Fatal Imbalance
The Crisis in Research and Development for Drugs for Neglected Diseases
Médecins Sans Frontières
Access to Essential Medicines Campaign

Médecins Sans Frontières (MSF) is an independent medical humanitarian organization committed to providing medical assistance to people in need regardless of race, religion, politics or gender, and to raising awareness of the plight of the people it helps. Too often, MSF volunteers, who work in over 80 countries worldwide, are left without adequate treatment options when the only available drugs are archaic, ineffective or toxic. To address this chronic emergency, MSF launched the Access to Essential Medicines Campaign in 1999.

Successes of the Campaign include raising awareness internationally of the access crisis, contributing to the dramatic fall in prices of antiretroviral drugs for the treatment of HIV/AIDS, securing a supply of discounted second-line drugs for multi-drug resistant tuberculosis, and ensuring the long-term production of four drugs for sleeping sickness. The Campaign has also helped put the problem of access to essential medicines on the international agenda.

The Drugs for Neglected Diseases Working Group

In October 1999, a group of concerned scientists, health professionals, and representatives from non-governmental organizations, the pharmaceutical industry, developing country governments and international organizations met in Paris to discuss stimulating the development and securing the availability of drugs for neglected diseases. Médecins Sans Frontières, the World Health Organization and the Rockefeller Foundation convened the meeting.

Following the meeting, the Drugs for Neglected Diseases (DND) Working Group was formed to continue the work begun at the conference by developing new ideas to restart research and development (R&D) of drugs for neglected diseases. The DND Working Group is a multi-disciplinary and independent group that includes researchers, drug development experts, and regulatory affairs professionals from the public and private sectors of developed and developing countries.

According to the DND Working Group mission statement, “It is the responsibility of society to address this public health failure, and seek new and creative strategies to solve this problem. Solutions and recommendations need to be sustainable, affordable, need-driven and involve input and active engagement of developing countries.”

The DND Working Group has studied the causes and proposed solutions for the R&D crisis. The group has also advocated for the active engagement and financial support of governments, private enterprises, foundations and international organizations to compensate for the failure of the market to provide drugs for neglected diseases. The work of the group focuses on the most neglected diseases, such as sleeping sickness and leishmaniasis, in addition to neglected diseases that are already receiving some renewed attention, such as tuberculosis and malaria. Close links have been established with other institutions such as the Special Programme for Research and Training in Tropical Diseases (TDR), which is based at the World Health Organization, and the Global Alliance for Tuberculosis Drug Development.

Part of the DND Working Group strategy is to fund and manage pilot drug development projects. The DND and TDR are working together to undertake several drug development projects that have not been completed due to lack of funds and human resources. Funding for these pilot projects is being provided partially by MSF and will be managed by drug development experts.

About the authors

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In spring 2001, the 20 top-grossing pharmaceutical companies in the world were surveyed about recent drug development activity. While the survey demonstrated some activity in neglected diseases, it indicated that private sector investment in this field was minimal. None of the responding companies has brought a drug to market in the last five years for any of the most neglected diseases included in the survey.

The DND Working Group has also explored the failure of the public sector to take a needs-based approach to managing drug development. Basic research leading to discovery of compounds—and thus potential drugs—has almost always been publicly funded. However, because politicians naturally respond to the needs of their constituencies, and because wealth is concentrated in industrialized countries, research money goes to the diseases primarily affecting these wealthier constituencies. While some government money has been devoted to diseases affecting developing countries, it is a pittance compared with overall spending on drug development. Private philanthropy has in recent years sought to fill in a bit of this gap, but it is not sufficient and cannot and should not take the place of public support.

Recent initiatives and policies seeking to redress the R&D imbalance are also outlined here. Public-private partnerships have been successful in mobilizing public and private sector expertise around certain diseases. Yet, to date, none of these provides an adequate strategy for developing drugs for the most neglected diseases.

Finally, recommendations for moving forward are presented, among them: that a well-defined and needs-driven research agenda be established at a global level; that governments fulfill their responsibility to become directly and proactively involved in searching for solutions; that funding be increased for research into neglected and most neglected diseases; and that a new not-for-profit initiative be explored as one way to address the shortage of R&D for the most neglected diseases.
The past 30 years have witnessed unprecedented transformations in global health, with, for example, life expectancy around the world rising by an average of four months every year. However, such impressive statistics should not obscure the fact that the benefits of the “global health revolution” have not been distributed evenly. Millions continue to die each year of preventable and treatable diseases. Communicable diseases killed 14 million people worldwide in 1999, mostly in developing countries. One cause of this is the vacuum in research and development (R&D) for medicines to treat the diseases of the poor.

There is a strong link between poverty and health. People from low- and middle-income countries carry a disproportionally high burden of disease, particularly with regard to communicable diseases. Those living in absolute poverty (on less than one dollar per day) are five times more likely to die before reaching the age of five, and two and a half times more likely to die between the ages of 15 and 59. Infectious and parasitic diseases account for 25% of the disease burden in low- and middle-income countries, compared to only 3% in high-income countries. According to the World Bank, eliminating communicable diseases would almost completely level the mortality gap between the richest 20% of the world population and the poorest 20%.

The vacuum in drug R&D for the diseases of the poor

Eliminating the mortality gap will likely remain an elusive goal, because R&D efforts are not addressing many of the communicable diseases that plague developing countries. An analysis of drug development outcomes over the past 25 years shows that only 15 new drugs were indicated for tropical diseases (11+2) and tuberculosis (2). These diseases primarily affect poor populations and account for 12% of the global disease burden. In comparison, 179 new drugs were developed for cardiovascular diseases, which represent 11% of the global disease burden (Figure 1A).

Too little money is going into health research that addresses the needs of the world’s poorest people. While it might be expected that health research would concentrate on the areas where the needs are greatest, the reality is quite different. Only 10% of global health research is devoted to conditions that account for 90% of the global disease burden—an imbalance that has been referred to as the 10/90 disequilibrium.

Heavy reliance on an increasingly consolidated and highly competitive multinational drug industry to generate new medicines has left the development of lifesaving drugs subject to the forces of a market economy. Currently, it is largely purchasing power that is defining research agendas and priorities, which means that poor people’s health needs are not being met.

This failure does not rest exclusively on the shoulders of the private sector. Governments hold the ultimate responsibility for ensuring that people’s basic health needs are met. They have the responsibility to take appropriate action when market forces fail to address these needs. In the past few decades, despite clear evidence of waning...
Tropical diseases are good examples of neglected diseases. A seriously disabling or life-threatening disease can be An examination of current research efforts in the pharmaceutical industry reveals that the pipeline of drugs for neglected diseases is virtually empty (see Figure 1C). In spring 2001, the DND Working Group and the Harvard School of Public Health sent written questionnaires to the world’s top 20 pharmaceutical companies to assess the level of R&D activity in several neglected diseases (sleeping sickness, leishmaniasis, Chagas disease, malaria and tuberculosis).9 Thirteen companies responded, eleven of which completed the questionnaire. Of the two others, one indicated no reportable research activities in infectious disease and the other said time constraints prevented completion of the survey. The eleven companies who responded fully include at least six of the top ten. Together the respondents represent nearly US$117 billion of the global pharmaceutical market, which is estimated at $406 billion for 2002.10

What kinds of needs does the pharmaceutical market cover?

A represents Global Diseases, such as cancer, cardiovascular diseases, mental illness and neurological disorders, which constitute the major focus of the R&D-based pharmaceutical industry. Although affecting developed and developing countries, most people in developing countries who have needs for drugs to treat these diseases cannot afford them, and are thus not covered by the pharmaceutical market.

B represents Neglected Diseases, such as malaria and tuberculosis (TB), for which the R&D-based pharmaceutical industry has only marginal interest. Although also affecting people in wealthy countries, for example TB patients or people who get malaria while travelling, these illnesses primarily affect people in developing countries.

C represents the Most Neglected Diseases, such as sleeping sickness, Chagas disease and leishmaniasis, which exclusively affect people in developing countries. Because most of these patients are too poor to pay for any kind of treatment, they represent virtually no market and for the most part fall outside the scope of the drug industry’s R&D efforts, and thus outside the pharmaceutical market.

Z represents the part of the pharmaceutical market for products addressing conditions other than those which are purely medical (such as cellulite, baldness, wrinkles, dieting, stress and jet-lag), which nonetheless represent a highly profitable market segment in wealthy countries.

What are neglected diseases?

A seriously disabling or life-threatening disease can be considered neglected when treatment options are inadequate or don’t exist, and when their drug-market potential is insufficient to readily attract a private sector response. Government response is also inadequate. In short, for neglected diseases, there has been a failure of the market and a failure of public policy. Neglected diseases mainly affect people in developing countries. Public research institutes in the industrialized world do not view these diseases as either a priority or a major threat to their populations, and research-based drug companies do not pursue promising compounds for drugs to treat these illnesses because of an inadequate return on investment.

A look at the dynamics of this market failure shows that a distinction between “neglected” and “most neglected” diseases can also be made. For the “most neglected” diseases, patients are so poor that they have virtually no purchasing power, and no amount of tinkering with market forces is likely to stimulate interest among drug companies. If the market is failing poor people suffering from neglected diseases, it has failed people suffering from the most neglected diseases even more (Figure 1B). Some examples of neglected diseases are malaria, tuberculosis, human African trypanosomiasis (sleeping sickness), South American trypanosomiasis (Chagas disease), Buruli ulcer, dengue fever, leishmaniasis, leprosy, lymphatic filariasis and schistosomiasis. All but the first two can be considered most neglected diseases.

Tropical diseases are good examples of neglected diseases. Of the 1,393 total new drugs approved between 1975 and 1999, only 1% (13 drugs) were specifically indicated for a tropical disease.8

An empty pipeline

An examination of current research efforts in the pharmaceutical industry reveals that the pipeline of drugs for neglected diseases is virtually empty (see Figure 1C). In spring 2001, the DND Working Group and the Harvard School of Public Health sent written questionnaires to the world’s top 20 pharmaceutical companies to assess the level of R&D activity in several neglected diseases (sleeping sickness, leishmaniasis, Chagas disease, malaria and tuberculosis).9

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10) Six respondents were among the top ten companies worldwide, by sales; two other respondents chose to remain anonymous.
### Number of companies (out of 11 respondents) with research and development activities targeting drugs for neglected diseases

<table>
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<tr>
<th>Disease</th>
<th>R&amp;D Spending</th>
<th>Screening</th>
<th>Pre-clinical or Clinical Development</th>
<th>Product to market in last five years</th>
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<td>0</td>
<td>0</td>
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</tr>
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<td>Chagas disease</td>
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<td>0</td>
<td>1</td>
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<tr>
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<td>1</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>tuberculosis</td>
<td>5</td>
<td>4</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>Other infectious diseases</td>
<td>9</td>
<td>N/A</td>
<td>8</td>
<td>6</td>
</tr>
</tbody>
</table>

**Methodology:** The survey was sent to the CEOs and/or Directors of Research of 20 pharmaceutical companies in Europe, Japan, and the US. The questionnaire inquired about overall resources devoted to infectious diseases, and specific resources devoted to particular neglected diseases. The survey stated that individual company names would not be disclosed when reporting the results. Results relied on self-reporting and reports were not independently validated.
Teno

Teno Worku is in Kahsay Abera hospital in Humera, Ethiopia. He doesn’t get any visitors. His only family, his mother, lives 300 km further south in Gondar. “I’m a commercial traveller and I pass through this region a lot. Five months ago I took ill. I had a headache and a fever, so I went back to Gondar to see a doctor. He treated me for malaria. But a month later I still hadn’t recovered.” Frail and emaciated, the 28-year-old looks at least ten years older than his age.

Eventually, Teno went to the Ethiopian capital of Addis Ababa for tests. Four months later, not a single doctor had been able to explain his symptoms. Bitterly disappointed and by now critically ill, he returned to his mother in Gondar, where, at long last, a doctor in a private clinic suspected kala azar and advised him to go to Humera.

“The doctor said that this hospital is specialized in kala azar. Tests showed that I did have the disease, and the injections were begun right away. I was very late starting the treatment but I’m getting a bit better every day,” he says, sounding as if he is trying to convince himself more than anyone else.

Teno is about to receive his twentieth injection. He grits his teeth and braces himself for a painful experience. The needle has to penetrate deep into the upper buttock in order to inject the fluid into the muscle tissue.
Bianga

Bianga had been ill for ten months. She had become too weak to work in the fields near her home in Omugo, Uganda, fetch water, or care for her six-year-old son, Lino.

At first, Bianga found herself sleeping all day long but lying wide awake at night. Her behavior changed: she would run out into the street, shouting loudly at the sky. At this point, her husband left her. Bianga and her son went to live with Bianga's elderly mother in her small hut. With no one in the family earning money and unable to produce their own food, they were penniless. Lino became malnourished.

Finally, in despair, Bianga's mother took her to the hospital to see if something could be done. The doctor discovered that she was suffering from sleeping sickness and had already reached the stage at which the parasite invades the brain. She was admitted directly to the treatment center where she was given a course of melarsoprol. Although the treatment was painful, she began to feel better. After the 20-day course, she was able to return home and pick up her life again.

After a month, Bianga began behaving strangely again, and Lino brought her to the hospital. It was discovered that she had gone into relapse. Her ankle had to be tied to the bed to prevent her from running away and getting lost. Bianga received another course of melarsoprol, but this time her condition did not show much improvement. With no other treatment available and little hope for recovery, she was sent home. For Bianga, some of the treatments that are only now becoming available arrived too late.
Dropped from the private research agenda

Over the last few decades, major progress in molecular biology and biotechnology has enabled the development of increasingly sophisticated medicines to cure a wide variety of diseases. Moreover, global expenditure on health R&D has increased dramatically and is still on the rise. For 2001, an estimated record US$70 billion will be invested globally in health R&D, with the U.S. private sector alone accounting for just under half of the spending at US$30.5 billion.1 While the public sector has traditionally been the major funder of health research, the private sector has recently taken the lead. Global health research priorities are changing accordingly.

In the "social contract" that has over the years emerged around drug development, industrialized countries rely on the pharmaceutical industry to develop and produce medicines, and governments attempt to ensure that the industry meets public needs through a variety of incentives. These incentives include the patent system, tax credits, and R&D grants, as well as subsidies provided by national health care or insurance systems to help pay for health commodities.

This balance between public and private capacity, investments, and interests has worked well to develop drugs for diseases such as heart disease and cancer, and has helped the pharmaceutical industry to prosper, with companies often having sales of hundreds of millions or even billions of dollars a year on a single drug. This profit-driven system has also mobilized R&D funds for "lifestyle" conditions such as impotence, baldness and obesity. By investing in these conditions or in "me-too" drugs (medicines that are only slightly different from existing compounds and are not considered to be true innovations or clinical advances), drug companies can also expect phenomenal sales figures.

According to the 2000 Fortune 500 ranking, pharmaceutical companies top the US industry performance list for return on investment, with a 39% return for shareholders.2 Furthermore, corporate mergers and consolidations have led to fierce competition between a shrinking number of players. To maintain expected profit levels, the R&D-based pharmaceutical industry focuses on the profit potential of wealthy markets. Figure 2A projections show that North America, Europe and Japan will account for 80% of the world pharmaceutical market in 2002 (with a total projected world value of $406 billion), while Africa, Asia, Latin America and the Middle East, representing 80% of the world's population, will account for only 20% of the pharmaceutical market.3


Lida

Lida weighs 35 kilograms. She says she feels “destroyed inside.” In her room in the department of “chronics” at the tuberculosis hospital in Guliripchi, Abkhazia, she waits for the results of an analysis of her sputum. She hopes for the impossible: the destruction of all the bacteria eating away at her lungs in spite of four successive treatments.

The tuberculosis from which Lida suffers is multi-drug resistant, no doubt contracted as a result of two previous treatments that were incomplete.

The first treatment, prescribed by the doctor at the iron and steel plant where Lida worked, included only two of the anti-tuberculosis drugs recommended by the World Health Organization protocol. As for the second, following crisis and war in her country, she didn’t have enough money to pay for the medication.

The two treatments that followed in the hospital were appropriate but arrived too late. A chronic patient, she would now need second-line medication, but that’s not available in Guliripchi. The complete treatment costs $15,000 and cures only up to 60-70% of the patients. There are terrible side effects, and hospitalization and treatment last up to 24 months.

During the year and a half that Lida has been in the hospital, she has put on a little bit of weight. She moves slowly from her bed to the window to breathe.

**MYTH:**

**The typical new drug, brought successfully to market, costs approximately US$500 million for research and development.**

This often quoted figure is based on a paper written by J.A. DiMasi and published in 1991. The DiMasi paper put the cost of developing a new drug at US$231 million. Subsequent studies used a higher opportunity cost of capital and changed other parameters, and the figure became $312 – $359 million. Adjusted to 2000 dollars the amount turns into $473 million. And this figure is quite simply rounded up to arrive at $500 million.

Yet the original study has several limitations, and the subsequent estimates based on the study inherit these flaws.

The initial calculation was based on several assumptions that can be disputed. Assumptions were made for the cost of pre-clinical studies. Assumptions were also made about the length of the R&D process, the opportunity cost of capital (in other words, potential revenue if the capital were invested elsewhere) and success rates. Additionally, the study puts the opportunity cost of capital (not actual spending) at half of total R&D costs, but it doesn’t take into account tax deductions or government grants awarded to a company for R&D expenses.

Aside from relying on assumptions, the initial study wasn’t representative of the ‘average’ drug, nor was it designed to be. The original study focused on drugs that were researched and developed exclusively by multinational pharmaceutical companies. Yet development of many drugs depends on major public involvement in both basic research and clinical trials. Calculating an average cost for R&D has limited usefulness, in any case, because costs may differ greatly between drugs for chronic diseases and drugs for acute infections, or between innovative drugs and me-too drugs.

Recent independent estimates on drug development costs vary. The group Public Citizen (using DiMasi’s original study as a base) computes the cash outlay for new drugs at $110 million, excluding opportunity cost but taking into account inflation and tax deduction; the Global Alliance for TB Drug Development (GATB) puts the cost of a new tuberculosis drug at around $40 million (excluding the cost of failure) using a chemical entity already identified. When the cost of failure is included, GATB estimates the cost at $76 – $115 million.

One final drawback to the original DiMasi study, and a limitation for these subsequent efforts to put a price tag on drug development, is the data: it came from confidential industry sources in the 1980s and has not been available to other researchers. To gain a better sense of what it would cost to develop a drug, access to actual data is essential.

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It is a matter of simple economics: potential return on investment, not global health needs, determines how companies decide to allocate R&D funds. According to the drug industry, the low purchasing power of developing countries – coupled with the high cost of R&D and drug registration – rationalizes their focus on wealthy country markets. Fierce market competition means that for diseases that primarily affect developing countries, neither promising drug leads nor research on new applications of existing drugs will be pursued.

Gaps in the drug development process

A closer examination of the drug development process shows exactly where the system breaks down. Developing a new drug from basic research can be a complex, capital-intensive and time-consuming activity. In order to produce one successful drug, thousands of candidate-compounds and successive selections based on biochemical properties, safety, clinical performance, and market considerations may be needed. Figure 2B outlines this process and identifies the gaps that occur when market prospects are low.

It is clear that the multinational pharmaceutical industry cannot be relied on to develop the medicines required to treat the diseases that affect the world’s poor. Governments are ultimately responsible for ensuring that people’s health needs are met. They must take action when the private sector or the market fails. The current crisis in R&D for neglected diseases is a result not only of the failure of the market, but also of public policy.
A Matter of public responsibility

Inadequate public policy has compounded the failure of the market to generate R&D for drugs for neglected diseases. Governments have the power to influence drug development, both through direct research funding and policies to influence the activities of the private sector. Not only can governments make a difference, they have a responsibility to do so. They should increase both their funding of and direct involvement in drug development for neglected diseases. But for the past 20 years, despite clear evidence of the decline in private sector interest in neglected diseases, government leaders have often stood by silently.

Government inaction compounds crisis

A needs-based approach and consolidated public funding of R&D for neglected disease drugs could have compensated for the market failure. Instead, public sector research has increasingly focused on diseases that affect wealthy countries. There is increasing pressure for publicly funded research to have commercial applications, further reinforcing the focus on lucrative diseases. Governments fund public research according to the health needs of their own constituencies. The end of colonial presence and declining military involvement in tropical countries has led to a further waning of interest in tropical diseases in the latter half of the 20th century.

Leaders in disease-endemic countries have also done little to improve the R&D situation for neglected diseases. In 1990, the Commission on Health Research for Development proposed that all governments allocate 2% of health expenditure to research. According to the Global Forum for Health Research and its partners, none of the low- and middle-income countries studied were making this level of contribution in 1998. Basic research that leads to the discovery of potential “drug leads” has almost always been publicly funded at universities, in-house government facilities, or research institutes in Europe, North America, and Japan. Since the beginning of the 20th century, publicly funded research has led to major drug lead discoveries in, for example, tuberculosis (streptomycin and rifampicin), other infectious diseases (various antibiotics), and cancer (various types of chemotherapy). More recently, publicly funded research has led to the discovery of antiretrovirals for the treatment of HIV/AIDS. Publicly funded genome research has also produced many drug leads.

Meanwhile, public sector policies increasingly view public research as an investment that needs to create economic value. Scientists are requested to not only publish their research and advance science, but also promote and actively pursue the possible commercialization of research findings (through active patenting and licensing strategies, research collaborations with industry, creation of spin-off companies, etc.). This so-called valorization of research has become an important policy objective of public research, especially in the biotechnology and health sector where financial returns are very attractive. Thus, the same market failure that detered the pharmaceutical industry from investing in neglected diseases also discourages the public research community.

While supporting basic and drug-lead discovery research, the public sector has rarely developed its own drug development expertise and capacity. It is the pharmaceutical industry that leads product development, from pre-clinical research through regulatory approval. However, the most innovative part of the process is the initial identification of lead compounds, which often happens in the public or academic research sectors. In these settings, publishing innovative research in high-ranking journals often makes a career and ensures continued funding. Not surprisingly, the most important gap in the drug R&D process for neglected diseases is between basic research and pre-clinical research (see Figure 2B, page 18).

Estimates of public, nonprofit, and foundation spending

Because there is a lack of basic, up-to-date information concerning spending on neglected disease R&D, the DND Working Group asked leading international experts on several neglected diseases to estimate the current level of investment in R&D:

**Leishmaniasis**

Dr. Farrokh Modabber, Director of the Infectious Disease Research Institute, Seattle, USA, estimates current research spending for leishmaniasis at US$20 million. Of the total, 15-20% is directly spent on drug development.9

**Malaria**

Dr. Catherine Davies of Wellcome Trust estimates that funds committed by major funders to malaria research in 1999 were over US$150 million (excluding US Department of Defence and French sources, for which detailed figures were not available). The equivalent figure for 2000 is over $200 million.10

Dr. Rob Ridley of the Medicines for Malaria Venture (MMV) says that, depending on how it is defined, drug discovery and development might constitute between 10-20% of the overall malaria research spending figure for 2000.11

**Sleeping sickness**

Mr. Felix Kuzoe, an expert in African trypanosomiasis (sleeping sickness) at the Special Programme for Research and Training in Tropical Diseases (TDR) estimates total research spending at as little as US$20 million in 2000. Of this total approximately $4 million (20% of the total) is devoted to drug development thanks mainly to a donation from the Gates Foundation. In 2001, the total research spending will increase to $21 million due to a recent donation from Aventis Pharma. This will increase the proportion for drug development to 24% (about $5 million) in 2001.12

**Tuberculosis**

Dr. Paul Nunn at TDR estimates research spending on tuberculosis by governments and private foundations during 2000 at US$143 million. Of this figure, only $37 million (27%) is devoted to drug development.13

Recent spending patterns

- The drug development outcomes for a particular disease clearly reflect the money invested in R&D. To gain an indication of the amounts currently being spent specifically on drug R&D for neglected diseases, the DND Working Group spoke with recognized experts on tuberculosis, malaria, sleeping sickness, and leishmaniasis. Based on their estimations (see above), government, nonprofit, and foundation funding for drug R&D appears to be little more than US$100 million per year for these four diseases combined. Putting this figure into perspective, total public spending on health research worldwide is estimated at $30 billion, of which $3.1 billion is devoted to cancer research in the US alone.5

TDR

- Another symptom of government indifference to the R&D crisis is the plight of the Special Programme for Research and Training in Tropical Diseases (TDR), the main international public body charged with research into tropical diseases. Established in 1975 as a joint program of the United Nations Development Program, the World Bank and the World Health Organization, TDR was intended to be a public sector response to pleas from countries where neglected diseases were endemic.

- TDR has two objectives. The first is to conduct research into new medicines to help control a defined group of tropical diseases.4 The second is to train scientists and strengthen institutions from disease-endemic countries and encourage them to play a larger role in the research process. TDR has achieved some considerable successes.7 Six of the thirteen drugs developed for tropical diseases between 1975 and 1999 were developed with TDR support, and the program has also raised awareness of tropical diseases and helped set the agenda for research. Even so, it has remained chronically under-funded. For many years the program has struggled by on about $30 million per year to fulfill a mandate for both research and training activities in the ten diseases it covers. Furthermore, TDR works within the UN system, abiding by international civil service norms, and the program is pulled by the differing priorities of its multiple sponsoring agencies. This is not an ideal management structure in a field where decisions on research and allocation of resources must be made quickly.

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6) Diseases currently in the TDR portfolio are leishmaniasis, onchocerciasis, schistosomiasis, lymphatic filariasis, Chagas disease, malaria, leprosy, African trypanosomiasis, tuberculosis and dengue.
The Rockefeller Foundation has also played a crucial role in neglected disease drug development. In the last few years, the Gates Foundation has given $25 million (over five years) to the Malaria Venture, $25 million (over five years) to the Global Alliance for TB Drug Development (GATB), and another $15 million to vaccine research for leishmaniasis. Previously, leishmaniasis vaccine trials had relied on small TDR resources plus in-kind donations from affected countries. The Gates donation has “completely changed the picture,” according to Dr. Farrokh Modabber, Director of the Infectious Disease Research Institute in Seattle (see page 21). The foundation also gave $15 million (over five years) for sleeping sickness and leishmaniasis drug development. Prior to this donation, the main funding for sleeping sickness drug development was less than $500,000 a year through TDR. The award of $15 million “has no precedence in the history of African trypanosomiasis,” according to Felix Kuzoe, an expert on sleeping sickness at TDR (see page 21). The Gates Foundation has also funded various research-related activities around several other neglected diseases.

The Rockefeller Foundation has also played a crucial role in raising awareness of global health issues and, in 2000, awarded $15 million to the public-private R&D initiative, the Global Alliance for Tuberculosis Drug Development (GATB).

The Bill and Melinda Gates Foundation, in addition to providing substantial funding for vaccines, has become a major force in neglected disease drug development. The foundation has given $25 million (over five years) to the Global Alliance for TB Drug Development (GATB), and another $15 million to vaccine research for leishmaniasis. The award of $15 million “has no precedence in the history of African trypanosomiasis,” according to Felix Kuzoe, an expert on sleeping sickness at TDR (see page 21). The Gates Foundation has also funded various research-related activities around several other neglected diseases.

Since the 1970s, some industries in developing countries have been developing new production processes through reverse engineering for medicines still under patent elsewhere in the world. This generic production has contributed to both industrial development and greater access to medicines through lower prices. With stronger patent protection, these countries will not be able to continue this practice.

**MYTH:**

There is little investment in tropical diseases because there is weak patent protection in the countries most affected by these illnesses. After 2006, when all countries will have implemented TRIPS (international trade rules that mandate minimum 20-year patents), drug development will increase in developing countries.

Drug development for neglected diseases will not automatically increase, no matter how strong the level of intellectual property protection, because private R&D is driven primarily by market potential. People who suffer from diseases like malaria, sleeping sickness and leishmaniasis, with or without strong patent protection in their countries, will not have the necessary purchasing power to constitute a market attractive to drug developers.

Intellectual property rights, including patents, are part of a complex legal and economic system that can motivate investment in R&D under certain circumstances. Protection of intellectual property in a country has historically followed industrial development. It is doubtful that the reverse will also occur – that industrial development will follow strong intellectual property protection. In fact, patents may actually hamper medical research activities in developing countries. Patents are often owned by private companies or research institutions, and, during the period of protection, put limits on research knowledge. Molecules that could be promising for the treatment of neglected diseases are consequently not easily accessible for research.

In addition, most developing countries are unlikely to significantly improve their R&D capacity solely on the basis of an expanded and stronger intellectual property rights regime. Even in industrialized countries, innovation is assisted by other incentives, including substantial government spending. Without significant government research spending, stronger patent protection may lead to higher prices without stimulating research.

Since the 1970s, some industries in developing countries have been developing new production processes through reverse engineering for medicines still under patent elsewhere in the world. This generic production has contributed to both industrial development and greater access to medicines through lower prices. With stronger patent protection, these countries will not be able to continue this practice.

21) UNCTAD. The TRIPS Agreement and Developing Countries. Geneva, 1996.
Traditionally, governments have played a positive role in developing drugs for communicable diseases. For example, with very few exceptions, today’s malaria drugs were initially discovered outside the private sector in universities or government labs – institutions known for their competence in identifying promising prospective drugs. The Walter Reed Army Institute of Research, for instance, with a small budget from the US Department of Defense, invented four important antimalarial drugs, which were then developed in collaboration with multinational drug companies.2

Despite the efforts of individual actors in the public sector, R&D into neglected diseases remains woefully inadequate. While the public sector is not powerless, it currently depends largely on the skills and expertise of the private sector to conduct final drug development. If the private sector is unwilling to take a drug through this final stage of development, it never leaves the laboratory. Recent proposals have sought to increase public sector involvement while increasing incentives for the private sector to move compounds beyond the laboratory and ultimately deliver them as drugs to patients.

In order to attract private sector R&D capacity back into needed areas, what are called “push” and “pull” mechanisms have begun to emerge as possible answers. “Push” mechanisms reduce costs and risks of R&D and can include tax credits, R&D grants, and support for clinical trials. “Pull” measures help create a market for drugs or increase their profitability. Two examples are the creation of purchase funds and “patent exchange,” whereby a company would invest in developing a drug for a neglected disease and

The International Conference on Harmonization: Is the bar being raised too high?

Getting a drug to market requires a complex series of evaluations and regulatory reviews to ensure that it meets quality, safety and efficacy standards. Approval of new drugs is done by national governments, which set standards. The United States, Japan, and the European Union have attempted to harmonize their standards through the creation of the International Conference on Harmonization (ICH), an initiative of drug regulatory authorities and research-based pharmaceutical industries. The goal of harmonization is to reduce drug development and regulatory review times.

The ICH is tightening requirements beyond those stipulated by the World Health Organization (WHO). The quality, efficacy and safety requirements that constitute the ICH guidelines deal specifically with drug development in a wealthy market, where cost is not a major issue and where safety is defined as near-zero risk. For neglected diseases, cost is a major issue, and the risk-to-benefit ratio in terms of quality, efficacy and safety should be put into the perspective of the gross public health failure of having no treatment at all.

These more stringent ICH guidelines raise costs and present barriers to drug development, particularly for small or medium-sized companies in developing countries. The risk is that the bar will be raised so high that only drugs developed in the industrialized world will be able to be marketed internationally. This will seriously hamper the development of R&D capacity in developing countries, which has been identified as a necessary component in the long-term solution to the R&D crisis. This potentially negative public health implication needs to be carefully weighed against the possible benefits of raising R&D requirements, which some have argued would be marginal and of little value to patients.

Many questions remain unanswered on the implications of the ICH guidelines. An independent and thorough technical review of ICH guidelines should be undertaken by WHO. If ICH is to become a global standard, it must be reexamined to ensure that it meets the needs of both developing and developed countries.

then, once the drug was approved, would have the right to extend the patent on one of its other, more profitable drugs. Both “push” and “pull” mechanisms are market-based measures that aim to increase the investment return for a drug to a level that will attract the private sector.

Orphan drug laws

Orphan drug laws are an example of a “push” mechanism. Orphan drug laws use tax credits and grants to promote research into drugs for diseases that affect a relatively small number of people (in the US this is set at 200,000 or fewer people).3 These rare diseases would otherwise represent a market return inadequate to motivate drug investment.

The Orphan Drug Act in the US (similar laws exist in Europe, Japan, Singapore and Australia) has successfully provided incentives for research into diseases such as cystic fibrosis.4 Some policymakers are recommending amending these kinds of laws to include neglected diseases in developing countries. However, it is critical to note that orphan drug legislation has succeeded because, in addition to tax incentives and government grants, companies can recoup costs by charging high prices for the drugs. One extreme example is the drug Ceredase, used to treat Gaucher’s disease, which was priced at hundreds of thousands of dollars per year of treatment.5 Since purchasing power is limited or non-existent among people with neglected diseases, the orphan drug mechanism alone is not likely to work. However, the concept may be useful if paired with other mechanisms, or modified to fit neglected diseases more specifically.

The history of this type of legislation also shows that it could be particularly effective in motivating small and medium-sized enterprises; in the US, over 50% of companies applying for orphan drug status are small and medium-sized.6 However, many of these companies depend on outside financing to support their R&D programs, and also need to maximize profits for their shareholders.

5) J ames Love, affidavit at the High Court of South Africa in the matter between Pharmaceutical Manufacturers’ Association of South Africa and Others and The President of South Africa and Others, and Treatment Action Campaign (Amicus Curiae), Case: 4183/98, 9 April 2001 (South Africa, 2001).
What about a “Pot of Gold”?

One commonly suggested “pull” strategy is the creation in advance of purchase funds for drugs for neglected diseases. The idea is to secure purchase funds through donors - the “pot of gold” waiting at the end of the drug development rainbow - in order to supplement an existing market, and thus “pull” companies into drug development. However, to prompt a major pharmaceutical company to invest, the existing market plus the “pot of gold” would need to compete with the average return on commercial sales, put at about US$265 million annually in 1998. This would be a great expense, and exist - are still likely to be overlooked. Still, the concept may be useful if paired with other mechanisms or modified to fit the most neglected diseases specifically.

Building capacity in developing countries

Building capacity in developing countries is another important strategy for stimulating R&D. Public health institutes in some developing countries are playing an increasingly important role in drug development. For example, the Thai government’s support

in a sense is “buying into” the existing drug development system by subsidizing shareholders’ needs for profits and other costs associated with private industry drug development.

This strategy could potentially work for some neglected diseases that affect large numbers of people, such as TB or malaria, because an existing market in wealthy countries would supplement the pot of gold (eg, TB in Europe or the malaria traveller market). For the most neglected diseases, a purchase fund by itself would likely be too costly for governments and other funders. Drugs for the most neglected diseases - again, those for whom a potential market does not

for malaria research has led to the development of an effective modern pharmaceutical version of artemisinin, a traditional Chinese medicine. In clinical trials, drugs using Thai artemisinin cured 90% of malaria cases, and elsewhere cut infection among children by 90% in camps for displaced people on the Thai/Burmese border. However, while this new formulation is saving lives in Thailand, it is not recognized as a legitimate treatment by international regulatory agencies because the research reporting methods used in Thailand do not match international agencies’ reporting requirements. In this case, “harmonization” regulations on drug R&D, which

If we introduce new medicines into poor countries, we will accelerate the development of resistance. We don’t necessarily need new drugs but we need to better use the ones we have.

Drug resistance is often perceived as a problem restricted to a few diseases in poor countries. It is, however, an inescapable phenomenon in both the industrialized and developing world, due to the normal genetic survival mechanism of most parasites, bacteria, and viruses. Resistance to drugs will inevitably develop, and can do so despite good drug management and high compliance to treatment.

For example, in the Moyo district of Uganda, sleeping sickness patients have been treated with the 50-year old drug melarsoprol for more than ten years. In spite of strict drug management and good compliance, recent studies have shown resistance in excess of 30%. In this case, although introducing combinations of drugs may forestall resistance, new drugs will also be needed.

In general, two things are required in the fight against drug resistance. Existing therapies must be used rationally in order to delay the onset of resistance, and new drugs must continuously be developed to create future therapeutic choices to face the inevitability of drug resistance. As with sleeping sickness, the neglect of tuberculosis and malaria drug research in the last thirty years has made treatment increasingly difficult and led to a situation where, in some instances, treatment is becoming less effective.

Finally, fear of inducing resistance has never been a sufficient reason to withhold necessary treatment in the industrialized world. It should not be considered justifiable in the developing world.

8) This $265 million refers to the 1998 average revenue of new launched drugs as calculated by Dr. Steve Arlington. Dr. Steve Arlington, “Pharma 2005: The Challenges” (paper presented at the American Society for Clinical Pharmacology and Therapeutics meeting, Orlando, Florida, March 7, 2001).


were created to meet the needs of wealthy markets, are hampering access to new treatments created in developing countries (see box page 25).

Drug research, development and production is increasing in, among other countries, Brazil, India, South Korea, Thailand, Malaysia and Argentina, countries that had not been considered in the past to have innovative R&D capacity. Some initiatives to build capacity in developing countries involve stimulating collaboration between the public and private sectors in those countries. For example, the International AIDS Vaccine Initiative (IAVI) is working directly with university scientists, governments and companies in South Africa, Kenya, Uganda, India and China. The IAVI has in particular identified India as an ideal location for “fast-tracking” vaccine development, given the country’s thriving pharmaceutical industry, experience in clinical trials and government commitment to research.\(^\text{11}\)

Regional ventures also attempt to maximize developing country capacity through inter-country collaboration. The International Vaccine Institute in South Korea is a non-profit organization that was created to develop vaccines for diseases prevalent in developing countries. The Institute has pooled the skills and knowledge of scientists in various developing countries, and has been identified as a possible model for drug development and production.\(^\text{12}\)

Another type of policy initiative that is often discussed as a potential solution to the R&D crisis is the public-private partnership (PPP). PPPs attempt to foster R&D for neglected diseases by mobilizing expertise, capacity, and funding from both the public and private sectors. Typically, the PPP plays a coordinating and management role around a disease-specific R&D agenda, tries to take advantage of appropriate push and pull mechanisms, and seeks a combination of public funding, philanthropic donations and in-kind donations from industry. Major examples of this kind of approach are the Medicines for Malaria Venture (MMV), the Global Alliance for TB Drug Development (GATB), and International AIDS Vaccine Initiative (IAVI). So far, no public-private partnerships have been designed specifically for developing drugs for the most neglected diseases.

Current government initiatives, “push” and “pull” mechanisms, building capacity for R&D in developing countries, and public-private partnerships are all only partial solutions to the continuing R&D crisis for neglected diseases. Many are new initiatives whose effectiveness will need to be evaluated over time. And all depend to a greater or lesser extent on market forces. None of them provides an adequate strategy for developing drugs for the most neglected diseases.

Pau

Pau no longer has a fever. Just one week ago, shakes, hot flushes, headaches and nausea began to overwhelm the frail body of this 14-year-old. Malaria. The third attack in three years. The small amount of chloroquine that she managed to find did not cure her: In Cambodia, malaria is now resistant to this medicine. The combination drug recommended by the health authorities is only available in health centers. The products sold on the private market are either fake or too expensive.

So Pau gathered the last bit of strength she had and walked for several hours to reach the health centre of Anlong Veng, the modest capital of this region in the northern Cambodia.

Like many settlers attracted by the lure of virgin land, Pau’s family lives in a poor hut on the side of the road that cuts through the forest.

Pau spends her day collecting bark off trees, which she sells to Thais. It seems that they make a kind of incense from it, intended to keep the mosquitoes away. At nightfall, when she has gone too far into the forest, she sleeps on the ground. It is at this hour that the mosquitoes attack.

Tomorrow Pau leaves the hospital. She will return to the forest – risking her life to earn a living.

Governments must lead in restarting R&D on diseases that are currently being ignored. They need to create and support new structures designed to develop essential medicines for diseases that are being sidelined by the private sector. The current model of profit-driven R&D should not be an exclusive model. Developing drugs as public goods should also be pursued.

6. Increased and reliable long-term funding for R&D into neglected diseases is urgently needed. The DND Working Group is exploring sustainable options to support R&D for neglected diseases through legal obligations. Governments can and do mandate industry spending in a wide range of areas. One example of a potential mandate would be an "essential research obligation" that would require companies to reinvest a percentage of pharmaceutical sales into R&D for neglected diseases, either directly or through public R&D programs.

A global treaty on R&D for neglected diseases could provide a framework for such mandates. Such a treaty should correct the imbalance that exists between private sector rights and obligations under present international treaties and agreements (eg, the World Trade Organization’s Agreement on Trade Related Aspects of Intellectual Property), and provide new legal options to make drugs for neglected diseases global public goods.

7. A complete cost analysis of the true costs of drug R&D should be carried out. Existing estimates on the costs of drug R&D vary widely and remain highly controversial. In order to address the R&D imbalance effectively and make informed funding decisions, policymakers need objective, accurate figures on the true costs of developing drugs. Calculating drug development costs within a commercial context, which will include items such as opportunity costs, will be dramatically different from calculating the funding needed to develop a drug in a non-commercial setting.

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**Recommendations for moving forward**

Since its formation in 1999, the DND Working Group has been studying the extent and causes of the R&D crisis in neglected diseases and analyzing potential solutions. This research has led to the following insights and recommendations:

1. Because drug development is done almost exclusively within the context of the proprietary pharmaceutical industry, investment in R&D is guided by market considerations. Therefore R&D for diseases that mainly affect the poor is stifled.

2. Public policy has failed to correct this failure, with the result that some diseases are being neglected.

3. The dynamics of the neglected are different depending on the number of people affected and their purchasing power. Therefore it is impossible to develop a single strategy to stimulate R&D. It is crucial to acknowledge the different dynamics of neglected and most neglected diseases: each category will need distinct strategies.

4. A well-defined and needs-driven R&D agenda is required to assist policy makers, funding agencies and the research community in setting priorities for developing safe, effective and affordable medicines. The World Health Organization (WHO), as the only legally mandated international governmental agency responsible for global health, should work toward establishing an essential R&D agenda. WHO should lead this process. The DND Working Group, with input from WHO, has begun by drafting agendas that prioritize R&D needs for leishmaniasis, sleeping sickness, and malaria. These documents analyze the disease burden, current research strategies, and existing and potential treatments for each of these diseases. A critical next step is for governments and international organizations to examine carefully how they can contribute to dislodging the bottlenecks that currently restrict development of new treatments.

5. Governments in both developed and developing countries need to take comprehensive action to compensate for the market failure in drug development for neglected and most neglected diseases.
8. Public funds for R&D into neglected diseases should be tied to guarantees of equitable access and affordability of the end product. Equitable access to medicines in developing countries should be a basic principle that guides policy initiatives from the start. If public funds are to be invested in correcting market failures in drug development, there must be guarantees that the new medicines developed are affordable to those who need them.

9. Focused capacity-building and technology transfer projects in developing countries should be encouraged as a direct way to increase R&D expertise and infrastructure. Long-term solutions to the current crisis in drug development for neglected diseases ultimately rest within developing countries. Therefore, the DND Working Group is cataloguing and examining means of increasing existing drug development capacity in developing countries, and is also working to promote technology transfer that will support sustainable drug development and production facilities.

10. An independent and thorough evaluation is needed of the current and future impact of the ongoing regulatory harmonization efforts (ICH process) on the ability of developing countries to increase their drug development efforts.

11. A new type of body is needed to contribute to drug development for the most neglected diseases. The DND Working Group is exploring the feasibility of a Not-for-Profit Initiative (DND NfPI) that would focus on drug development projects for neglected diseases. The DND Working Group’s analysis has concluded that current approaches to address the lack of R&D for neglected diseases do not sufficiently address the most neglected diseases. To ensure a sustainable solution, a new approach is needed that would systemically harness funding, new science and technology, and foster public-private cooperation for these diseases.

Encompassing the recommendations listed above, and based on the research of the DND Working Group, the vision of the proposed DND NfPI includes the following:

- Ensuring equitable access to effective, field-relevant and easy-to-use drugs for neglected diseases.
- Prioritizing the most neglected diseases, such as sleeping sickness, Chagas disease, and leishmaniasis.
- Using sound science and management techniques to pursue a vision of developing new drugs for neglected diseases.
- Collaborating closely with TDR, industry, and research institutes in developing and developed countries.
- Securing support of public and private resources over the long term, with the majority of funding coming from the public sector.
- Working with drug development experts in developing countries to build national capacity for future drug development.

It is hoped that the public sector will take a strong leadership role in the NfPI to establish its legitimacy and accountability to the public and provide it with the necessary funds.

Conclusion

Despite impressive advances in science and medicine, society has failed to allocate sufficient resources to battle the diseases that particularly affect people in poor countries. The vacuum in R&D for neglected and most neglected diseases means that doctors and nurses in developing countries still do not have effective medicines for many of the diseases they see every day. However, encouraging initiatives have emerged to counter the market and public policy failures that have led to this crisis. Many of these initiatives are new, and their effectiveness will need to be evaluated. For the most neglected diseases, implementation of new solutions, such as a not-for-profit initiative for developing drugs for neglected diseases, will be essential.