



# **MALAWI**

## The Pharmaceutical Sector Country Profile Survey

#### 1. Background and Rationale:

Pharmaceutical Sector Country Profiles aim to increase the availability of quality information on structures, processes and outcomes of health and pharmaceutical sectors of countries. This information will be collected through a questionnaire and is meant to be used by country decision-makers, health and pharmaceutical experts, international partners and the public through databases and published country, regional and global reports.

The information is categorized in nine sections, namely: (1) Health and Demographic data, (2) Health Services, (3) Medicines Policies, (4) Medicines Trade and Production, (5) Medicines Regulation, (6) Medicines Financing, (7) Pharmaceutical Procurement and distribution, (8) Selection and Rational Use and (9) Household data/access.

Every four years since 1999, health officials from the 193 WHO Member States have been invited to complete a standardized questionnaire (named Level I) reporting on the status of the national pharmaceutical situation. Level I indicators assessed structures and processes related to the pharmaceutical situation of a country. They were used to carry out a rapid assessment that would highlight strengths and weaknesses of countries pharmaceutical situations. 156 countries responded to the 2007 level I survey and the results were stored and available in a global WHO database and used to develop a global report as well as a number of regional and sub-regional reports. The Pharmaceutical Sector Country Profile questionnaire described here will replace the Level I tool for the 2011 Member States' survey. The aim of this new approach is to build on the achievements and lessons learnt from the Level I tools and surveys and to improve the quality and scope of information (e.g., outcomes and results indicators) and enhance the involvement and ownership of countries in the development of profiles. The new tool has been piloted in the 15 countries of the Southern African Development Community in 2009 and in 13 countries across the world in 2010. The of results these pilots available on-line at: are http://www.who.int/medicines/areas/coordination/coordination assessment/en/index.html

Another innovation of the 2011 survey is the collaboration between WHO and The Global Fund. In 2009, the Global Fund developed and introduced the Pharmaceutical and Health Product Management ("PHPM") Country Profile to gradually replace the Procurement and Supply Management ("PSM") Plan. In the course of 2010 both agencies have developed a joint Pharmaceutical Sector Country Profile questionnaire that includes key indicators of the

pharmaceutical sector and that will be used by both agencies as the sole tool for pharmaceutical sector data collection in countries. The information captured in the Pharmaceutical Sector Country Profile questionnaire will be used by the Global Fund during grant negotiations and signing, and will also support grant implementation. In addition to the Country Profile that provides an overview of countries' pharmaceutical sectors, the Global Fund will also use a second questionnaire that will focus in more detail on medicines procurement and supply.

### 2. What can Pharmaceutical Sector Country Profiles offer:

Completing this questionnaire will require the time of national experts and responsible officers but it is worthwhile as your country and your partners will benefit from it in a number of ways:

- The questionnaire offers a unique opportunity to consolidate, in one place, information that is available in different locations and institutions e.g. the National Medicines Regulatory Authority, Central Medical Stores, National Health Accounts, etc.
- II) The methodology proposed for filling in the questionnaire will ensure that good quality data are collected and that the source and date of information are known and reported.
- III) Data on structure, process and outcomes are collected, and the questionnaire has been pre-filled with data available in the public domain; indicators are divided into core and supplementary in order to make it easier to identify what is more important.
- IV) The data collected will highlight the strengths and weaknesses of the pharmaceutical sector and will be made available in a national database as official country information, for use by decision-makers, health and pharmaceutical experts, researchers and international partners and the public..
- V) The data collected could be transformed into a narrative report with robust data analysis and bibliographic references, that will summarize the medicines situation in the country.
- VI) Based on experiences from previous surveys, a detailed glossary of key definitions and a manual for use of the questionnaire have been developed and can be found at the end of the questionnaire.

### 3. The process of data collection and analysis:

**3.1 Data collection**. The Pharmaceutical Sector Country Profile questionnaire has already been filled in by WHO with reliable data available from global and country sources. We kindly ask you to review, to correct (if necessary) and to validate the information already included in the questionnaire, and also to fill in the gaps, based on reliable information available in your country.

In order to do this, we recommend that you involve the most appropriate respondents and responsible institutions to fill in the various components of the tool so that the questionnaire is completed within the given deadline, with good quality information. If during the data collection process, clarifications are needed, WHO Regional and Headquarters Offices will provide the necessary assistance and support, including for data quality issues.



**3.2 Official endorsement**. Once the questionnaire has been completed, the information contained in it should be officially endorsed and its disclosure authorized by a senior official in the Ministry of Health. This should be done by signing the formal endorsement form attached to the questionnaire. This will ensure that the quality of the information contained in the Pharmaceutical Sector Country Profile questionnaire is certified by the country.

**3.3 Data shared with the Global Fund**. Data collected from Global Fund priority countries will be shared with the Global Fund and it will be used as part of the Global Fund's own grant signing and implementation procedures.

**3.4 Data posted on key databases**. Data endorsed by the country will be posted on health databases (such as the WHO Global Health Observatory, <a href="http://www.who.int/gho/en/">http://www.who.int/gho/en/</a>), making it available to decision-makers, health and medicines experts and researchers, international partners and the public.

**3.5** Development of narrative Pharmaceutical Sector Country Profiles. Data provided within the country questionnaire can be used by the country to develop a narrative profile that will illustrate the national pharmaceutical sector. In order to do this, WHO has prepared a template profile (included in the CD-Rom shared with you) that can be easily used by countries and that will help presenting data in the form of tables, graphs and charts. Countries could seek support from WHO for the development of their narrative profile, which will be finalized and validated by the country that will own the copyright for it and will publish it as a national official document.

**3.6 Development of Regional and Global Reports.** The information provided by countries in the Pharmaceutical Sector Country Profile questionnaire will be analysed by WHO and used to produce regional and global reports on the pharmaceutical sector of countries in 2011. These reports will provide an overview of the progress made between 2007 and 2011, of the challenges that remain to be addressed and will include data analysis by technical areas, countries' income level and geographical location.

# Guidelines for countries on how to fill in the Pharmaceutical Sector Country Profile Questionnaire

### Please read these instructions carefully before starting data collection

<u>1. Macros:</u> the questionnaire has macros installed. A macro is a series of MS Word commands and instructions that are grouped together as a single command to accomplish a task automatically. For these macros to work properly, the macro security levels for MS Word on your computer should be set as 'low'. This can be easily adjusted by taking the following steps:

- 1. Open the Word document containing the instrument.
- 2. Go to 'Tools' > 'Macro' > 'Security'.
- 3. Click on the tab 'Security Level'.
- 4. Set the Security on 'Low' and click 'OK'.

After filling in the questionnaire, the setting should be restored to a higher level of security in order to protect your computer.

<u>2. Core and supplementary indicators:</u> the instrument consists of core and supplementary questions. Core questions cover the most important information, while supplementary questions deal with more specific information applicable to particular sections. Please note that core questions have been shaded with different coloured backgrounds for different sections of the instrument, while supplementary questions are all white. This should help you to distinguish between the different categories of indicators. Please try to fill in all the core questions for each section before moving to the supplementary ones. Remember that we are only asking you to collect information that is already available and you are not expected to conduct any additional survey(s).

<u>3. Prefilled data</u>: the answers to some of the questions have been prefilled by WHO HQ. Where this is the case, please verify this information as it may not be up-to-date. If you find that any of the prefilled responses are not correct, please change the value and document the source and year.

<u>4. Calculated fields:</u> for a few items, you will not be required to enter any value as these will be generated at WHO HQ using data entered into related fields. These fields have been clearly marked in red – please do not input any data into them or change data that are already in this field. For example, the per capita expenditure on health will be automatically calculated once the total health expenditure and population are entered into the questionnaire. This system is intended to improve the quality of answers and avoid you having to perform additional calculations. Calculated fields are protected and cannot be changed.

### 5. Possible answers:

*Checkbox 'Yes/No/Unknown':* tick one of the three options (only one answer is possible).

Multiple choice checkbox: tick any of the options that apply (multiple answers are sometimes possible).

*Percentage fields:* 0-100. Please use decimal points ('dots') for decimals (example: 98.11). Please do not use ranges (e.g. "3-5"). If you only have ranges, then use the median and otherwise the mean. In this instance, please detail what data you have used and what the range is in the comment boxes.

*Number fields:* unlimited number. Please use decimal points ('dots') for decimals (example: 29387.93). Please do not use ranges. If you only have ranges, then use the median and otherwise the mean. In this instance, please detail what data you have used and what the range is in the comment boxes.

<u>6. Comments:</u> comments fields allow the entry of free text to clarify or follow up on answers given. Please reference each comment by using the number of the question you are referring to (example: 2.01.02).

<u>7. Year of data</u>: year fields should be used to specify the year of the **data** used to answer the question. Only values between 1930 and 2011 will be accepted. Please use this column as follows:

- When the source refers directly to a specific document (for example: 'Medicines Act' or 'EML'), please put in the publication year of the document (note: only the year and not a specific date can be entered).
- When the source refers to a document that contains older data than the document itself, please put in the original year of the data. For example, when the total population for 2008 is extracted from the World Health Statistics 2010, please put 2008 in the 'year' column and 'World Health Statistics 2010' in the 'source' column.
- When the source of the information is not a document, but the informant himself/herself, please put in the current year.

8. Source of data: sources used for the answers given will be referenced in the narrative country profile and in the databases in which the information will be stored. Please specify your sources as clearly as possible by providing the name, year, and writer/publisher of the documents used. Also provide a web (URL) link to the documents, if available. If there is only a non-English version of the reference available, then please include it regardless of the language. Use the 'source' column to enter the name and year of the **source**, and use the "Comments and References" fields at the end of every section to list the sources. In case the source is not documented, then provide the name and title of the person and/or the entity they work for as a source of information. Examples are given below.

7.01.125	Which of the following <u>tender</u> methods are used in public sector procurement		1996 рон, 1996
7.01.12.01 S	National competitive tenders	Yes 🛛 No 🗋	
7.01.12.02 S	International competitive tenders	Yes 🛛 No 🗖	
7.01.12.03 S	Direct purchasing	Yes No	
7.01.135	Comments and References	National Drug Policy for South Africa , publis availabilt at: http://www.doh.gov.za/docs/poli	

<u>9. Documents:</u> you will see in the questionnaire that we would like you to collect and share a number of key country documents that we believe would greatly enrich the country's profile content and these documents could be made available through countries and WHO web pages. Please attach the following documents, if available:

- National Medicines Policy (NMP);
- NMP implementation plan;
- National Medicines Act;
- National pharmaceutical Human Resources report or strategic plan;
- Latest report on the national pharmaceutical market (any source);
- Pharmacovigilance national centre report (including an Adverse Drug Reaction (ADR), analysis report produced in the last two years);
- National pharmaceutical legislation or regulation;
- Annual report of quality control laboratories;

- Annual report of national regulatory authority;
- Legal provisions on medicines price regulations;
- Medicines procurement policy;
- National Essential Medicines List (EML);
- National Standard Treatment Guidelines (STGs);
- National strategy for antimicrobial resistance;
- Any other medicines pricing/availability surveys, household surveys and rational use surveys, in addition to the ones used to prefill the instrument.

The last page of the questionnaire contains a table with the list of key documents to be attached. Please fill it in by indicating the exact title, publisher and year for each attachment as shown in the example below.

Document	Exact title	Author	Publisher	Year	File
					name
Essential Medicines List	National	Ministry of	Ministry of	2009	EML.doc
	Medicines List	Health	Health		
National Medicines	National Drug	Federal Ministry	Federal Ministry	2005	NDP.pdf
Policy	Policy	of Health	of Health		

These documents will be published on the WHO web site's medicines library (<u>http://apps.who.int/medicinedocs/en/</u>) and will therefore have to be endorsed by the Ministry of Health prior to being made publicly available. You can send us these documents by e-mail as attachments or you can upload them into a protected web site. Please use the table at the end of the instrument to report the title, year and author of the documents attached.

<u>10. Attaching files to the questionnaire:</u> please place all files to be attached in a single folder on your computer. Name the documents as follows: <short name of the document>.doc (example: EML.doc). Then compress (ZIP) the files and attach the compressed file with the completed instrument to the email. If the total file size of the compressed file exceeds 7 MB, you can upload the documents in a protected file server called MedNet, which is managed by WHO. The procedure for doing this is very simple and please contact Mr Enrico Cinnella in WHO HQ, Geneva, (cinnellae@who.int) to be granted access to MedNet and to receive instructions on how to upload files. You can also upload documents to the WHO Medicines Documentation server at <a href="http://hinfo.humaninfo.ro/medicinedocs/">http://hinfo.humaninfo.ro/medicinedocs/</a>, though the documents will only appear on the Medicines Documentation site at the beginning of the following month.

<u>11. Manual for use of the questionnaire</u>: the manual contains detailed instructions on the questionnaire, on where to find information and how to answer questions.

Questions that may be particularly problematic are marked with the following icon:



12. <u>Glossary</u>: the glossary contains definitions for all key and/or problematic items in the instrument. It is highly recommended that you use the glossary, since exact definitions might differ between countries and institutions. The glossary is at the end of the file. When a question contains an item that is defined in the glossary, the terms will be marked in bold, underlined and written in blue font.

2.02 Healt	th Personnel and Infrastructure				
Core ques	tions <u>(click for help)</u>				
			Year	Source	
2.02.01	Total number of pharmacists licensed/registered to practice in your country				
2.02.02C	Pharmacists per 10,000 population	-			
2.02.03	Total number of pharmacists working in the public sector	-			definition of "pharmaceutical
2.02.04	Total number of <u>pharmaceutical</u> technicians and assistants				<ul> <li>technicians and assistants" is in the glossary</li> </ul>
2.02.05	A strategic plan for pharmaceutical human resource development is in place in your country?	Yes 🔲 No 🗌			Instructions are available for this specific question

<u>13. Respondents and acknowledgements</u>: at the beginning of every section there are fields available to fill in details about the respondent for that particular section. It is also possible to enter the details of multiple respondents. At the end of the instrument please add a list of contributors who should be acknowledged. Provide their names and the main organization(s) they work for.

<u>14. Endorsement of data</u>: A formal endorsement needs to be signed by a senior official in the Ministry of Health before the completed questionnaire is sent back to WHO. The endorsement form is included in the pack of CD-ROM documents you have received from WHO. Please present the endorsement form to a senior official in the Ministry of Health for signature, and for obtaining permission to use and publish the data.

<u>15. Process of creating a country profile document:</u> The data you will collect using this questionnaire can be used to develop a pharmaceutical sector country profile for the country. Examples of profiles are available on-line at <a href="http://www.who.int/medicines/areas/coordination/coordination\_assessment/en/index1.html">http://www.who.int/medicines/areas/coordination/coordination\_assessment/en/index1.html</a>

WHO has prepared a template profile (included in the CD) that can be easily used by countries and that will help presenting data in the form of tables, graphs and charts. Countries can use the generic template provided by WHO and add the information in the questionnaire. Below you can find an example of the template that shows how fields can be changed according to the specific responses provided by each country.

Country X 15/15 not a m	ember of the World Trade Org	anization. The country
has/has no patent law.	National Legislation has/has r	not been modified to
implement the TRIPS A	greement. Country X is/is not	eligible for the
transitional period to 2	016.	
The following (TRIPS) f law:	lexibilities and safeguards are	present in the national
Compulsory licensing p	rovisions that can be applied f	or Yes/NO
	h	

In each section of the questionnaire you will find some comment boxes that you can use to expand on the answer to one or more questions. The text of these comments can also be included in the profile in order to present the country situation in more detail.

In the questionnaire you are also asked to indicate the source and date of each piece of information you provide; these should be used to develop bibliographic references for the profile.

If you prefer, WHO can develop the narrative profile and the Organization will then share the document with the country, which will own/maintain the copyright for it and will be able to publish it as a national document.

Section (	Section 0 General Info				
0.01 Conta	0.01 Contact Info				
0.01.01	Country (precoded)	Malawi			
0.01.02	Name coordinator				
0.01.03	Address (Street, City)				
0.01.04	Phone number				
0.01.05	Email address				
0.01.06	Web address				
0.01.07	Institution				

Section 1	Health and Demographic data	l		
1.00 Respo	ndent Information Section 1			_
1.00.01	Name of person responsible for filling out Survey section 1			
1.00.02	Phone number			
1.00.03	Email address			
1.00.04	Other respondents for filling out this section			
1.01 Demo	graphic and Socioeconomic Indicate	ors		
Core quest	ions ( <u>click here for help</u> )			
			Year	Source
1.01.01	Population, total (,000)	13077	2008	NSO
1.01.02	Population growth rate (Annual %)	2.8	2008	NSO
1.01.03	Total <u>Gross Domestic Product</u> (GDP) (millions US\$)	4,974.86	2009	World Bank data
1.01.04	GDP growth (Annual %)	7.70	2009	World Bank data
1.01.05C	<u>GDP</u> per capita (US\$ current <u>exchange rate</u> )	867.063	2008	World Bank Data
1.01.06	Comments and References			
Suppleme	ntary questions ( <u>click here for help</u>			
			Year	Source
1.01.07S	Population < 15 years (% of total population)	46.2	2009	UN data

1.01.08S	Population > 60 years (% of total population)	5	2008	WHS
1.01.09S	Urban population (% of total population)	19	2008	WHS
1.01.10S	Fertility rate, total (Births per woman)	5.7	2010	DHS
1.01.11S	Population living with less than \$1.25/day (international PPP) (%)	73.86	2004	World Bank data
1.01.12S	Population living below nationally defined poverty line (%)	52.4	2005	World Bank data
1.01.13S	Income share held by lowest 20% of the population (% of national income)	6.98	2004	World Bank data
1.01.14S	Adult literacy rate, 15+ years (% of relevant population)	73	2011	WHS
1.01.15S	Comments and References			

## 1.02 Mortality and Causes of Death

## Core questions (<u>click here for help</u>)

			Year	Source
1.02.01	Life expectancy at birth for men (Years)	52	2008	WHS
1.02.02	Life expectancy at birth for women (Years)	54	2008	WHS
1.02.03	Infant mortality rate, between birth and age 1 (/1,000 live births)	65	2008	WHS
1.02.04	Under 5 mortality rate (/1,000 live births)	100	2008	WHS
1.02.05	Maternal mortality ratio (/100,000 live births)	510	2011	WHS
1.02.06	Please provide a list of top 10 diseases causing mortality			

1.02.06.01	Disease 1	Malaria
1.02.06.02	Disease 2	Pneumonia
1.02.06.03	Disease 3	Anaemia
1.02.06.04	Disease 4	HIV
1.02.06.05	Disease 5	Meningitis
1.02.06.06	Disease 6	Diarrhoea
1.02.06.07	Disease 7	Malnutrition
1.02.06.08	Disease 8	Upper Respiratory Infections
1.02.06.09	Disease 9	Hypertension
1.02.06.10	Disease 10	Cancer
1.02.07	Please provide a list of top 10 diseases causing morbidity	
1.02.07.01	Disease 1	Malaria
1.02.07.02	Disease 2	ARI
1.02.07.03	Disease 3	Skin Infection
1.02.07.04	Disease 4	Oral Conditions
1.02.07.05	Disease 5	Common Injuries and Wounds
1.02.07.06	Disease 6	Diarrhoea no-bloody V/5 yrs
1.02.07.07	Disease 7	Eye Infections
1.02.07.08	Disease 8	STI
1.02.07.09	Disease 9	Measles
1.02.07.10	Disease 10	Ear Infection
1.02.08	Comments and References	HMIS, 2009, 2010

			Year	Source
1.02.09S	Adult mortality rate for both sexes between 15 and 60 years (/1,000 population)	481	2008	WHS
1.02.10S	Neonatal mortality rate (/1,000 live births)	31	2010	Malawi DHS
1.02.11S	Age-standardized mortality rate by non-communicable diseases (/100,000 population)	796	2004	WHS
1.02.12S	Age-standardized mortality rate by cardiovascular diseases (/100,000 population)	376	2009	WHS
1.02.13S	Age-standardized mortality rate by cancer (/100,000 population)	140	2009	WHS
1.02.14S	Mortality rate for HIV/AIDS (/100,000 population)			
1.02.15S	Mortality rate for tuberculosis (/100,000 population)	23	2008	WHS
1.02.16S	Mortality rate for Malaria (/100,000 population)	95	2006	WHS
1.02.17S	Comments and References		I	1

Section 2	Health Services			
2.00 Respo	ondent Information Section 2			
2.00.01	Name of person responsible for filling out this section of the instrument			
2.00.02	Phone number			
2.00.03	Email address			
2.00.04	Other respondents for filling out this section			
2.01 Healt	h Expenditures		_	
Core quest	tions ( <u>click here for help</u> )			
			Year	Source
2.01.01.01	Total annual expenditure on health (millions NCU)	39,164.25	2008	NHA data
2.01.01.02	Total annual expenditure on health (millions US\$ average exchange rate)	277.67	2008	NHA data
2.01.02C	Total health expenditure as % of Gross Domestic Product	6.50		
2.01.03.01C	Total annual <u>expenditure on health</u> per capita (NCU)	2,638.03		
2.01.03.02C	Total annual expenditure on health per capita (US\$ average exchange rate)	18.70		
2.01.04.01	General government annual expenditure on health (millions NCU)	23,275.18	2008	NHA data
2.01.04.02	General government annual expenditure on health (millions US\$ average exchange rate)	165.02	2008	NHA data
2.01.05	Government annual expenditure on health as percentage of total government budget (% of total government budget)	11.92	2008	NHA data

2.01.06C       Government annual expenditure on health (s)       59.43       2008       NHA         2.01.07.01C       Annual per capita government expenditure on health (NCU)       1.567.77         2.01.07.02C       Annual per capita government expenditure on health (NCU)       1.12         2.01.07.02C       Annual per capita government expenditure on health (US\$ average exchange rate)       11.12         2.01.07.02C       Annual per capita government expenditure on health (US\$ average exchange rate)       40.57         2.01.08C       Private health expenditure as % of total expenditure on health) isservice or public health isservice or public health isservice or other sickness funds of total population       40.57         2.01.09       Population covered by a public health isservice or other sickness funds of total population)       60         2.01.10       Population covered by a public health isservice or other sickness funds of total population)       60         2.01.10       Population covered by expenditure enditure enditure (millions NCU)       60         2.01.11.02       Total pharmaceutical expenditure enditure endi				1	
2.01.07.02C       Annual per capita government expenditure on health (US\$ average exchange rate)       11.12         2.01.07.02C       Annual per capita government expenditure on health (US\$ average exchange rate)       11.12         2.01.08C       Private health expenditure (% of total expenditure on health)       40.57       2011       WHS         2.01.09       Population covered by a public health service or public health insurance or social health insurance, or other sickness funds of total population)       Image: Comparison of total population       Imag	2.01.06C	health as % of total expenditure on health (% of total expenditure on	59.43	2008	NHA
2.01.08C       Private health (US\$ average exchange rate)         2.01.08C       Private health expenditure as % of total expenditure on health)       40.57       2011       WHS         2.01.09       Population covered by a public health service or public health insurance or social health insurance, or other sickness funds of total population)       40.57       2011       WHS         2.01.09       Population covered by a public health insurance or social health insurance, or other sickness funds of total population) $\bigcirc$ $\bigcirc$ 2.01.10       Population covered by private health insurance (% of total population) $\bigcirc$ $\bigcirc$ 2.01.11.01       Total pharmaceutical expenditure (millions NCU) $\bigcirc$ $\bigcirc$ 2.01.12.01C       Total pharmaceutical expenditure per capita (NCU)       PREFILL CALC $\bigcirc$ 2.01.12.02C       Total pharmaceutical expenditure per capita (NCU)       PREFILL CALC $\bigcirc$ 2.01.12.02C       Total pharmaceutical expenditure per capita (NCU) $\bigcirc$ $\bigcirc$ 2.01.12.02C       Total pharmaceutical expenditure per capita (NCU) $\bigcirc$ $\bigcirc$ 2.01.12.02C       Total pharmaceutical expenditure per capita (NCU) $\bigcirc$ $\bigcirc$ 2.01.13C       Pharmaceutical expenditure per capita (US\$ current exchange rate) $\bigcirc$ $\bigcirc$ 2.01.13C	2.01.07.01C		1,567.77		
2.01.09       Population covered by a public health service or public health insurance or social health insurance, or other sickness funds of total population)       Image: Comparison of the comparison of the comparison of total population)         2.01.10       Population covered by private health insurance (% of total population)       Image: Comparison of total population)       Image: Comparison of total population)         2.01.10       Population covered by private health insurance (% of total population)       Image: Comparison of total population)       Image: Comparison of total population)         2.01.11.01       Total pharmaceutical expenditure (millions NCU)       Image: Comparison of total population       Image: Comparison of total population)         2.01.11.02       Total pharmaceutical expenditure (millions US\$ current exchange rate)       PREFILL CALC         2.01.12.01C       Total pharmaceutical expenditure per capita (NCU)       PREFILL CALC         2.01.12.02C       Total pharmaceutical expenditure per capita (US\$ current exchange rate)       PREFILL CALC         2.01.12.02C       Total pharmaceutical expenditure per capita (US\$ current exchange rate)       PREFILL CALC         2.01.13C       Pharmaceutical expenditure as a %       PREFILL CALC	2.01.07.02C	expenditure on health (US\$ average	11.12		
service or public health insurance or social health insurance, or other sickness funds of total population)       Image: Constraint of total population covered by private health insurance (% of total population)         2.01.10       Population covered by private health insurance (% of total population)       Image: Constraint of total population)         2.01.11.01       Total pharmaceutical expenditure (millions NCU)       Image: Constraint of total population         2.01.11.02       Total pharmaceutical expenditure per capita (NCU)       Image: PREFILL CALC         2.01.12.01C       Total pharmaceutical expenditure per capita (US\$ current exchange rate)       Image: PREFILL CALC         2.01.12.02C       Total pharmaceutical expenditure per capita (US\$ current exchange rate)       Image: PREFILL CALC         2.01.13C       Pharmaceutical expenditure as a %       Image: PREFILL CALC	2.01.08C	total health expenditure (% of total	40.57	2011	WHS
2.01.11.01       Total pharmaceutical expenditure (millions NCU)         2.01.11.02       Total pharmaceutical expenditure (millions US\$ current exchange rate)         2.01.12.01C       Total pharmaceutical expenditure per capita (NCU)         2.01.12.02C       Total pharmaceutical expenditure per capita (US\$ current exchange rate)         2.01.12.02C       Total pharmaceutical expenditure per capita (US\$ current exchange rate)         2.01.12.02C       Total pharmaceutical expenditure per capita (US\$ current exchange rate)         2.01.13C       Pharmaceutical expenditure as a %	2.01.09	service or public health insurance or social health insurance, or other sickness funds of total			
2.01.11.02       Total pharmaceutical expenditure (millions US\$ current exchange rate)         2.01.12.01C       Total pharmaceutical expenditure per capita (NCU)       PREFILL CALC         2.01.12.02C       Total pharmaceutical expenditure per capita (US\$ current exchange rate)       PREFILL CALC         2.01.13C       Pharmaceutical expenditure as a %       PREFILL CALC	2.01.10	private health insurance (%			
2.01.12.01C       Total pharmaceutical expenditure per capita (NCU)       PREFILL CALC         2.01.12.02C       Total pharmaceutical expenditure per capita (US\$ current exchange rate)       PREFILL CALC         2.01.13C       Pharmaceutical expenditure as a %       PREFILL CALC	2.01.11.01				
PREFILL CALC       2.01.12.02C     Total pharmaceutical expenditure per capita (US\$ current exchange rate)       2.01.13C     Pharmaceutical expenditure as a %	2.01.11.02				
2.01.13C     Pharmaceutical expenditure as a %	2.01.12.01C		PREFILL CALC		
	2.01.12.02C		PREFILL CALC		
	2.01.13C	Pharmaceutical expenditure as a % of GDP (% of GDP)	PREFILL CALC		
2.01.14C Pharmaceutical expenditure as a % of <u>Health Expenditure</u> (% of total health expenditure) PREFILL CALC	2.01.14C	of <u>Health Expenditure</u> (% of total	PREFILL CALC		
2.01.15.01 Total public expenditure on	2.01.15.01	Total public expenditure on			

	pharmaceuticals (millions NCU)			
2.01.15.02	Total public expenditure on pharmaceuticals (millions US\$ current exchange rate)			
2.01.16C	Share of public expenditure on pharmaceuticals as percentage of total expenditure on pharmaceuticals (%)	PREFILL CALC		
2.01.17.01C	Total public expenditure on pharmaceuticals per capita (NCU)	PREFILL CALC		
2.01.17.02C	Total public expenditure on pharmaceuticals per capita (US\$ current exchange rate)	PREFILL CALC		
2.01.18.01	Total private expenditure on pharmaceuticals (millions NCU)			
2.01.18.02	Total private expenditure on pharmaceuticals (millions US\$ current exchange rate)			
2.01.19	Comments and References			
Suppleme	ntary questions ( <u>click for help</u> )			
			Year	Source
2.01.20S	Social security expenditure as % of government expenditure on health (% of government expenditure on health)	0	2008	NHA
2.01.21S	Market share of generic pharmaceuticals [ <u>branded</u> and <u>INN</u> ] by value (%)			
2.01.22S	Annual growth rate of total pharmaceuticals market value (%)			
2.01.23S	Annual growth rate of generic pharmaceuticals market value (%)			

2.01.24S	Private <u>out-of-pocket</u> expenditure as % of private health expenditure (% of private expenditure on health)	28.34	2008	NHA data
2.01.25S	Premiums for private prepaid health plans as % of total private health expenditure (% of private expenditure on health)	15.62	2008	NHA data
2.01.26S	Comments and References			
2.02 Healt	n Personnel and Infrastructure			
	ions <u>(click for help)</u>			
			Year	Source
2.02.01	Total number of pharmacists licensed/registered to practice in your country	136	2009	SADC survey
2.02.02C	Pharmacists per 10,000 population	0.092		
2.02.03	Total number of pharmacists working in the public sector	10	2011	Ministry of Health
2.02.04	Total number of pharmaceutical         technicians and assistants	209	2010	MoH HR Strategic Plan
2.02.05	A strategic plan for pharmaceutical human resource development is in place in your country?	Yes 🗌 No 🗌		
2.02.06	Total number of physicians	257	2009	WHS
2.02.07C	Physicians per 10,000 pop	0.2		
2.02.08	Total number of <u>nursing and</u> <u>midwifery personnel</u>	3,896	2009	WHS
2.02.09C	Nurses and midwives per 10,000 pop	2.6		
2.02.10	Total number of hospitals	94	2009	SADC survey

2.02.11	Number of hospital beds per 10,000 pop	11	2009	WHS		
2.02.12	Total number of primary health care units and centers	1,396	2009	SADC survey		
2.02.13	Total number of licensed pharmacies	78	2010	РМРВ		
2.02.14	Comments and References					
Suppleme	Supplementary questions ( <u>click here for help</u> )					
	-		Year	Source		
2.02.15S	Starting annual salary for a newly registered <u>pharmacist</u> in the public sector (NCU)	513216	2008	Ministry of Health		
2.02.16S	Total number of pharmacists who graduated (first degree) in the past 2 years in your country	25		РМРВ		
2.02.17S	Are there <u>accreditation</u> requirements for pharmacy schools?	Yes 🛛 No		РМРВ		
2.02.18S	Is the Pharmacy Curriculum regularly reviewed?	Yes 🖾 No 🗌	2010	PMPB		
2.02.19S						

Section	3 Policy issues			
3.00 Res	pondent Information Section 4			
3.00.01	Name of person responsible for filling out this section of the instrument			
3.00.02	Phone number			
3.00.03	Email address			
3.00.04	Other respondents for filling out this section			
3.01 Poli	icy Framework			
	estions ( <u>click here for help</u> )			
			Year	Source
3.01.01	National Health Policy exists. If yes, please write year of the most recent document in the "year" field.	Yes 🛛 No 🗌	2007	SADC
3.01.02	National Health Policy Implementation plan exists. If yes, please write the year of the most recent document in the "year"	Yes 🛛 No 🗌	2007	MOH - Malawi
3.01.03	Please provide comments on the Health policy and its implementation plan		I	
3.01.04	National Medicines Policy official document exists. If yes, please write the year of the most recent document in the "year" field.	Yes 🛛 No 🗌	2009	MOH - Malawi
3.01.05	Group of policies addressing pharmaceuticals exist.	Yes 🗌 No 🗌		
3.01.06	National Medicines Policy covers the following components:			

3.01.06.01	Selection of Essential Medicines	⊠Yes		
3.01.06.02	Medicines Financing	⊠Yes		
3.01.06.03	Medicines Pricing	⊠Yes		
3.01.06.04	Medicines Procurement	⊠Yes		
3.01.06.05	Medicines Distribution	⊠Yes		
3.01.06.06	Medicines Regulation	⊠Yes		
3.01.06.07	Pharmacovigilance	⊠Yes		
3.01.06.08	Rational Use of Medicines	⊠Yes		
3.01.06.09	Human Resource Development	⊠Yes		
3.01.06.10	Research	⊠Yes		
3.01.06.11	Monitoring and Evaluation	⊠Yes		
3.01.06.12	Traditional Medicine	⊠Yes		
3.01.07	National medicines policy implementation plan exists. If yes, please write year of the most recent document.	Yes 🛛 No 🗌	2009	SADC survey
3.01.08	Policy or group of policies on clinical laboratories exist. If yes, please write year of the most recent document in the "year" field	Yes 🛛 No 🗌	2007	MOH Malawi
3.01.09	National clinical laboratory policy implementation plan exists. If yes, please write year of the most recent document in the "year" field	Yes 🖾 No 🗌	2009	MOH - Malawi
3.01.10	Access to essential medicines/technologies as part of the fulfillment of the right to health, recognized in the constitution or national legislation?	Yes 🗌 No 🗌		

3.01.11	There are official written guidelines on medicines donations.	Yes 🖾 No 🗌	2007	WHO level I
3.01.12	Is pharmaceutical policy implementation being regularly monitored/assessed?	Yes 🗌 No 🗌		
3.01.12.01	Who is responsible for pharmaceutical policy monitoring?			
3.01.13	Is there a national <u>good governance</u> policy?	Yes 🖂 No 🗌		
3.01.13.01	Multisectoral	⊠Yes		
3.01.13.02	For the pharmaceutical sector	⊠Yes		
3.01.13.03	Which agencies are responsible?			
3.01.14	A policy is in place to manage and sanction <u>conflict of interest</u> issues in pharmaceutical affairs.	Yes 🗌 No 🗍		
3.01.15	There is a formal code of conduct for public officials.	Yes 🗌 No 🗍		
3.01.16	Is there a <u>whistle-blowing</u> mechanism allowing individuals to raise a concern about wrongdoing occurring in the pharmaceutical sector of your country (ombudsperson)?	Yes 🗌 No 🗌		
3.01.16.01	Please describe:			
3.01.17	Comments and References			

Section 4	Section 4 Medicines Trade and Production				
4.00 Resp	ondent Information Section 4				
4.00.01	Name of person responsible for filling out this section of the instrument				
4.00.02	Phone number				
4.00.03	Email address				
4.00.04	Other respondents for filling out this section				
401 Intel	lectual Property Laws and Medicine				
	Core questions (click here for help)				
			X	2	
4.01.01	Country is a member of the World Trade Organization	Yes 🖾 No	Year 1995	Source WTO	
4.01.02	Legal provisions provide for granting of Patents on:		1958	Patent Law	
4.01.02.01	Pharmaceuticals	Yes 🛛 No			
4.01.02.02		Yes 🛛 No 🗌			
	Medical supplies	Yes 🛛 No 🗌			
4.01.02.04	Medical equipment				
4.01.03.01	Please provide name and address of the institution responsible for managing and enforcing intellectual property rights	Registrar of companies			
4.01.03.02	Please provide <u>URL</u>				
4.01.04	National Legislation has been modified to implement the <u>TRIPS</u> <u>Agreement</u>	Yes 🗌 No 🖾	2009	Registry General	
4.01.05	Current laws contain (TRIPS)	Yes 🗌 No 🛛	2009	SADC	

	flexibilities and safeguards			Survey
4.01.06	Country is eligible for the transitional period to 2016	Yes 🛛 No	2010	WTO
4.01.07	Which of the following (TRIPS) flexibilities and safeguards are present in the national law?		2009	SADC Survey
4.01.07.01	Compulsory licensing provisions that can be applied for reasons of public health	Yes 🗌 No 🖾		
4.01.07.02	Bolar exception	Yes 🗌 No 🔀		
4.01.08	Are <u>parallel importing</u> provisions present in the national law?	Yes 🗌 No 🔀	2009	SADC Survey
4.01.09	The country is engaged in initiatives to strengthen capacity to manage and apply intellectual property rights to contribute to innovation and promote public health	Yes 🗌 No 🗍		
4.01.10	Are there legal provisions for <u>data</u> <u>exclusivity</u> for pharmaceuticals	Yes 🗌 No 🗍		
4.01.11	Legal provisions exist for <u>patent</u> extension	Yes 🗌 No 🗌		
4.01.12	Legal provisions exist for linkage between patent status and <u>Marketing</u> <u>Authorization</u>	Yes 🗌 No 🗍		
4.01.13	Comments and References			
4.02 Manu	facturing			
	ions ( <u>click here for help</u> )			
			Year	Source
4.02.01	Number of licensed pharmaceutical manufacturers in the country	4	2009	PMPB
4.02.02	Country has manufacturing capacity		2007	WHO level I

r				
4.02.02.01	R&D to discover new active substances	Yes 🗌 No 🖾 Unknown 🗌		
4.02.02.02	Production of pharmaceutical starting materials ( <u>API</u> s)	Yes 🗌 No 🖾 Unknown 🗌		
4.02.02.03	Production of formulations from pharmaceutical starting material	Yes 🖾 No 🗌 Unknown 🗌		
4.02.02.04	Repackaging of finished dosage forms	Yes 🖾 No 🗌 Unknown 🗌		
4.02.03	Percentage of market share by value produced by domestic manufacturers (%)			
4.02.04	Comments and References			
Suppleme	ntary questions ( <u>click here for help</u>	<u>2</u> )		
			Year	Source
4.02.058	Percentage of market share by volume produced by domestic manufacturers (%)			
4.02.06S	Number of multinational pharmaceutical companies manufacturing medicines locally	0		РМРВ
4.02.07S	Number of manufacturers that are <u>Good Manufacturing Practice</u> (GMP) certified	3	2011	РМРВ
4.02.08S	Comments and References			

Section 5	Section 5 Medicines Regulation				
5.00 Respo	ndent Information Section 4				
5.00.01	Name of person responsible for filling out this section of the instrument				
5.00.02	Phone number				
5.00.03	Email address				
5.00.04	Other respondents for filling out this section				
5.01 Regula	atory Framework				
Core quest	ions ( <u>click here for help</u> )				
			Year	Source	
5.01.01	Are there legal provisions establishing the powers and responsibilities of the <u>Medicines</u> <u>Regulatory Authority</u> (MRA)?	Yes 🖾 No 🗌	2009	SADC survey	
5.01.02	There is a Medicines Regulatory Authority	Yes 🖾 No 🗌	2009	SADC survey	
5.01.03	If yes, please provide name and address of the Medicines regulatory authority	Pharmacy Medicines and Poisons Board o P.O. Box 30241 Lilongwe 3 Malawi			
5.01.04	The Medicines Regulatory Authority is:		2009	PMPB	
5.01.04.01	Part of MoH	□Yes			
5.01.04.02	Semi autonomous agency	⊠Yes			
5.01.04.03	Other (please specify)				

5.01.05	What are the functions of the National Medicines Regulatory Authority?			
5.01.05.01	Marketing authorization / registration	Yes 🖾 No 🗌		
5.01.05.02	Inspection	Yes 🖂 No 🗌		
5.01.05.03	Import control	Yes 🛛 No 🗌		
5.01.05.04	Licensing	Yes 🖾 No 🗌		
5.01.05.05	Market control	Yes 🗌 No 🛛		
5.01.05.06	Quality control	Yes 🖾 No 🗌		
5.01.05.07	Medicines advertising and promotion	Yes 🖾 No 🗌		
5.01.05.08	Clinical trials control	Yes 🗌 No 🗌		
5.01.05.09	Pharmacovigilance	Yes 🖾 No 🗌		
5.01.05.10	Other: (please explain)			
5.01.06	Number of the MRA permanent staff	34	2011	PMPB
5.01.06.01	Date of response	2010		
5.01.07	The MRA has its own website	Yes 🗌 No 🗌		
5.01.07.01	- If yes, please provide MRA Web site address (URL)			
5.01.08	The MRA receives external technical assistance	Yes 🖾 No 🗌	2010	GF Pharmaceu tical Sector Country Profile
5.01.08.01	If yes, please describe:	eg Harmanization of Drug Registration wit	th SADC Cou	ntries.
5.01.09	The MRA is involved in harmonization/ collaboration initiatives	Yes 🖾 No 🗌	2009	WHO level I
5.01.09.01	- If yes, please specify	There is collaboration between NMRAs from	om the entire	SADC

	Region to harmonize guidelines for registration of medicines.				
5.01.10	An assessment of the medicines regulatory system has been conducted in the last five years.	Yes 🖾 No 🗌	2006	РМРВ	
5.01.11	Medicines Regulatory Authority gets funds from regular budget of the government.	Yes 🖾 No 🗌	2007	WHO level I	
5.01.12	Medicines Regulatory Authority is funded from fees for services provided.	Yes 🛛 No 🗌	2007	WHO level I	
5.01.13	Medicines Regulatory Authority receives funds/support from other sources	Yes 🛛 No 🗌			
5.01.13.01	- If yes, please specify	Registration of Medicines/Pharmacies			
5.01.14	Revenues derived from <u>regulatory</u> <u>activities</u> are kept with the Regulatory Authority	Yes 🗌 No 🗍			
5.01.15	The Regulatory Authority is using a computerized information management system to store and retrieve information on registration, inspections, etc.	Yes 🖾 No 🗔			
5.01.16	Comments and References				
	eting Authorization (Registration)				
Core quest	ions ( <u>click here for help</u> )				
			Year	Source	
5.02.01	Legal provisions require a <u>Marketing</u> <u>Authorization</u> (registration) for all pharmaceutical products on the market	Yes 🖾 No 🗌	2007	WHO level I	
5.02.02	Are there any mechanism for	Yes 🖾 No 🗌	2010	GF Pharmaceu	

	exception/waiver of registration?			tical Country Profile
5.02.03	Are there mechanisms for recognition of registration done by other countries	Yes 🗌 No 🔀	2010	GF Pharmaceu tical Country Profile
5.02.03.01	If yes, please explain:	There are mechanisms for waiver of registration of medicines imported for emergencies, epidemics, specific programmes' requirements and for individual special cases.		
		There is no mechanism for recognition of r countries, but work is in progress for harm	-	
5.02.04	Explicit and publicly available criteria exist for assessing applications for Marketing Authorization of pharmaceutical products	Yes 🛛 No 🗌	2009	PMPB
5.02.05	Information from the <u>prequalification</u> programme managed by WHO is used for product registration	Yes 🖾 No 🗌	2010	GF Pharmaceu tical Country Profile
5.02.06	Number of pharmaceutical products registered in your country	4,500	2011	PMPB
5.02.07	Legal provisions require the MRA to make the list of registered pharmaceuticals with defined periodicity publicly available	Yes 🖾 No 🗌	2007	WHO level I
5.02.07.01	If yes, how frequently updated	Quarterly by Expert Medicines Committee		
5.02.07.02	If yes, please provide updated list or URL *	See attached		

5.02.08	Medicines registration always includes the <u>INN (International Non-</u> proprietary Names)	Yes 🖾 No 🗌	2007	WHO level I
5.02.09	Legal provisions require the payment of a fee for Medicines Marketing Authorization (registration) applications	Yes 🛛 No 🖾		
5.02.10	Comments and References			
Suppleme	entary questions ( <u>click here for hel</u>	<u>)</u> )		
			Year	Source
5.02.11S	Legal provisions require Marketing Authorization holders to provide information about variations to the existing Marketing Authorization	Yes 🗌 No 🗌		
5.02.12S	Legal provisions require publication of a <u>Summary of Product</u> <u>Characteristics (SPCs)</u> of the medicines registered	Yes 🗌 No 🗌		
5.02.13S	Legal provisions require the establishment of an expert committee involved in the marketing authorization process	Yes 🖾 No 🗌	2007	WHO level I
5.02.14S	Certificate for Pharmaceutical Products in accordance with the WHO Certification scheme is required as part of the Marketing Authorization application	Yes 🗌 No 🖾	2007	WHO level I
5.02.15S	Legal provisions require declaration of potential <u>conflict of interests</u> for the experts involved in the assessment and decision-making for registration	Yes 🗌 No 🗌		
5.02.16S	Legal provisions allow applicants to appeal against MRAs decisions	Yes 🗌 No 🗌		
5.02.17S	Registration fee - the amount per application for pharmaceutical product containing <u>New Chemical</u>	USD 500	2011	PMPB

	Chemical Entity (NCE) (US\$)			
5.02.18S	Registration fee - the Amount per application for a <u>generic</u> pharmaceutical product (US\$)	USD 500	2011	PMPB
5.02.19S	Time limit for the assessment of a Marketing Authorization application (months)	4	2009	PMPB
5.02.20S	Comments & References			
5.03 Regul	atory Inspection			
Core Quest	ions( <u>click here for help</u> )			
			Year	Source
5.03.01	Legal provisions exist allowing for appointment of government pharmaceutical inspectors	Yes 🖾 No 🗌	1988	PMP AcT
5.03.02	Legal provisions exist permitting inspectors to inspect premises where pharmaceutical activities are performed	Yes 🖾 No 🗌	2007	WHO level I
5.03.02.01	If yes, legal provisions exist requiring inspections to be performed	Yes 🖾 No 🗌		
5.03.03	Inspection is a pre-requisite for licensing of:			
5.03.03.01	Public facilities	Yes 🗌 No 🗌		
5.03.03.02	Private facilities	Yes 🛛 No 🗌		
5.03.04	Inspection requirements are the same for public and private facilities	Yes 🖾 No 🗌		
5.03.05.01	Local manufactures are inspected for GMP compliance	Yes 🖾 No 🗌	2010	GF Pharmaceu tical Country

				Profile
5.03.05.02	Private wholesalers are inspected	Yes 🖾 No 🗌		
5.03.05.03	Retail distributors are inspected	Yes 🖾 No 🗌		
5.03.05.04	Public pharmacies and stores are inspected	Yes 🖾 No 🗌		
5.03.05.05	Pharmacies and dispensing points of health facilities are inspected	Yes 🖾 No 🗌		
5.03.05.06	Please provide details on frequency of inspections for the different categories of facilities	Manufacturers - Biannually Reyailer and Wholesalers - Biannually diagnostics and Dispensing Clinics - Quar	terly	
5.03.06	Comments and References		·	
				_
5.04 Impor	rt Control			
Core Quest	ions ( <u>click here for help</u> )			
		-	Year	Source
5.04.01	Legal provisions exist requiring authorization to import medicines	Yes 🗌 No 🖾		
5.04.02	Legal provisions exist allowing the sampling of imported products for testing	Yes 🗌 No 🖾		
5.04.03	Legal provisions exist requiring importation of medicines through authorized ports of entry	Yes 🗌 No 🗌		
5.04.04	Legal provisions exist allowing inspection of imported pharmaceutical products at the authorized ports of entry	Yes 🗌 No 🗌		
5.04.05	Comments and References			
5.05 Licens	sing			
			Year	Source

5.05.01	Legal provisions exist requiring manufacturers to be licensed	Yes 🖾 No 🗌	2007	WHO level I
5.05.02	Legal provisions exist requiring both domestic and international manufacturers to comply with <u>Good</u> <u>manufacturing Practices (GMP)</u>	Yes 🖾 No 🗌	2008	PMP
5.05.02.01	If no, please explain			
5.05.03	GMP requirements are published by the government.	Yes 🖾 No 🗌	2009	PMPB
5.05.04	Legal provisions exist requiring importers to be licensed	Yes 🖾 No 🗌	2007	WHO level I
5.05.05	Legal provisions exist requiring wholesalers and distributors to be licensed	Yes 🛛 No 🗌	2007	WHO level I
5.05.06	Legal provisions exist requiring wholesalers and distributors to comply with <u>Good Distributing</u> <u>Practices</u> When filling in this part, please also fill in the relevant questions in the procurement and distribution section (Section 7)	Yes 🗌 No 🖂		
5.05.07	National Good Distribution Practice requirements are published by the government	Yes 🗌 No 🖾	2009	CMS
5.05.08	Legal provisions exist requiring pharmacists to be registered	Yes 🖾 No 🗌	1988	PMPB
5.05.09	Legal provisions exists requiring private pharmacies to be licensed	Yes 🖾 No 🗌	1988	PMPB
5.05.10	Legal provision exist requiring public pharmacies to be licensed	Yes 🖾 No 🗌	1988	PMPB
5.05.11	National Good Pharmacy Practice Guidelines are published by the government	Yes 🛛 No 🗌		

5.05.12	Legal provisions require the publication of a list of all licensed pharmaceutical facilities	Yes 🖾 No 🗌	1988	PMPB
5.05.13	Comments and References		1	
5.06 Mark	et Control and Quality Control			
Core Quest	tions ( <mark>click here for help</mark> )			
			Year	Source
5.06.01	Legal Provisions for regulating the pharmaceutical market exist	Yes 🖾 No 🗌		PMPB
5.06.02	Does a laboratory exist in the country for Quality Control testing?	Yes 🖾 No 🗌	2010	GF Pharmaceu tical Country Profile
5.06.02.01	If yes, is the laboratory part of the <u>MRA</u> ?	Yes 🖾 No 🗌		
5.06.02.02	Does the regulatory authority contract services elsewhere?	Yes 🗌 No 🖂		
5.06.02.03	If yes, please describe			
5.06.03	Is there any national laboratory accepted for collaboration with <u>WHO</u> <u>prequalification Programme</u> ? Please describe.	No		
5.06.04	Medicines are tested:		2010	GF Pharmaceu tical Country Profile
5.06.04.01	For quality monitoring in the public sector (routine sampling in pharmacy stores and health facilities)	Yes 🖾 No 🗌		
5.06.04.02	For quality monitoring in private sector (routine sampling in retail	Yes 🛛 No 🗌		
	outlets)			
------------	---	------------	------	----------------
5.06.04.03	When there are complaints or problem reports	Yes 🖾 No 🗌		
5.06.04.04	For product registration	Yes 🛛 No 🗌		
5.06.04.05	For public procurement prequalification	Yes 🗌 No 🖂		
5.06.04.06	For public program products prior to acceptance and/or distribution	Yes 🖾 No 🗌		
5.06.05	Samples are collected by government inspectors for undertaking <u>post-marketing</u> <u>surveillance</u> testing	Yes 🖾 No 🗌	2007	WHO level I
5.06.06	How many Quality Control samples were taken for testing in the last two years?			
5.06.07	Total number of samples tested in the last two years that failed to meet quality standards			
5.06.08	Results of quality testing in past two years are publicly available	Yes 🗌 No 🖾	2009	PMPB
5.06.09	Comments and References			
5.07 Medic	ines Advertising and Promotion			
Core Quest	tions ( <u>click here for help</u> )			
			Year	Source
5.07.01	Legal provisions exist to control the promotion and/or advertising of prescription medicines	Yes 🖾 No 🗌	2007	WHO level I
5.07.02	Who is responsible for regulating, promotion and/or advertising of medicines? Please describe:	PMPB		
5.07.03	Legal provisions prohibit direct advertising of prescription medicines	Yes 🖾 No 🗌	2007	WHO level

	to the public			I
5.07.04	Legal provisions require a pre- approval for medicines advertisements and promotional materials	Yes 🖾 No 🗌	2009	PMPB
5.07.05	Guidelines/Regulations exist for advertising and promotion of non- prescription medicines	Yes 🗌 No 🔀		
5.07.06	A national code of conduct exists concerning advertising and promotion of medicines by marketing authorization holders and is publicly available	Yes 🗌 No 🔀		
5.07.06.01	If yes, the <u>code of conduct</u> applies to domestic manufacturers only, multinational manufacturers only, or both			
	Domestic only	Yes		
	Multinational only	□Yes		
	Both	□Yes		
5.07.06.02	If yes, adherence to the code is voluntary	Yes 🗌 No 🗌		
5.07.06.03	If yes, the code contains a formal process for complaints and sanctions	Yes 🗌 No 🗌		
5.07.06.04	If yes, list of complaints and sanctions for the last two years is publicly available	Yes 🗌 No 🗌		
5.07.07	Comments and References			

5.08 Clinic	al trials			
Core Quest	tions ( <u>click here for help</u> )			
			Year	Source
5.08.01	Legal provisions exist requiring authorization for conducting <u>Clinical</u> <u>Trials</u> by the MRA	Yes 🖾 No 🗌		
5.08.02	Legal provisions exist requiring the agreement by an <u>ethics committee/</u> <u>institutional review board</u> of the Clinical Trials to be performed	Yes 🛛 No 🗌		
5.08.03	Legal provisions exist requiring registration of the clinical trials into international/national/regional registry	Yes 🛛 No 🗌		
5.08.04	Comments and References			
Supplementar	y questions ( <u>click here for help</u> )			
			Year	Source
5.08.05S	Legal provisions exist for GMP compliance of investigational products	Yes 🖾 No 🗌		
5.08.06S	Legal provisions require sponsor, investigator to comply with <u>Good</u> <u>Clinical Practices (GCP)</u>	Yes 🖾 No 🗌		
5.08.07S	National GCP regulations are published by the Government.	Yes 🖾 No 🗌		
5.08.08S	Legal provisions permit inspection of facilities where clinical trials are performed	Yes 🛛 No 🗌		
5.08.09S	Comments and References		L	
		·		
5.09 Contr	olled Medicines			
Core Ques	tions ( <u>click here for help</u> )			
			Date	Source
5.09.01	The country has adopted the			

	following conventions:			
5.09.01.01	Single Convention on Narcotic Drugs, 1961	Yes 🖾 No 🗌	1965	Internation al Narcotics Control Board, 2010
5.09.01.02	The 1972 Protocol amending the Single Convention on Narcotic Drugs, 1961	Yes 🖾 No 🗌	1973	Internation al Narcotics Control Board, 2010
5.09.01.03	Convention on Psychotropic Substances 1971	Yes 🖾 No 🗌	1980	Internation al Narcotics Control Board, 2010
5.09.01.04	United Nations <u>Convention against</u> the Illicit Traffic in Narcotic Drugs and <u>Psychotropic Substances</u> , 1988	Yes 🖾 No 🗌	1995	Internation al Narcotics Control Board, 2010
5.09.02	Laws for the control of narcotic and psychotropic substances, and precursors exist	Yes 🖾 No 🗌		DDA
5.09.03	Annual consumption of Morphine (mg/capita)	9066000.000000	2010	РМРВ
5.09.04	Comments and References			
Suppleme	ntary questions ( <u>click here for help</u>	<u>)</u>		
			Year	Source
5.09.05S	The legal provisions and regulations for the control of narcotic and psychotropic substances, and precursors have been reviewed by a WHO International Expert or Partner Organization to assess the balance between the prevention of abuse and access for medical need	Yes 🗌 No 🖾 Unknown 🗋		

5.09.05.01S	If yes, year of review			
5.09.06S	Annual consumption of Fentanyl (mg/capita)	1000mg	2009	Internation al Narcotics Control Board, 2010
5.09.07S	Annual consumption of Pethidine (mg/capita)	34,192,500 mg	2009	Internation al Narcotics Control Board, 2010
5.09.08S	Annual consumption of Oxycodone (mg/capita)	5,760,000mg		
5.09.09S	Annual consumption of Hydrocodone (mg/capita)	5,760,000mg		
5.09.10S	Annual consumption of Phenobarbital (mg/capita)	1,870,634,000mg		
5.09.11S	Annual consumption of Methadone (mg/capita)	375,000mg		
5.09.12S	Comments and References			

## 5.10 Pharmacovigilance

Core Questions (<u>click here for help</u>)

			Year	Source
5.10.01	There are legal provision in the Medicines Act that provides for <u>pharmacovigilance</u> activities as part of the MRA mandate	Yes 🗌 No 🖾	2009	SADC Survey
5.10.02	Legal provisions exist requiring the <u>Marketing Authorization</u> holder to continuously monitor the safety of their products and report to the MRA	Yes 🗌 No 🖾		
5.10.03	Legal provisions about monitoring Adverse Drug Reactions (ADR) exist	Yes 🗌 No 🛛		

	in your country			
5.10.04	A national pharmacovigilance centre linked to the MRA exists in your country	Yes 🖾 No 🗌	2010	PMPB
5.10.04.01	If a national pharmacovigilance centre exists in your country, how many staff does it employ full- time	2		
5.10.04.02	If a national pharmacovigilance center exists in your country, an analysis report has been published in the last two years.	Yes 🗌 No 🔀		
5.10.04.03	If a national pharmacovigilance center exists in your country, it publishes an ADR bulletin	Yes 🗌 No 🔀		
5.10.05	An official standardized form for reporting ADRs is used in your country	Yes 🖾 No 🗌		
5.10.06	A national Adverse Drug Reactions database exists in your country	Yes 🖾 No 🗌		
5.10.07	How many ADR reports are in the database?	5		PMPB
5.10.08	How many reports have been submitted in the last two years?	5		PMPB
5.10.09	Are ADR reports sent to the WHO database in Uppsala?	Yes 🗌 No 🔀	2009	SADC Survey
5.10.09.01	If yes, number of reports sent in the last two years			
5.10.10	Is there a national ADR or pharmacovigilance advisory committee able to provide technical assistance on causality assessment, risk assessment, risk management, case investigation and, where necessary, crisis management	Yes 🛛 No 🗌		

	including crisis communication?			
5.10.11	Is there a clear communication strategy for routine communication and crises communication?	Yes 🗌 No 🖾		
5.10.12	In the absence of a national pharmacovigilance system, ADRs are monitored in at least one public health program (for example TB, HIV, AIDS)?	Yes 🗌 No 🔀		
5.10.13	Please describe how you intend to enhance the Pharmacovigilance system	To scrutinize the general public through media		
5.10.14	Comments and References			
Suppleme	ntary questions ( <u>click here for hel</u>	<u>)</u>		
			Year	Source
5.10.15S	Feedback is provided to reporters	Yes 🛛 No 🗌		
5.10.16S	The ADR database is computerized	Yes 🖾 No 🗌		
5.10.17S	Medication errors (MEs) are reported	Yes 🖾 No 🗌		
5.10.18S	How many MEs are there in the ADRs database?	2	2010	РМРВ
5.10.19S	There is a <u>risk management plan</u> presented as part of product dossier submitted for Marketing Authorization?	Yes 🖾 No 🗌		
5.10.20S	In the past two years, who has reported ADRs?			
5.10.20.01S	Doctors	⊠Yes		
5.10.20.02S	Nurses	□Yes		
5.10.20.03S	Pharmacists	Yes		
5.10.20.04S	Consumers	Yes		

5.10.20.05S	Pharmaceutical Companies	⊠Yes	 
5.10.20.06S	Others, please specify whom		
5.10.21S	Was there any regulatory decision based on local pharmacovigilance data in the last 2 years?	Yes ⊠ No□	
5.10.22S	Are there training courses in pharmacovigilance?	Yes 🖾 No	
5.10.22.01S	If yes, how many people have been trained in the last two years?	810	
5.10.23S	Comments and References		

Section 6	Medicines Financing			
6.00 Respo	ndent Information Section 5			
6.00.01	Name of person responsible for filling out this section of the instrument			
6.00.02	Phone number			
6.00.03	Email address			
6.00.04	Other respondents for this sections			
6.01 Medic	ines Coverage and Exemptions			
Core Quest	ions ( <mark>click here for help</mark> )			
			Year	Source
6.01.01	Do the followings receive medicines free of charge:		2007	WHO level I
6.01.01.01	Patients who cannot afford them	Yes 🛛 No		
6.01.01.02	Children under 5	Yes 🖾 No		
6.01.01.03	Pregnant women	Yes 🛛 No		
6.01.01.04	Elderly persons	Yes 🖾 No		
6.01.01.05	Please describe/explain your yes answers for questions above			
6.01.02	Is there a public health system or social health insurance scheme or public programme providing medicines free of charge for :		2007	WHO level I
6.01.02.01	All medicines included in the EML	Yes 🛛 No 🗌		
6.01.02.02	Any non-communicable diseases	Yes 🖾 No 🗌		
6.01.02.03	Malaria medicines	Yes 🖾 No 🗌		
6.01.02.04	Tuberculosis medicines	Yes 🛛 No 🗌		
6.01.02.05	Sexually transmitted diseases	Yes 🖾 No 🗌		

	medicines			
6.01.02.06	HIV/AIDS medicines	Yes 🖾 No 🗌		
6.01.02.07	Expanded Program on Immunization (EPI) vaccines	Yes 🖾 No 🗌		
6.01.02.08	If others, please specify			
6.01.02.09	Please describe/explain your yes answers for questions above			
6.01.03	Does a national health insurance, social insurance or other <u>sickness</u> <u>fund</u> provide at least partial <u>medicines</u> <u>coverage</u> ?	Yes 🖾 No 🗌	2009	SADC Survey
6.01.03.01	Does it provide coverage for medicines that are on the EML for inpatients	Yes 🗌 No 🗌		
6.01.03.02	Does it provide coverage for medicines that are on the EML for outpatients	Yes 🗌 No 🗌		
6.01.03.03	Please describe the medicines benefit of public/ <u>social insurance schemes</u>			
6.01.04	Do private health insurance schemes provide any medicines coverage?	Yes 🗌 No 🗌		
6.01.04.01	If yes, is it required to provide coverage for medicines that are on the <u>EML</u> ?	Yes 🗌 No 🗌		
6.01.05	Comments and References			
( 02 P-1				
	nts Fees and Copayments			
Core Quest	ions ( <mark>click here for help</mark> )			
			Year	Source
6.02.01	In your health system, at the point of delivery, are there any <u>co-</u> <u>payment</u> /fee requirements for	Yes 🖾 No 🗌	2007	WHO level i

	consultations			
6.02.02	In your health system, at the point of delivery, are there any co- payment/fee requirements for medicines	Yes 🗌 No 🖾	2007	WHO level I
6.02.03	In practice, (even though this may be contrary to regulations) is revenue from fees or sales of medicines sometimes used to pay the salaries or supplement the income of public health personnel in the same facility?	Yes 🗌 No 🖾	2007	WHO level I
6.02.03.01	Please describe the patient fees and copayments system			
6.02.04	Comments and References			
6 03 Pricin	g Regulation for the Private Sector			
	ions (click here for help)			
une quest				
			Maan	0
6.03.01	Are there legal or regulatory provisions affecting pricing of medicines	Yes 🗌 No 🖾	Year 2007	Source WHO level I
6.03.01 6.03.01.01	provisions affecting pricing of	Yes 🗌 No 🖾 Yes 🗌 No 🖾		
	provisions affecting pricing of medicines If yes, are the provisions aimed at			
6.03.01.01	provisions affecting pricing of medicines If yes, are the provisions aimed at <u>Manufacturers</u> If yes, are the provisions aimed at	Yes 🗌 No 🖾		
6.03.01.01 6.03.01.02	provisions affecting pricing of medicines If yes, are the provisions aimed at <u>Manufacturers</u> If yes, are the provisions aimed at <u>Wholesalers</u> If yes, are the provisions aimed at	Yes 🗌 No 🖾		

6.03.03	Regulations exists retail medicine price should be publicly a	e information		Yes 🗌 No 🖂			2009	SADC Survey
6.03.03.01	-if yes, please expla information is made available							
6.03.04	Comments and Ref	erences						
6.04 Prices	, Availability and A	Affordabili	ty		_	-	-	_
Core Quest	ions ( <mark>click here fo</mark>	<mark>r help</mark> )						
							Year	Source
6.04.01-04	Please state if a me survey using the W methodology has b the past 5 years in <b>If yes</b> , please india survey and use the table <b>If no</b> , but other sur prices and availabi conducted, please fill in this section, b comment box to we results and attach to questionnaire	HO/HAI been conduct your country cate the yea results to fil veys on med lity have been do not use t ut rather use rite some of	ted in y. r of the II in this dicines en hem to e the the	Yes 🗌 No 🗍	Unknown 🗌		2004	WHO
	Basket Of ke	ey medicin	es	Public procurement	Public patient	Private patient		
	Availability (one or both of)	Mean (%)	Orig		6.04.01.01	6.04.01.03		
			LPG		6.04.01.02	6.04.01.04		
		Median (%)	Orig		6.04.02.01	6.04.02.03		

			LPG		6.04.02.02	6.04.02.04		
	Price	Median Price Ratio	Orig	6.04.03.01	6.04.03.03	6.04.03.05		
			LPG	6.04.03.02	6.04.03.04	6.04.03.06		
	Affordability Days' wages of the lowest paid govt worker	Number of days' wages	Orig		6.04.04.01	6.04.04.03		
	for standard treatment with co-trimoxazole for a child respiratory infection		LPG		6.04.04.02	6.04.04.04		
C 04 05	Comments and Ref	oroncos						
6.04.05		erences						
			_		_			
6.05 Price	e Components and A stions ( <u>click here fo</u>	ffordabilit	J					
6.05 Price	e Components and A	ffordabilit	IJ				Year	Source
6.05 Price	e Components and A	ffordabilit r help) rvey of med nas been	icines	Yes 🛛 No 🗋	Unknown 🗌		Year	Source
6.05 Price Core Ques	e Components and A stions (click here for Please state if a sur price components h conducted in the pa	ffordabilit r hein) rvey of med has been hast 5 years in percentage acturer Sellir nsurance ar and final me f key medici	icines n your <u>mark-</u> ng nd dicine	Yes 🛛 No 🗆	Unknown 🗌		Year	Source

6.05.04	Comment and References			
Supplem	entary questions ( <u>click here for help</u>			
6.05.05S	Median percentage contribution of MSP/CIF to final medicine price for a basket of key medicines in the public sector (Median % contribution)			
6.05.06S	Median percentage contribution of MSP/CIF to final medicine price for a basket of key medicines in the private sector (Median % contribution)			
6.05.07S	Median manufacturer selling price (CIF) as percent of final medicine price for a basket of key medicines (%)			
6.05.08S	Median wholesaler selling price as percent of final medicine price for a basket of key medicines (%)			
6.05.09S	Median pharmacist <u>mark-up</u> or <u>dispensing fee</u> as percent of retail price for a basket of key medicines (%)			
6.05.10S	Median percentage contribution of the wholesale mark-up to final medicine price for a basket of key medicines (in the public and private sectors) (%)			
6.05.11S	Median percentage contribution of the retail mark-up to final medicine price for a basket of key medicines (in the public and private sectors) (%)			
6.05.12S	Comment and References			
6.06 Dutio	es and Taxes on Pharmaceuticals (Ma	rket)		
Core Ques	stions ( <u>click here for help</u> )			
			Year	Source

6.06.01	There are <u>duties</u> on imported <u>active</u> <u>pharmaceutical ingredients (APIs)</u>	Yes 🛛 No 🗌	2009	SADC Survey
6.06.02	There are duties on imported <u>finished</u> products	Yes 🗌 No 🖾	2007	WHO level I
6.06.03	VAT (value-added tax) or any other tax is levied on finished pharmaceuticals products	Yes 🗌 No 🛛	2009	SADC Survey
6.06.04	There are provisions for tax exceptions or waivers for pharmaceuticals and health products	Yes 🛛 No 🗌	2010	GF Pharmaceu tical Country Profile
6.06.05	Please specify categories of pharmaceuticals on which the taxes are applied and describe the exemptions and waivers that exist	VAT charged		
6.06.06	Comments and References			
Suppleme	ntary questions ( <u>click here for help</u>			
			Year	Source
6.06.07S	Duty on imported active pharmaceutical ingredients, APIs (%)			
6.06.08S	Duty on imported finished products (%)	NONE		
6.06.09S	VAT on pharmaceutical products (%)			
6.06.10S	Comments and References			

Section 7 Pharmaceutical procurement and distribution						
7.00 Respo	ondent Information Section 6					
7.00.01	Name of person responsible for filling out this section of the instrument					
7.00.02	Phone number					
7.00.03	Email address					
7.00.04	Other respondents for filling out this section					
7.01 Public	c Sector Procurement					
	tions (click.here.for.help)					
Core Quest						
7.01.01	Public sector procurement is:		Date	Source		
7.01.01.01	Decentralized	∐Yes				
7.01.01.02	Centralized and decentralized	⊠Yes				
	•					
7.01.01.03	Please describe	Central Medical Stores procures medicines too procure medicines from other sources w				
7.01.02	If public sector <u>procurement</u> is wholly or partially centralized, it is under the responsibility of a <u>procurement agency</u> which is:					
7.01.02.01	Part of MoH	Yes 🖾 No 🗌				
7.01.02.02	Semi-Autonomous	Yes 🗌 No 🗌				

7.01.02.03	Autonomous	Yes 🗌 No 🗌		
7.01.02.04	A government procurement agency which procures all public goods	Yes 🖾 No 🗌		
7.01.03	Public sector requests for tender documents are publicly available	Yes 🖾 No 🗌	2009	CMS
7.01.04	Public sector tender awards are publicly available	Yes 🖾 No 🗌	2009	CMS
7.01.05	Procurement is based on prequalification of suppliers	Yes 🗌 No 🗌		
7.01.05.01	If yes, please describe how it works	Advert for pre-qualification for suppliers to su type.	upply specific	c product
7.01.06	Comments and References			
Suppleme	ntary questions ( <u>click here for he</u>	( <mark>ql¢</mark>		
			Year	Source
7.01.07S	Is there a written public sector procurement policy?. If yes, please write the year of approval in the "year" field	Yes 🖾 No 🗌	2003	SADC Survey
7.01.08S	Are there legal provisions giving priority in public procurement to goods produced by local manufacturers?	Yes 🖾 No 🗌	2009	SADC Survey
7.01.09S	The key functions of the procurement unit and those of the tender committee are clearly separated	Yes 🖾 No 🗌	2007	WHO level I
7.01.10S	A process exists to ensure the quality of products procured	Yes 🖾 No 🗌	2009	CMS
7.01.10.01S	If yes, the quality assurance process includes <u>pre-qualification</u> of products and suppliers	Yes 🖾 No 🗌		
7.01.10.02S	If yes, explicit criteria and procedures exist for pre-	Yes 🛛 No 🗌		

	qualification of suppliers			
7.01.10.03S	If yes, a list of pre-qualified suppliers and products is publicly available	Yes 🖾 No 🗌		
7.01.11S	List of samples tested during the procurement process and results of quality testing are available	Yes 🖾 No 🗌	2009	CMS
7.01.12S	Which of the following <u>tender</u> methods are used in public sector procurement:	<u> </u>	2009	CMS
7.01.12.01S	National competitive tenders	Yes 🗌 No 🗌		
7.01.12.02S	International competitive tenders	Yes 🖾 No 🗌		
7.01.12.03S	Direct purchasing	Yes 🖾 No 🗌		
7.01.13S	Comments and References			
	1			
	c Sector Distribution			
	c Sector Distribution tions ( <u>click here for help</u> )			
			Year	Source
		Yes 🛛 No 🗌	Year	Source
Core Quest	tions (click here for help) The government supply system department has a Central Medical	Yes 🖾 No 🗌	Year	Source
Core Quest 7.02.01	tions (click here for help) The government supply system department has a Central Medical Store at National Level Number of public warehouses in the secondary tier of public distribution	Yes ⊠ No □ Yes □ No ⊠	Year  Year  2009	Source
Core Quest 7.02.01 7.02.02	tions (click here for help) The government supply system department has a Central Medical Store at National Level Number of public warehouses in the secondary tier of public distribution (State/Regional/Provincial) © There are national guidelines on			

	it accredit public distribution facilities?			
7.02.05	List of GDP certified warehouses in the public sector exists	Yes 🗌 No 🖂	2009	CMS
7.02.06	List of GDP certified distributors in the public sector exists	Yes 🗌 No 🔀	2009	CMS
7.02.07	Comments and References			
Suppleme	ntary questions ( <u>click here for he</u>	elp)		
			Year	Source
7.02.08S	Which of the following processes is in place at the Central Medical Store:			
7.02.08.01S	Forecasting of order quantities	Yes 🖾 No 🗌		
7.02.08.02S	Requisition/Stock orders	Yes 🖾 No 🗌		
7.02.08.03S	Preparation of picking/packing slips	Yes 🖾 No 🗌		
7.02.08.04S	Reports of stock on hand	Yes 🖾 No 🗌		
7.02.08.05S	Reports of outstanding order lines	Yes 🖾 No 🗌		
7.02.08.06S	Expiry dates management	Yes 🖾 No 🗌		
7.02.08.07S	Batch tracking	Yes 🖾 No 🗌		
7.02.08.08S	Reports of products out of stock	Yes 🖾 No 🗌		
7.02.09S	Percentage % availability of key medicines at the Central Medical Store			
7.02.10S	Average stock-out duration for a basket of medicines at the Central Medical Store, in days			
7.02.11S	Routine Procedure exists to track the expiry dates of medicines at the Central Medical Store	Yes 🗌 No 🗌		

7.02.12S	The Public Central Medical Store is GDP certified by a licensing authority	Yes 🗌 No 🗍		
7.02.13S	The Public Central Medical Store is <u>ISO</u> certified	Yes 🗌 No 🗌		
7.02.14S	The second tier public warehouses are GDP certified by a licensing authority	Yes 🗌 No 🗌		
7.02.15S	The second tier public warehouses are ISO certified	Yes 🗌 No 🗌		
7.02.16S	Comments and References			
7 02 Datas	o Soctor Distribution			
7.03 Priva	e Sector Distribution			
	ions ( <u>click here for help</u> )			
			Year	Source
		Yes 🛛 No 🗌	<u>Year</u> 1988	Source PMPB
Core Quest	t <b>ions (<u>click here for help</u>)</b> Legal provisions exist for licensing	Yes ⊠ No □ Yes □ No ⊠		
Core Quest	Legal provisions exist for licensing wholesalers in the private sector Legal provisions exist for licensing			
Core Quest 7.03.01 7.03.02	Legal provisions exist for licensing wholesalers in the private sector Legal provisions exist for licensing distributors in the private sector List of <u>GDP</u> certified wholesalers in	Yes 🗌 No 🖾		

Section 8	Selection and rational use			
8.00 Respo	ndent Information Section 7			
8.00.01	Name of person responsible for filling out this section of the instrument			
8.00.02	Phone number			
8.00.03	Email address			
8.00.04	Other respondents for filling out this section			
0.01 Notice	a l Churrathurra			
	nal Structures			
Core Quest	ions ( <u>click here for help</u> )			
			Year	Source
8.01.01	National <u>essential medicines list</u> (EML) exists. If yes, please write year of last update of EML in the "year" field	Yes 🖾 No 🗌	2009	Ministry of Health
8.01.01.01	If yes, number of medicines on the EML (no. of <u>INN</u> )			
8.01.01.02	If yes, there is a written process for selecting medicines on the EML	Yes 🗌 No 🗌		
8.01.01.03	If yes, the EML is publicly available	Yes 🗌 No 🗌		
8.01.01.04	If yes, is there any mechanism in place to align the EML with the <u>Standard Treatment Guidelines</u> (STG)	Yes 🖾 No 🗌		
8.01.02	National Standard Treatment Guidelines (STGs) for most common illnesses are produced/endorsed by the MoH. If yes, please insert year of last update of STGs in the "year" field	Yes 🛛 No 🗌	2009	МОН
8.01.03	STGs specific to Primary care exist. Please use the "year" field to	Yes 🗌 No 🛛	2007	WHO level

	write the year of last update of primary care guidelines			I
8.01.04	STGs specific to Secondary care (hospitals) exists. Please use the "year" field to write the year of last update of secondary care STGs.	Yes 🖾 No 🗌	1998	WHO level I
8.01.05	STGs specific to Paediatric conditions exist. Please use the "year" field to write the year of last update of paediatric condition STGs	Yes 🖾 No 🗌	2009	МОН
8.01.06	% of public health facilities with copy of EML (mean)- Survey data	100%	2011	Ministry of Health
8.01.07	% of public health facilities with copy of STGs (mean)- Survey data	100%	2011	Ministry fo Health
8.01.08	A public or independently funded national medicines information centre provides information on medicines to prescribers, dispensers and consumers	Yes 🗌 No 🔀	2007	WHO level I
8.01.09	Public education campaigns on rational medicine use topics have been conducted in the previous two years	Yes 🖾 No 🗌	2011	Ministry of Health
8.01.10	A survey on rational medicine use has been conducted in the previous two years	Yes 🗌 No 🗌		
8.01.11	A national programme or committee (involving government, civil society, and professional bodies) exists to monitor and promote rational use of medicines	Yes 🗌 No 🔀	2007	WHO level I
8.01.12	A written National strategy exists to contain <u>antimicrobial resistance</u> . If yes, please write year of last update of the strategy in the "year" field	Yes 🗌 No 🖾	2007	WHO level I

8.01.13	Comments and References			
Suppleme	ntary questions ( <u>click here for he</u>	<mark>dip</mark> )		
			Year	Source
8.01.14S	The <u>Essential Medicines List (EML</u> ) includes formulations specific for children	Yes 🖾 No 🗌		
8.01.15S	There are explicitly documented criteria for the selection of medicines in the EML	Yes 🖾 No 🗌	2009	МОН
8.01.16S	There is a formal committee or other equivalent structure for the selection of products on the National EML	Yes 🖾 No 🗌	2007	WHO level I
8.01.16.01S	If yes, <u>conflict of interest</u> declarations are required from members of national EML committee	Yes 🗌 No 🖾		
8.01.17S	National medicines formulary exists	Yes 🖾 No 🗌	2007	WHO level I
8.01.18S	Is there a funded national inter- sectoral task force to coordinate the promotion of appropriate use of antimicrobials and prevention of spread of infection?	Yes 🗌 No 🖾	2007	WHO level I
8.01.19S	A national reference laboratory/or any other institution has responsibility for coordinating epidemiological surveillance of <u>antimicrobial resistance</u>	Yes 🖾 No 🗌	2007	WHO level I
8.01.20S	Comments and References		1	
8.02 Presc	ribing			
Core Quest	ions ( <u>click here for help</u> )			
			Year	Source

8.02.01	Legal provisions exist to govern the licensing and prescribing practices of prescriber	Yes 🛛 No 🗌	2007	WHO level I
8.02.02	Legal provisions exist to restrict dispensing by prescribers	Yes 🗌 No 🖾	2009	РМРВ
8.02.03	Do prescribers in the private sector dispense medicines?	Yes 🖾 No 🗌	2007	WHO Level 1
8.02.04	Regulations require hospitals to organize/develop <u>Drug and</u> <u>Therapeutics Committees (DTCs)</u>	Yes 🖾 No 🗌	2007	WHO level I
8.02.05	Do more than half of <u>referral</u> <u>hospitals</u> have a DTC?	Yes 🖾 No 🗌 Unknown 🗌	2007	WHO Level 1
8.02.06	Do more than half of <u>general</u> <u>hospitals</u> have a DTC?	Yes 🖾 No 🗌 Unknown 🗌	2007	WHO Level 1
8.02.07	Do more than half of regions/provinces have a DTC?	Yes 🗌 No 🖾 Unknown 🗌	2007	WHO Level 1
8.02.08	The core medical training curriculum includes components on:		2007	WHO level I
8.02.08.01	Concept of <u>EML</u>	Yes 🖾 No 🗌		
8.02.08.02	Use of <u>STGs</u>	Yes 🖾 No 🗌		
8.02.08.03	Pharmacovigilance	Yes 🗌 No 🖂		
8.02.08.04	Problem based pharmacotherapy	Yes 🖾 No 🗌		
8.02.09	Mandatory continuing education that includes pharmaceutical issues is required for doctors (see <u>physician</u> )	Yes 🗌 No 🖾	2007	WHO level I
8.02.10	Mandatory continuing education that includes pharmaceutical issues is required for <u>nurses</u>	Yes 🖾 No 🗌		
8.02.11	Mandatory continuing education that includes pharmaceutical issues	Yes 🗌 No 🖾	2007	WHO level

	is required for paramedical staff			
8.02.12	Prescribing by <u>INN</u> name is obligatory in:		2007	WHO level I
8.02.12.01	Public sector	Yes 🖾 No 🗌		
8.02.12.02	Private sector	Yes 🗌 No 🖂		
8.02.13	Average number of medicines prescribed per patient contact in public health facilities (mean)	3	2009	МОН
8.02.14	% of medicines prescribed in outpatient public health care facilities that are in the national EML (mean)	100	2009	МОН
8.02.15	% of medicines in outpatient public health care facilities that are prescribed by INN name (mean)			
8.02.16	% of patients in outpatient public health care facilities receiving antibiotics (mean)	70	2009	КСН
8.02.17	% of patients in outpatient public health care facilities receiving injections (mean)	25	2009	КСН
8.02.18	% of prescribed drugs dispensed to patients (mean)	90	2009	КСН
8.02.19	% of medicines adequately labelled in public health facilities (mean)	100	2009	МОН
8.02.20	Comments and References			
Supplementary questions (click here for help)				
			Year	Source
8.02.21S	A professional association code of conduct exists governing professional behaviour of doctors	Yes 🗌 No 🗍		
8.02.22S	A professional association code of conduct exists governing	Yes 🗌 No 🗌		

	professional behaviour of nurses			
8.02.23S	Diarrhoea in children treated with Oral Rehydration Solution (ORS) (%)	100	2009	МОН
8.02.24S	Comments and References		1	
8.03 Dispe	nsing			
Core Quest	tions (click here for help)			
			Year	Source
8.03.01	Legal provisions exist to govern dispensing practices of pharmaceutical personnel	Yes 🖾 No 🗌	2007	WHO Level I
8.03.02	The basic pharmacist training curriculum includes components on:		2007	WHO level I
8.03.02.01	Concept of EML	Yes 🖾 No 🗌		
8.03.02.02	Use of STGs	Yes 🖾 No 🗌		
8.03.02.03	Drug Information	Yes 🗌 No 🗌		
8.03.02.04	Clinical pharmacology	Yes 🗌 No 🗌		
8.03.02.05	Medicines supply management	Yes 🗌 No 🗌		
8.03.03	Mandatory continuing education that includes rational use of medicines is required for pharmacists	Yes 🗌 No 🖾	2007	WHO level I
8.03.04	Generic substitution at the point of dispensing in public sector facilities is allowed	Yes 🛛 No 🗌	2007	WHO level I
8.03.05	Generic substitution at the point of dispensing in private sector facilities is allowed	Yes 🛛 No 🗌	2007	WHO level I
8.03.06	In practice, (even though this may be contrary to regulations) are	Yes 🖾 No 🗌 Unknown 🗌	2007	WHO Level

	antibiotics sometimes <u>sold over-</u> <u>the-counter</u> without any prescription?			1
8.03.07	In practice, (even though this may be contrary to regulations) are injections sometimes sold over-the- counter without any prescription?	Yes 🖾 No 🗌 Unknown 🗌	2007	WHO Level 1
8.03.08	Comments and References			
Suppleme	ntary questions ( <u>click here for he</u>	elp)		
			Year	Source
8.03.09S	A professional association <u>code of</u> <u>conduct</u> exists governing professional behaviour of pharmacists	Yes 🖾 No 🗌	2009	РМРВ
8.03.10S	In practice, (even though this may be contrary to regulations) do the following groups of staff <i>sometimes</i> prescribe <u>prescription-only</u> <u>medicines</u> at the primary care level in the public sector?			
8.03.10.01S	Nurses	Yes 🗌 No 🗌 Unknown 🗌		
8.03.10.02S	Pharmacists	Yes 🗌 No 🗌 Unknown 🗌		
8.03.10.03S	Paramedics	Yes 🗌 No 🗌 Unknown 🗌		
8.03.10.04S	Personnel with less than one month training	Yes 🗌 No 🗌 Unknown 🗌		
8.03.11S	Comments and References			

Section	9 Household data/access				
9.00 Respondent Information section 8					
9.00.01	Name of person responsible for filling out this section of the instrument				
9.00.02	Phone number				
9.00.03	Email address				
9.00.04	Other respondents for filling out this section				
9.01 Data	ı from Household Surveys				
	stions (dick here for help)				
			Year	Source	
9.01.01	What household surveys have been undertaken in the past 5 years to assess access to medicines?		Total	Course	
9.01.02	Adults with acute condition in two- week recall period who took all medicines prescribed by an authorized prescriber (%)	91.1	2003	WHS	
9.01.03	Adults with acute conditions not taking all medicines because they cannot afford them (%)	4.6	2003	WHS	
9.01.04	Adults (from poor households) with an acute health condition in two- week recall period who took all medicines prescribed by an authorized prescriber (%)	90.7	2003	WHS	
9.01.05	Adults (from poor households) with an acute condition in two-week recall period who did not take all medicines because they cannot afford them (%)	5.9	2003	WHS	

9.01.06	Adults with chronic conditions taking all medicines prescribed by an authorized <u>prescriber</u> (%)	92.0	2003	WHS
9.01.07	Adults (from poor households) with chronic conditions not taking all medicines because they cannot afford them (%)	0.0	2003	WHS
9.01.08	Adults (from poor households) with chronic conditions who usually take all medicines prescribed by an authorized prescriber (%)	96.1	2003	WHS
9.01.09	Children (from poor households) with an acute condition in two-week recall period who took all medicines prescribed by an authorized prescriber (%)	91.5	2003	WHS
9.01.10	Percentage of people who obtained the medicines prescribed in the 15 days before the interview (%)			
9.01.11	People who obtained prescribed medicines for free in the 15 days before the interview (%)			
9.01.12	Comments and References			
Suppleme	entary questions ( <u>click here for he</u>			
			Year	Source
9.01.13S	Adults with acute conditions not taking all medicines because the medicines were not available (%)	91.5	2003	WHS
9.01.14S	Adults with chronic conditions not taking all medicines because they cannot afford them (%)	4.0	2003	WHS
9.01.15S	Adults with chronic conditions not taking all medicines because the medicines were not available (%)	93.6	2003	WHS
9.01.16S	Children with acute conditions taking all medicines prescribed by	89.7	2003	WHS

	an authorized prescriber (%)			
9.01.17S	Children with acute conditions not taking all medicines because they cannot afford them (%)	12.5	2003	WHS
9.01.18S	Children with acute conditions not taking all medicines because the medicines were not available (%)	79.4	2003	WHS
9.01.19S	Children (from poor households) with acute conditions not taking all medicines because they cannot afford them (%)	3.7	2003	WHS
9.01.20S	Comments and References		1	<u>-</u>

Glossary