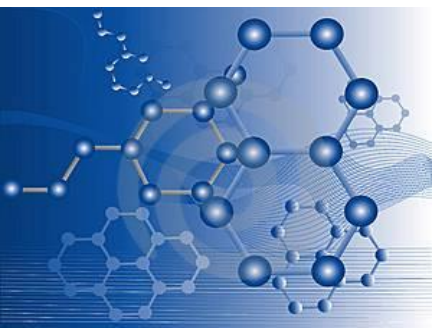


ПРОЦЕДУРА СОВМЕСТНОЙ РЕГИСТРАЦИИ ЛЕКАРСТВЕННЫХ СРЕДСТВ

ОПЫТ НЕБОЛЬШОЙ СТРАНЫ

Лилит Казарян

Научный центр экспертизы
лекарств и медицинских
технологий при МЗ РА



Научный центр
экспертизы лекарств РА

РЕСПУБЛИКА АРМЕНИЯ



Территория

29 800 км²

Население

2 963 000 (2020)

Объем рынка

130 млн US \$



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экспертизы лекарств РА

РЕГУЛИРОВАНИЕ В СФЕРЕ ЛЕКАРСТВ

ЦЕЛЬ

осуществление национальной лекарственной политики для обеспечения

ЭФФЕКТИВНОСТИ

БЕЗОПАСНОСТИ

КАЧЕСТВА

ДОСТУПНОСТИ

лекарств



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РЕГУЛИРОВАНИЕ ЛЕКАРСТВ ТРЕБУЕТ, ПРЕЖДЕ ВСЕГО, СБАЛАНСИРОВАННОСТИ

Защита здоровья населения посредством всесторонней оценки конкретного продукта

Содействие общественному здравоохранению путем предоставления необходимых продуктов без неоправданных задержек

Внедрение гибких подходов к расширению более быстрого доступа к лекарственным средствам без ущерба для строгих стандартов безопасности, качества и эффективности



Научный центр
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2016

ИЗМЕНЕНИЕ СУЩЕСТВУЮЩЕЙ ПРАВОВОЙ БАЗЫ В ЦЕЛЯХ ОБЕСПЕЧЕНИЯ ДОСТУПА К ОСНОВНЫМ ЛЕКАРСТВЕННЫМ СРЕДСТВАМ

ЦЕЛЬ

Избежать

- длительных непредсказуемых сроков оценки
- дублирования усилий регулирующих органов

Свести к минимуму

- Специфичные национальные требования



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экспертизы лекарств РА

ПРАВОВЫЕ И НОРМАТИВНЫЕ ИНСТРУМЕНТЫ ДЛЯ ПОДДЕРЖКИ ДОСТУПНОСТИ

- ◎ Маркировка и листок-вкладыш для пациентов - разрешение нескольких языков (русский, английский), перевод PИ и SmPC со стороны центра
- ◎ Сострадательное использование - бедаквилин, деламанид
- ◎ Сборы - снижение сборов для оценки лекарственных средств с низким оборотом
- ◎ Процедура экстренного импорта незарегистрированных основных лекарственных средств, одобренных другими регулирующими органами или прошедших преквалификацию ВОЗ
- ◎ Сокращенные процедуры оценки, ускоренный обзор на основе предварительного одобрения другими регулирующими органами или преквалификации ВОЗ

(участие в процедурах сотрудничества ВОЗ)



НАЦИОНАЛЬНЫЕ ПРОЦЕДУРЫ РЕГИСТРАЦИИ





- **Сотрудничающий центр ВОЗ по международному мониторингу лекарственных средств, Упсала, с 1997 года**
- **Статус наблюдателя в Европейской фармакопейной комиссии, с 2008 года**
- **Статус наблюдателя в Фармакопейной конвенции США, с 2013 года**
- **Процедура совместной регистрации с ВОЗ, с 2014 года**
- **Статус наблюдателя в ИСН, с 2018 года**



Процедура совместной регистрации



World Health Organization

WHO - Prequalification of Medical Products (IVDs, Medicines, Vaccines and Immunization Devices, Vector Control)

Contact us | Glossary & Acronyms | FAQ

Home | **PRODUCT STREAMS** | EVENTS | NEWS | ABOUT | Search

M Medicines

- + About Medicines Prequalification
- + What We Do
 - Documents A-Z
- + Prequalification Pipeline
- + Prequalified Lists
 - FPPs and APIs Eligible for Prequalification ("EOIs")
- + Prequalification Procedures & Fees: FPPs, APIs & QCLs
- + Post-qualification Procedures & Fees: APIs, FPPs, QCLs

Accelerated Registration of Prequalified FPPs

Finished pharmaceutical products (FPPs) that are WHO-prequalified have been evaluated and inspected according to international standards. But they must still be approved for use by the national medicines regulatory authorities (NMRAs) of the countries for which market entry is sought. Repeating assessment and inspection of those FPPs not only consumes scarce regulatory resources but also extends the time needed to make them available to patients. WHO has therefore designed a collaborative procedure that both enables NMRAs to make use of work already carried out by WHO and to strengthen their own regulatory oversight processes, in line with international best practices. Of greatest interest to manufacturers is that application of the procedure enables faster registration.

The procedure (*Collaborative Procedure between the World Health Organization Prequalification of Medicines Programme and National Medicines Regulatory Authorities in the Assessment and Accelerated National Registration of WHO-prequalified Pharmaceutical Products*) is open to NMRAs in all WHO Member States and holders of prequalified FPPs, on a voluntary basis, and its principles are a model for other regulatory collaborative initiatives.

Information for

- Manufacturers
- Regulatory agencies
- Quality control laboratories
- Procurement agencies

Participating Countries

- Armenia
- Azerbaijan
- Belarus

Преквалификационные отчеты



WHO - Prequalification of Medical Products (IVDs, Medicines, Vaccines and Immunization Devices, Vector Control)

Contact us | Glossary & Acronym

- ↑
- PRODUCT STREAMS ▾
- EVENTS
- NEWS
- ABOUT

M Medicines

- + About Medicines Prequalification
- + What We Do
 - Documents A-Z
- + Prequalification Pipeline
- + Prequalified Lists
 - FPPs and APIs Eligible for Prequalification ("EOIs")
- + Prequalification Procedures & Fees: FPPs, APIs & QCLs
- + Post-prequalification Procedures & Fees: APIs, FPPs, QCLs

Prequalification Reports

Transparency is a key principle of WHO prequalification. For each generic finished pharmaceutical product (FPP) that is prequalified WHO therefore posts on this website a WHO Public Assessment Report which can be accessed via this website:

- [WHO Public Assessment Reports \(WHOPARs\)](#)

A WHO Public Inspection Report (WHOPIR) is posted following inspection of the manufacturing site of a finished pharmaceutical product (FPP), of an active pharmaceutical ingredient, of a contract research organization (that is linked to prequalification of an FPP), or of a quality control laboratory. WHOPIRs are posted only for those sites that have passed inspection and can be accessed via this website:

- [WHO Public Inspection Reports \(WHOPIRs\)](#)

Additional important outputs of WHO prequalification, of particular value to regulators and procurers are:

Information for

- Manufacturers
- Regulatory agencies
- Quality control laboratories
- Procurement agencies

[PRODUCT STREAMS](#)[EVENTS](#)[NEWS](#)[ABOUT](#)

M

[Medicines](#)

Overview of WHO Public Assessment Report (WHOPAR)

Часть 1 - Резюме.

Часть 2 - Все принятые формы выпуска (включая фото)

Часть 3 – Информация для пациента, рекомендованная ВОЗ-PQ.

Часть 4 - Краткое характеристика продукта, рекомендованная ВОЗ-PQ.

Часть 5- Маркировка

Часть 6- Обсуждение, статус во время преквалификации

Часть 7 - Этапы до преквалификации

Часть 8 - Шаги после преквалификации

Part 5 - Label

Part 6 - Discussion (status at the time of prequalification)

MedNet



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MedNet  World Health Organization

Communities for Scientific collaboration,
Information exchange and sharing

Доступ к закрытым отчетам ВОЗ



Communities for Scientific collaboration,
Information exchange and sharing

Welcome to the Community for WHO Collaborative Registration

The collaborative registration procedure serves to facilitate and accelerate national registration of pharmaceutical products which the WHO Prequalification of Medicines Team (WHO/PQT) has already assessed and prequalified. WHO/PQT assessment and inspection reports are shared with participating National Medicines Regulatory Authorities (NMRAs) at the manufacturer's request. The decision about national registration of a product is then expected to be issued within 90 days of information-sharing.

This site serves for sharing of prequalification information with designated regulatory focal points. Access to each relevant product subcommunity is by invitation, subject to a signed confidentiality undertaking and with the prequalification holder's written consent.

More information is found on the WHO/PQT website (<http://apps.who.int/prequal>) under "Collaborative registration".

Sub-communities

HIDE INACTIVE

[5th CRP Meeting Ghana Nov 2017](#)

[7th CRP Annual Meeting, Bangkok, November 2019](#)

[CRP Meeting Denmark May 2018](#)

[CRP Meeting M.M. A. 2016](#)

[6th CRP Meeting Kenya Oct 2018](#)

[8th CRP annual meeting, 23 to 27 November 2020](#)

[CRP Meeting Denmark, May 2019](#)

[P. 2020](#)

Communities I moderate

Communities I participate in HIDE INACTIVE

MedNet

[WHO Collaborative registration procedure](#)

[8th CRP annual meeting, 23 to 27 November 2020](#)

[CRP Meeting Denmark, May 2019](#)

[CRP Meeting Moldova April 2019](#)

[7th CRP Annual Meeting, Bangkok, November 2019](#)

[CRP Meeting Denmark May 2018](#)

[5th CRP Meeting Ghana Nov 2017](#)

[6th CRP Meeting Kenya Oct 2018](#)

[HA722](#)

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[TB070](#)

[TB134](#)

[TB159](#)

[TB178](#)

[TB179](#)

[TB184](#)

[TB226](#)

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All content	24
0 - WHO PQ documents (click on subfolder names to filter)	
1 - Quality assessment-related	7
2 - Bioequivalence assessment-related	3
3 - Assessment of post-prequalification variations	2
4 - Final assessment outcome & latest QIS	7
5 - GMP inspections	
6 - Bioequivalence trial inspections	
Armenia	2
Botswana	1
Mozambique	2

- Appendix2_TRS996_2016_Annex8 consent form.pdf BOTSWANA
- HA722-Appendix2_TRS996_2016_Annex8 consent form (1).pdf
- HA722-Appendix3_PartA_WHO_TRS_996_2016_Annex8 expres
- Appendix3_PartB_HA722.pdf ARMENIA
- Appendix 2_Signed.pdf ARMENIA
- 108HA722Q2.pdf 1 - QUALITY ASSESSMENT-RELATED
- 105HA722Q2.pdf 1 - QUALITY ASSESSMENT-RELATED
- 115-114HA722Q2_LP.pdf 4 - FINAL ASSESSMENT OUTCOME & LATEST QIS
- 120-119HA722Q2.pdf 4 - FINAL ASSESSMENT OUTCOME & LATEST QIS
- 113HA722Q2_LP.pdf 4 - FINAL ASSESSMENT OUTCOME & LATEST QIS
- HA722_QIS_FINAL Feb2021.pdf 4 - FINAL ASSESSMENT OUTCOME & LAT
- 120HA722Q2_LP.pdf 4 - FINAL ASSESSMENT OUTCOME & LATEST QIS
- HA722_QIS_FINAL.pdf 4 - FINAL ASSESSMENT OUTCOME & LATEST QIS
- 114HA722_QIS_FINAL.pdf 4 - FINAL ASSESSMENT OUTCOME & LATEST QI
- 109HA722Q2.pdf 1 - QUALITY ASSESSMENT-RELATED

СТАТУС ЗАЯВЛЕНИЙ ПО ПСР

ОДОБРЕННЫЕ



9

НА РАССМОТРЕНИИ



1

В ОЖИДАНИИ

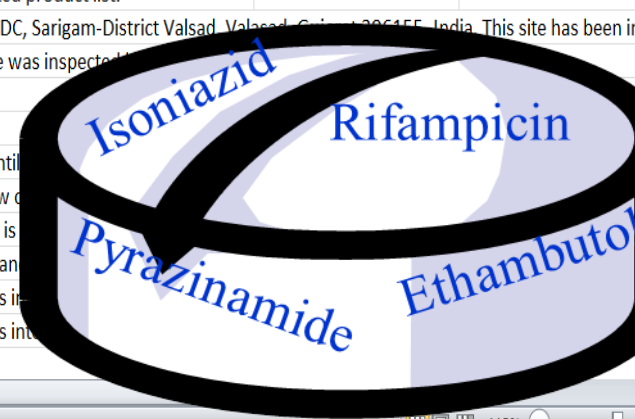


1



Препараты, зарегистрированные в рамках процедуры совместной регистрации ВОЗ

Products registered through the WHO collaborative registration procedure							
List Number 64, updated 12 July 2021							
► Notes and disclaimer							
► Update history							
WHO PQ number	Notes	Product	Prequalification holder	Country of registration	Region	Registration date	Registration number
547		Isoniazid + Rifampicin Tablets 75mg + 150mg	Lupin Ltd	Armenia	EURO	8.Dec.16	16213
552		Ethambutol + Isoniazid + Pyrazinamide + Rifampicin Tablets 275mg + 7	Lupin Ltd	Armenia	EURO	8.Dec.16	16215
564		Ethambutol Tablets 400mg	Macleods Pharmaceuticals Ltd	Armenia	EURO	6.Oct.16	16017
576		Pyrazinamide Tablets 400mg	Macleods Pharmaceuticals Ltd	Armenia	EURO	6.Oct.16	16014
588		Isoniazid Tablets 100mg	Macleods Pharmaceuticals Ltd	Armenia	EURO	6.Oct.16	16015
592		Isoniazid Tablets 300mg	Macleods Pharmaceuticals Ltd	Armenia	EURO	6.Oct.16	16016
610		Isoniazid + Rifampicin Dispersible tablets 30mg + 60mg	Lupin Ltd	Armenia	EURO	5.Oct.17	17059
611		Isoniazid + Pyrazinamide + Rifampicin Dispersible tablets 30mg + 150m	Lupin Ltd	Armenia	EURO	8.Dec.16	16214
622		Ethambutol hydrochloride Tablets 100mg	Macleods Pharmaceuticals Ltd	Armenia	EURO	6.Oct.16	16018
701	Notes:	1 The product registered in-country is technically the same as RH013, except that it includes 7 inert tablets. Collaborative registration was based on shared prequalification information for RH013.					
702		2 Registration details to be confirmed - the registration number and/or date was not yet available at the time of publishing the updated product list.					
703		3 The product was registered with an additional manufacturing site: Famy Care Ltd, Plot no 1606/1609, Sarigam Industrial Estate GIDC, Sarigam-District Valsad, Valsad, Gujarat-386155, India. This site has been inspected.					
704		4 The product was registered with an additional manufacturing site (PLOT NO. 1606/ 1609 GIDC, Sarigam, Valsad, Gujarat). This site was inspected.					
705		5 [Not used - Note regarding conditional registration in Tanzania, subsequently deleted]					
706		6 [Not used - Note regarding shelf life for a product added 12 Feb 2016, deleted 24 Feb 2016]					
707		7 WHO-PQT found data integrity issues with bioequivalence study. Applicant was informed that importation would be suspended until issues are resolved.					
708		8 The product has been registered without considering Anuh Pharma Limited, Boisar, India as a supplier for Pyrazinamide API in view of the quality issues.					
709		9 Of the packaging presentations that have been prequalified by WHO, only the Aluminum foil strip of 5 tablets (1 foil strip in a box) is available.					
710		10 Of the packaging presentations that have been prequalified by WHO, only the blister pack of 10 tablets (10 blister packs in a box) and the blister pack of 28 tablets (24 blister packs in a box) are available.					
711		11 Of the packaging presentations that have been prequalified by WHO, only the blister pack of 28 tablets (24 blister packs in a box) is available.					
712		12 Of the packaging presentations that have been prequalified by WHO, only the blister pack of 10 tablets (10 blister packs in a box) is available.					
713		13 Name change of PQ holder from Jai Pharma to Mylan Laboratories Limited					



ПРОТИВОТУБЕРКУЛЕЗНЫЕ ПРЕПАРАТЫ

Национальный перечень основных лекарственных средств на 2021 год	Типовой перечень основных лекарственных средств ВОЗ на 2021 год	Статус/процедура регистрации
ethambutol	ethambutol	CRP
ethambutol + isoniazid	-----	Nonregistered
ethambutol + isoniazid + pyrazinamide + rifampicin	ethambutol + isoniazid + pyrazinamide + rifampicin	CRP
ethambutol + isoniazid + rifampicin	ethambutol + isoniazid + rifampicin	Nonregistered
isoniazid	isoniazid	CRP
isoniazid + pyrazinamide + rifampicin	isoniazid + pyrazinamide + rifampicin	CRP
isoniazid + rifampicin	isoniazid + rifampicin	CRP
-----	isoniazid + rifapentine	Nonregistered
pyrazinamide	pyrazinamide	CRP
rifabutin	rifabutin	Nonregistered
rifampicin	rifampicin	National/Standard
rifapentine	rifapentine	Nonregistered



Դեղերի և բժշկական տեխնոլոգիաների փորձագիտական կենտրոն

isoni

Search In

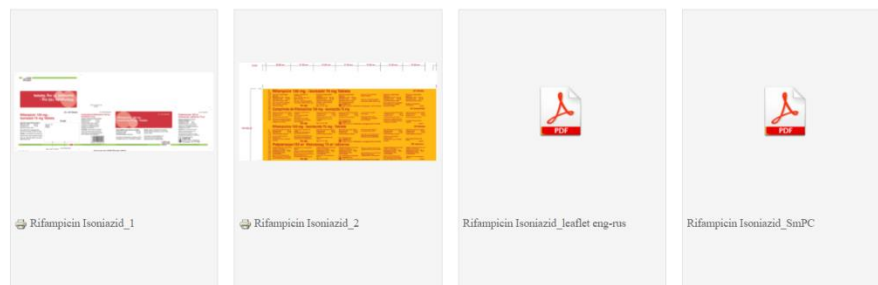
Generic name ▾

English ▾

New search

Login

N	Trade name	Generic name	Dosage form	Dosage strength	Country	Manufacturer	Registered till	Dispensing by	Under Control	Membership country of origin
1	Rifampicin Isoniazid	rifampicin, isoniazid	tablets film-coated	150mg+75mg; (672/24x28) in blister	India	Lupin Limited, A-28/1, M.I.D.C., Chikalthana, Aurangabad 431 210 for Lupin Limited, 159, CST Road, Kalina Santacruz(East), Mumbai 400098, India	08.12.2021	Prescription	-	
2	Rifampicin Isoniazid Pyrazinamide	rifampicin, isoniazid, pyrazinamide	tablets	60mg+30mg+150mg; (84/4x16) in blister						
3	Rifampicin Isoniazid Pyrazinamide Ethambutol hydrochloride	rifampicin, isoniazid, pyrazinamide, ethambutol (ethambutol hydrochloride)	tablets film-coated	150mg+75mg+400mg+275mg; (672/24x28/) in blister						
4	Rifampicin 60mg-Isoniazid 30mg Tablets	rifampicin, isoniazid	tablets dispersible	60mg+30mg; (84/14x6) in strip						



ПРОБЛЕМЫ

- компании не подают заявки для регистрации или перерегистрации, или пострегистрационные изменения
- существуют расхождения между информацией, предоставленной заявителем и оцененной референтным органом
- продукт с действительным МА не выводится на рынок
- **Новая региональная сеть: Евразийский экономический союз**
 - Формат досье
 - Информация о продукте: структура и содержание
 - Совместные оценки
 - Предварительная инспекция



РЕГУЛИРОВАНИЕ ЛЕКАРСТВ ТРЕБУЕТ, ПРЕЖДЕ ВСЕГО, СБАЛАНСИРОВАННОСТИ

Обеспечение более раннего доступа лекарственных средств

Создание стабильной и сильной системы фармаконадзора для своевременного выявления любого риска



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ОСНОВНЫЕ НАПРАВЛЕНИЯ ВНЕДРЕНИЯ СПР



- Установление правовых положений, позволяющих ускорить процесс регистрации
- Разработка руководящих принципов, в том числе по управлению изменениями после утверждения
- Разработка внутренних СОП
- Разработка шаблонов проверки
- Подготовка экспертов-оценщиков
- Назначение координаторов для связи с ВОЗ
- Обновление руководства





Документы, одобренные Комитетом экспертов ВОЗ по спецификациям фармацевтических препаратов

<https://extranet.who.int/prequal/content/who-technical-report-series>

Annex 9

Annex 6

Good practices of national regulatory authorities implementing the collaborative registration procedure for medical products

1. Background
2. Aims and objectives
3. Scope
4. Glossary
5. Key principles
6. Essential elements of a registration system (in the context of collaborative registration procedures)

References

- Appendix 1** An example of information to applicants for registration via the WHO collaborative registration procedure
- Appendix 2** Verification for product submitted under the WHO collaborative procedure
- Appendix 3** Abridged/abbreviated review for product submitted under the WHO collaborative procedure
- Appendix 4** Additional information to be included in the screening checklist
- Appendix 5** Example of a national regulatory authority reliance model approach information, documentary evidence and assessment activity
- Appendix 6** Model acknowledgement or approval letter for variations of product registered through the WHO collaborative procedure

Guidance on good practices for desk assessment, compliance with good manufacturing practices, laboratory practices and good clinical practices for regulatory decisions

Background

1. Introduction
2. Aim and objectives of the guidance
3. Scope of the guidance
4. Glossary
5. Essential elements of desk assessment

- 5.1 High-level support and cooperation
- 5.2 Commonality of quality management systems in inspectorates
- 5.3 Convergent standards of good practices
- 5.4 Reliability and accuracy of information
- 5.5 Management tools to support consistent and objective assessment
- 5.6 Risk-based assessment of available information
- 5.7 Mutual trust and confidence among inspectorates
- 5.8 Quality assurance of the desk assessment process
- 5.9 Communication of assessment outcomes

Annex 11

Collaborative procedure in the assessment and accelerated national registration of pharmaceutical products and vaccines approved by stringent regulatory authorities

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Взгляд в будущее

- Постепенное расширение сферы применения ПСР, охватывая новые продукты, включая медицинские изделия и вакцины
- Продвижение более широкого использования принципов признания на региональном уровне, направленное на сокращение дублирования, укрепление потенциала, содействие гармонизации нормативных положений и установление доверия



ПРЕИМУЩЕСТВА ПРОЦЕДУРЫ СОВМЕСТНОЙ РЕГИСТРАЦИИ



Возможность обмениваться информацией (НРО-ВОЗ) во время оценки (ОТЧЕТЫ об оценке, ОТЧЕТЫ об изменениях, QIS)



Сокращенный обзор и экономия ресурсов (сравнительная оценка, без инспекций и лабораторных испытаний)



Сокращение сроков регистрации и улучшение доступа к качественным лекарствам



Повышение потенциала регулирования и гармонизация практики регулирования: подробные отчеты, участие в тренингах



Контакт:
prequalreg@who.int

**9th Ежегодное
совещание по
процедуре совместно**

Регистрации

виртуальное совещание (Zoom)

**13 – 17
декабря
2021**



**БЛАГОДАРЮ ЗА ВНИМАНИЕ
И СОТРУДНИЧЕСТВО!!!**

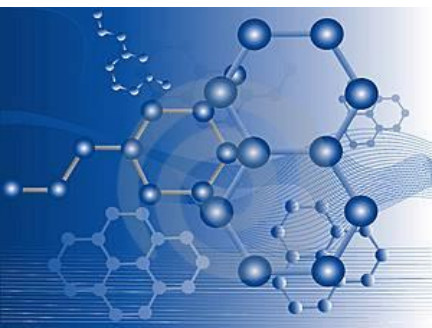


**ՇՆՈՐՀԱԿԱԼՈՒԹՅՈՒՆ
ՈՒՇԱԴՐՈՒԹՅԱՆ ԵՎ ՀԱՄԱԳՈՐԾԱԿՑՈՒԹՅԱՆ
ՀԱՄԱՐ**



**Научный центр
экспертизы лекарств РА**

Вопросы



Научный центр
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