1 July 2015

URGENT MEDICINE RECALL (Class III Type B)

Dear Customer

Johnson & Johnson (Pty) Ltd, in consultation with the Medicines Control Council, is recalling the following batches of BENYLIN® products. This is a Class III Type B recall and is intended to reach all distribution points.

This recall ONLY affects the following TWO batches:

<table>
<thead>
<tr>
<th>Name of product</th>
<th>Dosage form</th>
<th>South Africa registration details</th>
<th>Pack size</th>
<th>Batch Number</th>
<th>Expiry date</th>
</tr>
</thead>
<tbody>
<tr>
<td>Benylin Original</td>
<td>Liquid</td>
<td>G829 (Act 101/1965)</td>
<td>100 ml</td>
<td>297842</td>
<td>05/2017</td>
</tr>
<tr>
<td>Benylin with Codeine</td>
<td>Liquid</td>
<td>G830 (Act 101/1965)</td>
<td>100 ml</td>
<td>297917</td>
<td>05/2017</td>
</tr>
</tbody>
</table>

Reason for recall:
These batches were manufactured for the Mozambique market, but inadvertently delivered to the South African market due to a setup error in our ordering system. As a result the enclosed package inserts in these packs are available in Portuguese language only and therefore do not meet the English/Afrikaans language requirements for the South African market. The information on both cartons and labels however, is reflected in English and Afrikaans.

For ease of reference we have attached images of the products concerned.

Action:
1. Please stop all distribution and/or dispensing and remove any of the product with the above-mentioned batch numbers from stores, storage facilities and shelves immediately.
2. Quarantine and await contact from wholesaler for upliftment and credit process.

This recall is being conducted with the knowledge of the Medicines Control Council. We appreciate your immediate attention and cooperation and sincerely regret any inconvenience this action may cause.

Yours sincerely

CHRISTO FOUCHE
Customer Director

KAREN SLATER
Responsible Pharmacist