach even, though the destination country could legally distribute it [44].

Currently, the efficacy of medicine is generally checked, and the desired quality achieved through the following initiatives:
Interventions to Improve Access to Medicine in Developing Countries: Mapping WHO’s Building Blocks and Supply Chain Functions

- Product Authentication uses colour shifting inks, holograms and chemical markers embedded in drugs to identify legitimate medicines [45].
- E-coding is another method followed by many countries to help identify substandard drugs. For example, China uses an 'e-coding' system whereby each party within the supply chain is required to send a 'signal' to confirm receipt or dispatch to a regulatory database [46].
- Track and trace devices and technology.
- National enforcement of strict sanctions and penalties.

These general measures have not yielded the results LMICs. Therefore, some of the interventions that can be adopted to improve quality of medicine are:

- Educating patients and healthcare workers on the accurate identification and use of the medicine through means that meet a range of needs, including language, literacy and cultural, demographic and environmental needs.
- Creation of international marking systems, so that medicine does not need to be re-marked when entering a new country, saving supply chain costs and lead times.
- Stakeholders should have policies, procedures and resources in place to carry out effective drug recalls to protect the public from a defective or potentially harmful product.
- Sharing of information and communication by identifying emerging markets, particularly, when it comes to the implementation of segmented pricing and product registration filings, and to transferring knowledge, expertise and other capacities to the local manufacture of pharmaceutical products.
- Inclusion of LMICs in international frameworks.
- More research regarding substandard medicine is needed as the prevalence of substandard medicine is still the result of ‘informed guesses’ [40].
- An international code of practice to aid in coordinating regulatory, customs and law enforcement agencies.

Improving obtainability of medicine

In public health, the availability of medicine is considered a crucial issue. However, most of the studies overlook obtainability, since it is determined by a host of factors both within and outside the country’s control. Obtainability can be influenced by challenges in the country’s health system, infrastructure, legal barriers, socio-cultural influences, etc. [47]. This leads to patient unawareness and adherence to non-standardised practice. Again, the availability of healthcare personnel per capita is very low in LMICs, reducing the possibility of preventative medicine and diagnosis. Doctors and other healthcare personnel may be concentrated in urban areas, leaving rural populations without easy access to basic healthcare. In rural areas, it may be very time-consuming or expensive to reach the nearest health point or distribution facility [48]. It is also observed that the wholesale market is excessively fragmented with poor traceability [47]. Even, if the medicine makes it to the local community centre, access may be physically restricted due to lack of basic infrastructure.

The existing initiatives to overcome these challenges are:

- Task-shifting is a tool embraced to develop the existing human capacity and provide a more sustainable solution. This process involves the movement of specific tasks from highly qualified workers to those with less training and fewer qualifications, in order to make more efficient use of the available human resources for health [49].
- Interventions through grassroots organisations (GROs), made up of interested parties coming together for self-help (as opposed to interested parties coming together to help others, as tends to happen in the sector) [48].

Once again, these initiatives have not yielded adequate results in LMICs. Some interventions that can be adopted to improve the obtainability of medicine are:

- Acceptance and identification of supply chain issues which are resulting in poor rates of medicine obtainability at the last-mile. This can be achieved through capacity planning and development at the various levels. Initiatives in this area range from training on good distribution practice, proper warehousing, forecasting and cold chain requirements, to projects that use technology to track stock and prevent stockouts.
- Strong political will to support the public health systems and supply chains to overcome bottlenecks by creating more distribution hubs, even in rural areas.
- Tailored mechanisms for change using contextualised and issue-driven technology, avoiding the blind adoption of modern technology without recognition of grassroot realities.
- Aligning global, regional and country-level supply planning processes with demand for the products by making efforts to understand product distribution and demand behaviour in different countries and applying the information for timely supply.
- The socio-economic determinants of health must be recognised in order to define a holistic framework.

Improving access to new medicine

It is critical for all LMICs to have access to new drugs and medicine. This can be done through proper advance planning on the part of the various stakeholders and through a clear understanding of the access plan. Pharmaceutical companies can voluntarily enter licensing agreements under pro-access terms to facilitate generic entry [50]. This can be achieved through planning for and aligning with other actors along supply chains to ensure the timely supply of good quality products and prevent stockouts.

Planning can also be done through submitting products to WHO’s prequalifications process to allow for UN procurement and accelerate the registration process in countries with weak national regulatory authorities [51]. Access plans can also be implemented through local partners to make the new products widely accessible. Generally, access plans first take the shape of commitments made

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during the early stages of development before being turned into concrete strategies or agreements. This is a necessary first step, and efforts must be made to ensure these commitments are turned into action.

Matching the interventions to WHO building blocks and supply chain functions

It is well-recognised that access to medicine is not only a matter of life and death; it also enhances the quality of life and it is important for overall improved standards of living. Access to medicine is a complex and multi-dimensional issue calling for holistic solutions. Measures must include improving supply chains, understanding social determinants of health, encouraging policy coherence, implementing proactive intellectual property regimes, and ensuring that health delivery systems are appropriate to those they serve. The following table summarises the proposed interventions and their areas of focus. We have identified the following areas of focus:

- **Analytical/Scientific:** These interventions help in generating new ideas and foster innovations to improve access to medicines.
- **Structural:** These interventions establish processes, procedures, and platforms to enhance and facilitate activities.
- **Collaborative:** These interventions enhance co-ordination among the various stakeholders and technical partners.
- **Legislative:** These interventions aim to bring changes in the legislative frameworks and boost political will.
- **Workforce development:** These interventions facilitate building the capacity of the workforce.

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<thead>
<tr>
<th>Access to medicine</th>
<th>Interventions</th>
<th>Area of focus</th>
<th>WHO building block</th>
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</thead>
<tbody>
<tr>
<td>Improve the availability of medicine</td>
<td>Tradable patent</td>
<td>Analytical/Scientific</td>
<td>Information and Research</td>
</tr>
<tr>
<td></td>
<td>Systematically planning during clinical development</td>
<td>Structural</td>
<td>Information and Research</td>
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<td></td>
<td>Refocusing of the direction of the Major stakeholders</td>
<td>Collaborative</td>
<td>Leadership and Governance, Service delivery</td>
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<td></td>
<td>State-run programmes</td>
<td>Legislative</td>
<td>Leadership and Governance, Service delivery</td>
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<td></td>
<td>Differential pricing</td>
<td>Structural</td>
<td>Finance</td>
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<tr>
<td>Improve affordability of medicine</td>
<td>Increased transparency between the pharmaceutical sector and the country government</td>
<td>Collaborative and Legislative</td>
<td>Leadership and Governance</td>
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<td></td>
<td>Political will</td>
<td>Legislative</td>
<td>Leadership and Governance</td>
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<td></td>
<td>Backdoor deals and evergreening need to be stopped</td>
<td>Collaborative and Legislative</td>
<td>Leadership and Governance</td>
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<td></td>
<td>Time-bound targets for filling to register new products</td>
<td>Analytical/Scientific</td>
<td>Information and Research</td>
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<td>Setting priority research and development account to focus on disease gaps</td>
<td>Analytical/Scientific</td>
<td>Information and Research</td>
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<td></td>
<td>Mobilizing new and innovative means of financing</td>
<td>Structural</td>
<td>Finance</td>
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<td>Socio-economic factors</td>
<td>Structural</td>
<td>Leadership and Governance</td>
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<tr>
<td>Improving the quality of medicine</td>
<td>Educating patients and healthcare workers</td>
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<td>Health Workforce</td>
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<td>International marking systems</td>
<td>Collaborative, legislative</td>
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<td>Effective drug recalls</td>
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<td>Information and Research</td>
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<td>Sharing of information and communication</td>
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<td>International frameworks and code of practice</td>
<td>Legislative</td>
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<td>Research regarding substandard medicines</td>
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<td>Information and Research</td>
</tr>
<tr>
<td>Improving the obtainability of medicine</td>
<td>Capacity planning and development</td>
<td>Workforce development</td>
<td>Health Workforce</td>
</tr>
<tr>
<td></td>
<td>Strong political will to support the public health systems and supply chains</td>
<td>Legislative, Structural</td>
<td>Leadership and Governance, Service delivery</td>
</tr>
<tr>
<td></td>
<td>Tailored mechanisms for change</td>
<td>Collaborative</td>
<td>Leadership and governance</td>
</tr>
<tr>
<td></td>
<td>Aligning global, regional and country-level supply planning processes with demand</td>
<td>Structural</td>
<td>Service delivery, Information and Research</td>
</tr>
<tr>
<td></td>
<td>Socio-economic determinants of health</td>
<td>Structural</td>
<td>Leadership and Research, Service delivery</td>
</tr>
</tbody>
</table>

Table 2: Proposed interventions and the area of focus in developing nations.
We also link each intervention to a WHO Building Block (refer to Table 2) and the corresponding supply chain management functions (refer to Table 3). This mapping will support stakeholders get a broader picture and envision policies, which will contribute towards comprehensive solutions to strengthen the public health supply chains in developing nations. Establishing better rationality to decisions and building it on relevant constraints will support developing nations propel towards stronger health systems and public health supply chains.

<table>
<thead>
<tr>
<th>WHO Building Block</th>
<th>Supply chain functions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Information and Research</td>
<td>Selection of medicines, quantification, procurement, inventory management</td>
</tr>
<tr>
<td>Leadership and Governance</td>
<td>Policies, co-ordination, transparency, accountability, change management, stakeholder management</td>
</tr>
<tr>
<td>Finance</td>
<td>Funding (Donor and Government), drug revolving fund, investments</td>
</tr>
<tr>
<td>Health Workforce</td>
<td>Staffing, skills, supervision, performance management, working conditions, motivation, incentives</td>
</tr>
<tr>
<td>Service delivery</td>
<td>Procurement, warehousing, distribution tools, transportation, technology, last mile delivery</td>
</tr>
</tbody>
</table>

Table 3: WHO Building Blocks and supply chain functions.

Conclusion

Access to medicines requires efficient public supply chains where each component of the cycle is aligned with the performance of the other. If one component is not managed correctly, other components are bound to be adversely impacted. With an efficient system in place, procurement, and distribution of appropriate medicine and equipment, rational diagnostic, therapeutic practices can be promoted in LMICs. The integration of strategies into contextualised frameworks will create a sustainable roadmap for better access to medicine. Developing nations are the new frontiers for growth, and hence provide opportunities to improve the quality of human care and innovations. This creates a platform for the various stakeholders to enter, understand, and develop effective policies to have a more positive and significant effect on the majority of the population. Interventions to improve universal healthcare through better access to medicine will benefit LMICs with improved health systems and solutions to local needs. Since the disease patterns and the strength of public health systems differ across countries, tailored interventions which incorporate training for healthcare workers and health financing can help overcome country-specific barriers. Inclusive involvements will aim to include people living on very low incomes to improve access to specific medicine or other health products. This will make a tremendous difference and provide medicine which the poor people cannot afford and at the same time reduce the out-of-pocket expenditure.

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