TRIPS, R&D AND ACCESS TO MEDICINES:

A GUIDE
TO THE POST 2005 WORLD

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Médecins Sans Frontières (MSF) Campaign for Access to Essential Medicines

Based on a January 18th 2005 presentation to Members of European Parliament by Ellen ‘t Hoen

The magnitude of the AIDS crisis has drawn attention to the fact that millions of people in the developing world do not have access to the medicines that are needed to treat disease or alleviate suffering. Each day, close to eight thousand people die of AIDS in the developing world. The reasons for the lack of access to essential medicines are manifold: logistical supply and storage problems, substandard drug quality, inappropriate selection of drugs, wasteful prescription and inappropriate use, inadequate production, prohibitive prices and lack of financing for health care.

In many cases, however, high drug prices are the main barrier to needed treatments. Prohibitive drug prices are often the result of strong intellectual property protection. Governments in developing countries that attempt to bring down the price of medicines have come under pressure from industrialised countries and the multinational pharmaceutical industry. For instance, in 2001, 39 drug companies took the South African government to court over its medicines act designed to provide South Africans with affordable drugs. More recently, Guatemala has come under pressure to implement "TRIPS-plus" data protection rules.

The 1995 World Trade Organization (WTO) Trade-Related Aspects of Intellectual Property Rights (TRIPS) Agreement sets out minimum standards for the protection of intellectual property, including patents on pharmaceuticals. These standards derive from wealthy Western nations and are not necessarily appropriate for developing countries. The TRIPS Agreement has come under fierce criticism for this "one-size-fits-all" principle because of the effects of increased levels of patent protection on drug prices.

MSF has witnessed the impact of patents on the prices and availability of medicines, in particular newer medicines, and has documented the patent practices in the countries where it works¹. It should be no surprise that patent protection translates into high drug prices: patents create monopolies and monopolies lead to higher drug prices. As soon as the monopoly ceases to exist, prices come tumbling down. Figure 1 on page 2 shows the effect of generic competition on the price of first-line AIDS triple therapy.

¹ "Drug patents under the spotlight - Sharing practical knowledge about pharmaceutical patents." Médecins Sans Frontières (MSF), June 2004.
In addition to their impact on prices, patents may also hamper the development and availability of recommended formulations. An example is the problem of developing fixed-dose combinations (e.g. the "three-in-one" pill for AIDS treatment) when the patents of the individual components are held by different companies. These fixed-dose combinations are particularly important in AIDS treatment. Some recommended fixed-dose combinations are now available from Indian producers because until 2005 pharmaceutical product patents were not granted in that country and therefore did not create a barrier to formulating these products.

Why do we have patents?

The patent system is a social policy tool which aims to stimulate innovation. The idea is that by providing limited exclusivity to the "inventors" of products, which comes at a price, innovation will be promoted and society as a whole will benefit from the availability of new and improved products.

Patents and Research and Development (R&D)

However, a major problem of the current patent system is the imbalance between rights and obligations: the patent system is intended as a stimulus for innovation, but there is no mechanism for directing that innovation, and as a result many diseases are totally ignored. Drug R&D, which is almost exclusively confined to the private sector, is skewed towards areas that promise a profitable return. This is a logical consequence of the patent-driven R&D mechanism our societies rely on these days. Thus in the last 25 years, almost 1,400 new medicines have been developed, but only 1% of these were for tropical diseases (see figure 2). These diseases
kill tens thousands of people every year, but because they occur mostly in the developing world, they do not represent a profitable market for industry.²

**Figure 2. Drug Development Outcome**

1975-1999: **1393** new chemical entities marketed

- Tropical diseases: 13
- Tuberculosis: 3

Patent protection has increased over the last 20 years, but the mean innovation rate has fallen, with an increase in the number of ‘me-too drugs’ of little or no therapeutic gain, as shown in figure 3 below. This global crisis in innovation has a disproportionately heavy impact on the needs of people in developing countries.

**Figure 3. Innovation in France 1981-2001**

Adapted from Prescrire International, April 2001/Vol 10, n° 52 p 54

By adopting the Doha Declaration on TRIPS and Public Health in 2001, the WTO recognised some of the concerns raised by developing countries regarding access to medicines. The Doha Declaration lays out the flexibilities contained in the TRIPS agreement which countries can use to overcome the barriers posed by patents. It also extends the "transition period" - during which Least Developed Countries (LDCs) are not obliged to enforce or grant patents on pharmaceuticals products - until 2016.

However, in recent years, we have seen a systematic dismantling of the Doha Declaration through bilateral trade agreements with the United States, which include so-called "TRIPS plus" provisions: these annul the achievement of Doha and confirm the lack of political support for the use of TRIPS flexibilities.

**Access to medicines in the post 2005 world**

Following the full implementation of the TRIPS Agreement as of January 1st 2005 in India and the few other developing countries not yet granting pharmaceutical patents, access to affordable new drugs is expected to become more difficult. For example, most of the ARVs currently available at affordable prices come from India. Successful AIDS programmes such as those of Brazil and Thailand were possible because key pharmaceuticals were not patent-protected and could be produced locally at much lower costs.

From 2005 onwards, all new drugs may be subject to at least 20 years of patent protection in all but the least developed countries and the occasional non-WTO country such as Somalia, Palestine or Macedonia. A number of developing countries that are presently scaling up AIDS treatment have expressed their concern to the World Health Organization about the effects of TRIPS implementation in India.³

Because TRIPS implementation will affect both producers in key manufacturing countries and countries that are dependent on these manufacturers for raw materials, prices will be kept high and new medicines will be made inaccessible for the majority of the population in developing and least developed countries. Generic producers will also be blocked from developing fixed dose combinations until the relevant patents on the individual components of the combinations expire. In other words, access to essential medicines could become dramatically more difficult in the coming years if no action is taken.

Faced with these new challenges, the public health safeguards affirmed in the Doha Declaration, such as compulsory licensing or government use, will become even more important. It is imperative that producing countries such as Brazil, Thailand and India routinely make use of compulsory licenses or "government use" provisions, including allowing the export of these medicines, to enable generic competition to drive prices down. Strong political resolve will be needed to do this.

**Production and export of generic medicines**

In 2003, the WTO adopted the "August 30th decision" which allows the export of medicines produced under a compulsory license – this is restricted in the TRIPS agreement by the requirement that a compulsory license be ‘predominantly for the domestic market’. The August 30th solution is needlessly complex, and is not likely to remove the real threat of dwindling

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³ Letter from Dr Jim Kim, Director WHO HIV/AIDS department to the Minister of Health of India dd. December 17th 2004.
generic production in countries such as India. The mechanism is based on a drug by drug, country by country and case by case decision-making process which ignores the fact that economies of scale are needed to attract interest from producers. Without the pull of a viable market for generic pharmaceutical products, manufacturers cannot rationally be expected to want to take part in the production for export system.

Certain countries with production capacities have taken the initiative to implement the decision. Given the complexity of the WTO solution, one would expect that the implementation by potential exporters would at least be straightforward, without introducing extra barriers.

Unfortunately, this is not necessarily the case. For example, in Canada, the implementation of the August 30th decision contained limitations that were rejected by WTO Members at the time of negotiating the solution, such as a list of eligible countries, as well as a limited list of approved medicines that can be produced and exported in generic form to developing countries. But the medicines list does not include the fixed-dose AIDS drug combinations which are recommended by WHO and are vital for scaling up AIDS treatment in developing countries. Although it is foreseen that the list can be reviewed, the Canadian experience shows that new medicines have been excluded from the list following lobbying from the drug industry. For example, the company Bayer successfully lobbied to keep its pneumonia therapy, moxifloxacin, off the list of medicines.

Even if these issues are resolved, implementation in good faith of a text that is basically flawed cannot possibly yield real solutions.

Regrettably, these hollow measures are often hailed as great progress, and the public and policymakers are led to believe that access problems have been resolved, that affordable medicines will now become available and that no further action is needed. Such an approach would be disastrous.

Once again, the AIDS crisis shows us why. First-line triple therapy is now available for as little as US$140 per patient per year. But resistance to first-line ARVs is as inevitable in poor countries as it is in rich countries. When patients need to switch to second-line treatment, they will face treatment costs as high as US$5,000 per patient per year.

<table>
<thead>
<tr>
<th></th>
<th>3TC/d4T/NVP (1st line)</th>
<th>TDF+ddI+LPV/r (2nd line)</th>
<th>2nd line vs 1st line</th>
</tr>
</thead>
<tbody>
<tr>
<td>Western country6</td>
<td>US$8773/year</td>
<td>US$13151/year</td>
<td>1.5 times more expensive</td>
</tr>
<tr>
<td>Developing countries</td>
<td>US$154/year</td>
<td>US$3950/year</td>
<td>26 times more expensive</td>
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<tr>
<td>Cipla Triomune7</td>
<td></td>
<td>Originator products</td>
<td></td>
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<tr>
<td>Reduction</td>
<td>- 98 %</td>
<td>- 70 %</td>
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4 A country in need of a certain product which is not available (for example because it is not marketed, or because the price is too high) will have to inform the WTO about its intention to import and indicate the type of product and quantities needed and - if it is not for governmental use and / or an emergency - will have to seek a voluntary license from the patent holder(s). A potential producer in an exporting country needs to be identified. This producer - assuming there is one willing to invest in production for a limited market (the quantities need to be defined beforehand based on the request from one or several countries) - must then request and obtain a compulsory license from its national authorities, adapt its production line and capacities, and pay royalties to the patent-holder.


6 Australian EXW prices: "Schedule of Pharmaceutical Benefits for Approved Pharmacists and Medical Practitioners, May 2004. Exchange rate used for conversion (1Australian $=0.72213 US$, May 1, 2004)

7 Clinton Foundation price (FOB) + 10% due to transportation and importation taxes.
This discrepancy needs to be tackled urgently. The price of first-line drugs came down dramatically because countries that did not grant pharmaceutical product patents were able to produce generics and stimulate competition. The challenge will be much greater for second-line drugs. Now that key manufacturing countries will no longer be able to produce generic versions of new drugs, bringing down the price of a single source product is going to be much more difficult.

**Sources of affordable versions of new medicines drying up**

While the 2001 Doha Declaration on TRIPS and Public Health offers measures to access existing generics, much more needs to be done to ensure the production of generic second-line drugs.

In the post 2005 era where all drugs may be patented in most countries in the world, a lot more action will need to be taken to ensure that drug prices are set at a level the people and communities who need them can afford. Essential medicines are not a luxury whose availability can be left to private market forces only, but an essential component of the fulfillment of the right to health.

It is easy to get lost in the legal details, but it is crucial not to lose the human picture in this discussion. The fact is that effective medicines that dramatically increase the life expectancy of people living with AIDS became available in Europe and North America a decade ago. Today, 40 million people in the developing world are infected with HIV, and six million people need access to these medicines NOW. Only 700,000 do. The result is that, in the next 24 hours, another 8,000 people will have died of AIDS.

**Recommendations for political decision makers**

The way in which medicines are researched, developed and sold today leads to grave inequities. Global rules that affect the R&D and availability of medicines should be driven by health needs rather than industrial or commercial considerations. Faced with the rise of infectious diseases such as AIDS, TB, and malaria, and the increasing marginalisation of health problems that do not affect the developed world, strong voices are needed now more than ever to defend global public health.

Ensuring access to the fruits of innovation for even the poorest patients and promoting health R&D as a global public good requires global action. We ask political leaders worldwide to put R&D for neglected diseases at the top of their agenda and ensure sufficient, sustainable and long-term financing to address the R&D needs and work towards a change in the way health R&D priorities are set and financed.

We urge governments to address the dismantling of the Doha declaration, which is advancing insidiously through US-initiated Free Trade Agreements. Strong political support and technical assistance is needed for countries that use the TRIPS flexibilities. All producing countries must ensure that their patent policies allow the production and export of generic versions of newer medicines.

We urge the international community and political leaders to come together and to step up the effort to ensure access to new essential medicines now that medicines may be patented everywhere in the world. Without extra effort we will rapidly lose the progress that has been made in the area of access to medicines. We only need to look at the prices of second-line AIDS treatment to see how urgently action is needed.