Provisional agenda

1. Opening of the meeting

2. Adoption of the agenda and method of work

3. Update by the Secretariat on the activities and budget to implement the workplan of the Member State mechanism

4. Update on the list of prioritized activities for 2018–2019
   (A) Develop and promote training material and guidance documents to strengthen the capacity of national/regional regulatory authorities for the prevention and detection of and response to substandard and falsified medical products
   (B) Expand and maintain the global focal point network among national medicines regulatory authorities to facilitate cooperation and collaboration
   (C) Improve understanding of Member States on detection technologies, methodologies and “track and trace” models
   (D) Increase Member States’ knowledge of the links between substandard and falsified medical products and access to quality, safe, efficacious and affordable medical products
   (E) Develop and leverage existing activity for effective risk communication and make recommendations for awareness campaigns on substandard and falsified medical products
   (F) Enhance Member States’ capacity to expand awareness, effectiveness, impact and outreach in their work on substandard and falsified medical products
   (G) Promote shared understanding among Member States from a public health perspective regarding medical products in transit
   (H) Identify and develop appropriate strategies to understand and address the distribution or supply of substandard and falsified medical products via the internet

5. WHO participation in the Global Steering Committee for Quality Assurance of Health Products

6. Update on WHO activities for regulatory systems strengthening

7. Update on governance issues
8. Proposed dates of the eighth meeting of the Member State mechanism
9. Report of the Member State mechanism
10. Closure of the meeting