Assuring vaccine quality: Overview of Prequalification

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Outline

• Overview on Prequalification
• Strategic priorities
• Activities to facilitate access of vaccines:
  • Path forward
  • Programmatic suitability for PQ
• Technical assistance and capacity building
Prequalification

- Ensures the quality, efficacy and safety of medicines, vaccines and immunization devices and diagnostics.

- Medicines:
  - Prequalification programme for medicines (finished dosage forms)
  - Prequalification of active pharmaceutical ingredients (APIs)
  - Prequalification of quality control (QC) laboratories
  - Expanding access to priority essential medicines: HIV/AIDS, tuberculosis, Malaria, Reproductive Health and some other disease categories (e.g. NTD)

- Vaccines and immunization devices:
  - Ensures that candidate vaccines are suitable for the target population and meet the needs of the programme
Prequalification is NOT stand alone activity
Many other technical work areas support and link to prequalification (medicines, vaccines, diagnostics and medical devices)

- Outside EMP – Disease oriented departments/programs, IVB Department, Strategic Advisory Group of Experts (SAGE) on Immunization; Regional and Country Offices
- Inside EMP – Norms and standards work/Quality Assurance, Safety/Vigilance, Activities to combat SFFC medical products, NRA strengthening, Policy, Innovation and technology transfer
Prequalification

WHO uses the same scientific principles to assess the products safety, quality and efficacy/performance as well-resourced national regulators:

- scientific assessment of documentary evidence for quality, safety and efficacy
- Assessment of suitability for use of the vaccine in the intended settings
- site inspections for GMP, GLP and GCP
- control of variations to products and their manufacturing processes
- post-approval monitoring of quality and safety
Extensive multilayer collaboration: working with regulators ... for regulators

- Not duplicating work done by stringent regulatory authorities
  - SRA approval of new and generic products – abridged procedure
  - US FDA tentative approvals – based on confidentiality agreement including in the PQ products list
  - European Medicines Agency (EMA) – Art 58 ... and beyond
  - Collaboration with EDQM, in particular in the area of APIs (confidentiality agreements with US FDA, EDQM, EMA ...)

- Active participation and involvement of
  - Regulatory authority experts from well resourced and less resourced settings WORKING TOGETHER for common goal
Why is Vaccines PQ important for user countries and their NRAs?

It represents a source of vaccines of "assured quality"
In addition the evaluation is focused on programmatic needs
WHO follows up on complaints and reports of AEFIIs and publishes the outcome of investigations
WHO monitors the quality of prequalified vaccines on a continuing basis, through testing of samples, reassessment of the products, targeted audits, and delists vaccines if they do not meet the established specifications and/or standard
Opportunity for NRAs in user countries to save resources to focus on other priorities, since registration can be granted through a facilitated and shortened procedure
Technical assistance and capacity building

Meetings with manufacturers at early stages of vaccine development. Advice on product characteristics and clinical development.
PQ briefing workshops
Support to IFPMA and DCVMN
Support to regulatory networks: DCVRN, AVAREF
Supply Security

Monitor closely the performance of prequalified vaccines including FU audits and conducting production capacity assessments.

Actively seek for additional sources for priority vaccines.

Secure the supply base for priority vaccines for developing countries.

Establish risk mitigation strategies in case of failure of NRA.
### Access

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<th>Single standard of quality (WHO recommended requirements)</th>
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<td>Consolidated investigation, reporting and communication in response to quality or safety concerns</td>
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<td>Implementation of an expedited/facilitated registration procedure for prequalified vaccines in receiving countries</td>
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Facilitate access to quality products for developing countries.
Purpose of WHO vaccines prequalification programme

A service provided to UN purchasing agencies.
Provides independent opinion/advice on the quality, safety and efficacy of vaccines for purchase
Ensures that candidate vaccines are suitable for the target population and meet the needs of the programme
Ensures continuing compliance with specifications and established standards of quality
Who Prequalification Programme

Principles

- GMP
- Clinical data
- Consistency of final product characteristics
- Meeting WHO requirements and tender specifications
- Reliance on NRA
Pre-conditions for PQ evaluation

Reliance on the National Regulatory Authority (NRA) of the exporting country

NRA must be assessed as functional as a result of successful evaluation using the WHO NRA assessment tool

NRA’s functional status needs to be sustained over time

Continued regulatory oversight by NRA is required as well as communication with WHO about potential problems with the vaccine

Agreements are established with the NRAs for information exchange when a vaccine is about to be prequalified
Pre-conditions for PQ evaluation

- Vaccine is licensed/registered by the responsible NRA (Article 58 Scientific opinion by EMA accepted)
- WHO guidelines/recommendations approved by the ECBS are available (published in the WHO Technical Report Series)
- Listed in the vaccine priority list – “no priority” vaccines will not be reviewed
Prequalification process

- Scientific review of quality dossier
- Scientific review of clinical data
- Testing of samples
- Consultation with responsible NRA
- Site audit to manufacturing facilities

Current revision of procedure in place from January 2012
Prequalification process: timelines (excluding applicant response times)

1. Submission of application for PQ
2. Screening (30 days + 90 days if there is critical PSPQ non-compliance)
   - 270 days internal time
3. Streamlined based on SRA approval and sharing of NRA reports
   - 90 days internal time
4. Submission of variation
5. Screening
   - 90 days internal time
Role of NRA during PQ process

As part of the evaluation procedure, consultation with NRA

To discuss regulatory status of the concerned vaccine/s
Clinical performance in country of manufacture if used
Quality evaluation, outcome of recent GMP inspections
Compliance with specifications (trends from lot release data)
Regulatory actions
Informal agreement for information sharing with WHO recorded in Consultation report
Programmatic suitability and its assessment

- Vaccines produced in developed countries may not have taken into account programmatic challenges in developing countries.
  - Examples:
    - Non auto-disable prefilled syringe presentations
    - Stability of components in the event of cold chain breakdown

- WHO PQT has always considered programmatic suitability but it was in 2012 that a written guidance (PSPQ) was developed and put in place.
Monitoring performance of PQd vaccines

Targeted testing by WHO contracted labs: Once a year testing of samples of lots shipped to countries to ensure continuing compliance with specifications

Monitoring and resolution of complaints and reports of AEFIs (with collaboration of the responsible NRA)

Reassessments frequency defined on risk analysis basis
Summary

- Prequalification system ensures the quality, efficacy and safety of medicines, vaccines and immunization devices and diagnostics for global use.
- Assessment includes focus on programmatic needs
- Facilitates access to assured quality products for developing countries.
Relevant PQ information

http://www.who.int/immunization_standards/vaccine_quality/pq_system/en/
http://www.who.int/immunization_standards/vaccine_quality/pq_suppliers/en/
http://www.who.int/immunization_standards/vaccine_quality/ps_pq/en/
http://www.who.int/immunization_standards/vaccine_quality/expedited_review/en/