



PMI RESEARCH & DEVELOPMENT

Clinical Study Report Signature Pages

Study Title: A randomized, controlled, open-label, 3-arm parallel group, multi-center study to demonstrate reductions in exposure to selected smoke constituents in apparently healthy smokers switching to the Tobacco Heating System 2.2 Menthol (THS 2.2 Menthol) or observing smoking abstinence, compared to continuing to use menthol conventional cigarettes, for 5 days in confinement and prolonged by 86 days in an ambulatory setting ZRHM-REXA-08-US

Study Number:

Product Name: Tobacco Heating System 2.2 Menthol (THS 2.2 Menthol)

Principal Investigator and Affiliation: Dr William Lewis, Covance Dallas Site
1341 W. Mockingbird Ln., Suite 400E
Dallas, TX 75247
Dr H. Frank Farmer, Covance Daytona Beach Site
1900 Mason Ave., Suite 140
Daytona Beach, FL 32117

Sponsor: Philip Morris Products S.A.
PMI Research & Development
Quai Jeanrenaud 5
2000 Neuchâtel, Switzerland

Sponsor Signatories: Christelle Haziza, PhD, Manager P1 Clinical Program, Clinical Scientist
Guillaume de La Bourdonnaye, MEng, MSc, Biostatistician
Andrea Donelli, Clinical Scientist
Ruben Rosoky, MD PhD MFPM, Medical Safety Officer

Version: 1.0

Date: 25 May 2016

Confidentiality Statement

This document is confidential. Disclosure of any of its contents to third parties is not permitted except by the prior written consent of Philip Morris Products S.A.



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Study Number:

ZRHM-REXA-08-US

I have read the Clinical Study Report referenced above and confirm that to the best of my knowledge it accurately describes the conduct and results of the study.

Signed: W Lewis

Date: 26 May 2016

Dr William Lewis



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Signed: _____

Date: _____

12 Jul 2016

Dr Hugh Coleman (Sub-Investigator)

Dr Hugh Coleman, a sub-Investigator on the Study has reviewed and signed off the Clinical Study Report as the Principal Investigator is no longer with the Company.

**SIGNATURES OF SPONSORS RESPONSIBLE PERSONNEL**

This Clinical Study Report was subject to critical review and has been approved by the Sponsor.


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
We, the undersigned, confirm that to the best of our knowledge this clinical study report accurately describes the experimental methods and results of the study.

Signed: 
Christelle Naziza, PhD
Manager P1 Clinical Program, Clinical Scientist

Date: 30 May 2016

Signed: 
Guillaume de La Bourdonhaye, MEng, MSc
Biostatistician

Date: 31 MAY 2016

Signed: 
Andrea Donelli
Clinical Scientist

Date: 26 May 2016

Signed: _____
Ruben Rosoky, MD PhD MFPM
Medical Safety Officer

Date: _____



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Christelle Haziza, PhD
Manager P1 Clinical Program, Clinical Scientist

Date: _____

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Guillaume de La Bourdonnaye, MEng, MSc
Biostatistician

Date: _____

Signed: _____
Andrea Donelli
Clinical Scientist

Date: _____

Signed: _____
Ruben Rosoky, MD PhD MFPM
Medical Safety Officer

Date: 26 MAY 2016