

Date

(00-0000) Page: 1/2

## **Committee on Technical Barriers to Trade**

## **NOTIFICATION**

The following notification is being circulated in accordance with Article 10.6.

1. Notifying Member: BRAZIL

If applicable, name of local government involved (Articles 3.2 and 7.2):

2. Agency responsible: ANVISA – Brazilian Health Surveillance Agency
Name and address (including telephone and fax numbers, e-mail and web-site
addresses, if available) of agency or authority designated to handle comments
regarding the notification shall be indicated if different from above: National
Institute of Metrology, Quality and Technology-INMETRO

Telephone: +(55) 21 2563.2821 Telefax: +(55) 21 2502.6542

Email: <u>barreirastecnicas@inmetro.gov.br</u>
Web-site: <u>www.inmetro.gov.br/barreirastecnicas</u>

The comments to this Draft Regulation shall be sent to

http://formsus.datasus.gov.br/site/formulario.php?id aplicacao=17911

- 3. Notified under Article 2.9.2 [ X ], 2.10.1 [ ], 5.6.2 [ ], 5.7.1 [ ], other:
- Products covered (HS or CCCN where applicable, otherwise national tariff heading.
   ICS numbers may be provided in addition, where applicable): Centres of

Pharmaceutical Equivalence

- **Title, number of pages and language(s) of the notified document:** Draft Technical Resolution no 95, October 9<sup>th</sup> 2014, regarding the petitions of request for qualification, license renewal, post-license changes, outsourcing trial, suspensions and cancellations of Centres of Pharmaceutical Equivalence (22 pages, Portuguese).
- **Description of content:** Draft Technical Resolution no 95, October 9<sup>th</sup> 2014, regarding the petitions of request for qualification, license renewal, post-license changes, outsourcing trial, suspensions and cancellations of Centers of Pharmaceutical Equivalence (22 pages, Portuguese).

This draft Technical Resolution establishes the minimum requirements and procedures to be attended by laboratories that intend to become Centers for Pharmaceutical Equivalence, and by the ones that are already licensed, to the request for qualification, license renewal, post-license changes, outsourcing trial, suspensions and cancellations of Centres of Pharmaceutical Equivalence.

The Centres of Pharmaceutical Equivalence already qualified have 120 days after the publication of the regulation to make the necessary adjustments, in order to comply with this Draft Technical Resolution.

This Draft Techical Resolution revokes article  $4^{th}$  and its paragraphs  $1^{st}$  and  $2^{nd}$ , of the Resolution RDC n. 41/2000.

- 7. Objective and rationale, including the nature of urgent problems where applicable: Protection of Human Health
- **8. Relevant documents:** Brazilian Official Journal (Diário Oficial da União), October 10<sup>th</sup> 2014; Section 1, p. 45, Draft Resolution (Consulta Pública) number 95, October 9<sup>th</sup> 2014, issued by Brazilian Health Surveillance Agency Anvisa. When adopted, it will be published at the Brazilian Official Journal. Available in Portuguese.
- 9. Proposed date of adoption: Proposed date of entry into force:

to be determined after the end of the consultation period.

to be determined after the end of the consultation period.

- **10. Final date for comments:** November 17<sup>th</sup>, 2014.
- 11. Texts available from: National enquiry point [ ] or address, telephone and fax numbers, e-mail and web-site addresses, if available of the other body:

Agency Responsible Brazilian Health Surveillance Agency – ANVISA SIA, Trecho 5, Área Especial 57 Brasília – DF / Brazil

CEP: 71.205-050

Phone.: 55 61 3462-5402 Website: <a href="https://www.anvisa.gov.br">www.anvisa.gov.br</a>

 $\underline{\text{http://portal.anvisa.gov.br/wps/wcm/connect/cb9eab8045c7dcc89f87ffd10ee53f37/Cons}}$ 

ulta+P%C3%BAblica+n%C2%B0+95+GGMED.pdf?MOD=AJPERES