Azithromycin for system suitability

The European Pharmacopoeia is the official intergovernmental body responsible for establishment of quality standards for medicines in Europe. Compliance with the standards is mandatory for any medicine to be sold in Europe. In many cases, to test compliance, pharmaceutical manufacturers have to use a reference substance.

All substances supplied by the European Directorate for the Quality of Medicines are supplied exclusively as European Pharmacopoeia Reference for use as standards or reference materials in tests and assays carried out in accordance with the official methods of the European Pharmacopoeia and for no other purpose.

**Purpose of Material:** this shipment contains drug samples for investigational use only and is intended for laboratory testing only.

**NOT FOR FOOD**

**NOT FOR HUMAN CONSUMPTION**

**FOR LABORATORY USE ONLY**

I give you the origin for the following substance:

<table>
<thead>
<tr>
<th>Code catalogue</th>
<th>Batch number</th>
<th>Origin</th>
</tr>
</thead>
<tbody>
<tr>
<td>Y0000641</td>
<td>10</td>
<td>Synthetic</td>
</tr>
</tbody>
</table>

- The material is not of animal origin;
- The material is chemically synthesized;
- The above material does not contain any animal or cell culture derived products or additives such as albumin or serum.
- The above material was not derived from any animal or cell culture derived products.

Best regards,

**Dr. Fanny MOUTIER-GAME**

*Head of Dispatching Section*

Division of Reference Standards and Samples