

## **Clinical Study Synopsis for Public Disclosure**

This clinical study synopsis is provided in line with **Boehringer Ingelheim's Policy on Transparency and Publication of Clinical Study Data**.


The synopsis - which is part of the clinical study report - had been prepared in accordance with best practice and applicable legal and regulatory requirements at the time of study completion.

The synopsis may include approved and non-approved uses, doses, formulations, treatment regimens and/or age groups; it has not necessarily been submitted to regulatory authorities.

A synopsis is not intended to provide a comprehensive analysis of all data currently available regarding a particular drug. More current information regarding a drug is available in the approved labeling information which may vary from country to country..

Additional information on this study and the drug concerned may be provided upon request based on **Boehringer Ingelheim's Policy on Transparency and Publication of Clinical Study Data**.

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<b>Name of Company:</b> Boehringer Ingelheim		<b>Statement on discontinuation of the study</b>	 <b>Boehringer Ingelheim</b>
<b>BI Proprietary Name:</b> NA		<b>EudraCT No.:</b> 2011-001801-29	
<b>BI Investigational Product:</b> Tiotropium + BI54903		<b>Page:</b> 1	
<b>Report Date:</b> NA	<b>Trial No. / Doc. No.:</b> 1298.3	<b>Dates of Trial:</b> NA	<b>Date of Revision:</b> NA
<p align="center"><b>Proprietary confidential information</b></p> <p>© 2018 <b>Boehringer Ingelheim International GmbH or one or more of its affiliated companies. All rights reserved.</b>  This document may not - in full or in part - be passed on, reproduced, published or otherwise used without prior written permission</p>			
<b>Title of Trial:</b>		A single dose, randomised, placebo-controlled, double-blind, 5-way crossover (employing an incomplete block design), efficacy (including 24-h pulmonary function tests) and safety comparison of Tiotropium/B1 54903 FDC ethanolic inhalation solution via Respimat® (doses of 1.23 µg/363.6 µg, 2.46 µg/363.6 µg or 4.93 µg/363.6 µg) versus free combination of Tiotropium aqueous inhalation solution via Respimat® (doses of 0, 2.5 µg, 5 µg or 10 µg) plus B1 54903 ethanolic inhalation solution via Respimat® (dose of 363.6 µg ) in patients with asthma	
<b>Trial Sites:</b>		NA	
<b>Publications:</b>		NA	
<b>Clinical Phase:</b>		II	
<b>Statement on discontinuation of the study:</b>		Discontinued by <b>Boehringer Ingelheim</b> during preparation of the trial. No patient entered the study, therefore no results / data are available.	