



Clinical Study Synopsis for Public Disclosure

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
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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|---|---|---|--|---|
| Name of company: Boehringer Ingelheim | | Tabulated Trial Report | |  Boehringer Ingelheim Synopsis No.: |
| Name of finished product: Not applicable | | EudraCT No.: 2009-015676-95 | | |
| Name of active ingredients: Linagliptin, metformin | | Page: 1 of 5 | | |
| Module: | | Volume: | | |
| Report date: 16 AUG 2010 | Trial No. / U No.: 1288.2 / U10-2276-01 | Dates of trial: 08 JAN 2010 – 31 MAR 2010 | Date of revision: Not applicable | |
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| Title of trial: | | Bioequivalence of a 2.5 mg linagliptin / 500 mg metformin fixed dose combination tablet compared with single linagliptin 2.5 mg and metformin 500 mg tablets administered together in healthy male and female volunteers (an open-label, randomised, single-dose, two-way crossover, Phase I trial) | | |
| Principal Investigator: | | [REDACTED] | | |
| Trial site: | | Human Pharmacology Centre of Boehringer Ingelheim, Ingelheim, Germany | | |
| Publication (reference): | | Data of this trial have not been published. | | |
| Clinical phase: | | I | | |
| Objectives: | | To demonstrate bioequivalence of a 2.5 mg linagliptin / 500 mg metformin fixed dose combination (FDC) tablet compared to single tablets of linagliptin 2.5 mg and metformin 500 mg administered together | | |
| Methodology: | | Open-label, randomised, single-dose, 2-way crossover design | | |
| No. of subjects: | | <p>planned: entered: 96</p> <p>actual: entered: 95</p> <p>Treatment A (FDC tablet): treated: 94 analysed (for primary endpoints): 94</p> <p>Treatment B (single tablets): treated: 95 analysed (for primary endpoints): 95</p> | | |
| Diagnosis and main criteria for inclusion: | | Healthy volunteers, male and female, age 21 to 50 years, body mass index (BMI) range: 18.5 to 29.9 kg/m ² | | |
| Test product: | | Linagliptin/metformin FDC tablet | | |
| dose: | | 2.5 mg linagliptin and 500 mg metformin | | |
| mode of admin.: | | Peroral with 240 mL water in a standing position after an overnight fast of at least 10 h | | |
| batch no.: | | 902832 | | |

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| Reference therapy: | Linagliptin tablet and metformin (Glucophage®) tablet | | | |
| dose: | 2.5 mg linagliptin and 500 mg metformin | | | |
| mode of admin.: | Peroral with 240 mL water in a standing position after an overnight fast of at least 10 h | | | |
| batch no.: | Linagliptin: B081004241, metformin: 250045 (Merck Pharma GmbH) | | | |
| Duration of treatment: | Single dose in each treatment period separated by a wash-out phase of at least 35 days | | | |
| Criteria for evaluation: | | | | |
| Clinical pharmacology: | Primary endpoints: AUC ₀₋₇₂ and C _{max} for linagliptin; AUC _{0-∞} and C _{max} for metformin Secondary endpoints: AUC _{0-∞} for linagliptin; AUC _{0-tz} , %AUC _{tz-∞} , AUC _{t1-t2} , t _{max} , λ _z , t _{1/2} , MRT _{po} , CL/F, V _z /F for both analytes | | | |
| Safety: | Physical examination, vital signs (blood pressure, pulse rate), 12-lead electrocardiogram (ECG), laboratory tests, adverse events (AEs), tolerability assessment | | | |
| Statistical methods: | | | | |
| Primary endpoints and key secondary endpoints: point estimators (geometric means [gMean]) of the median intra-subject ratios and their 2-sided 90% confidence intervals (CIs). Statistical model: analysis of variance (ANOVA) on log-transformed parameters including effects for 'sequence', 'subjects nested within sequences', 'period' and 'treatment'. Confidence intervals were based on the residual error from ANOVA. Other parameters: descriptive statistics and tabulated frequencies. | | | | |

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SUMMARY – CONCLUSIONS:

Clinical pharmacology results:


The study population consisted of 95 healthy volunteers, 48 male and 47 female. All subjects were White and aged between 21 and 50 years (mean: 36.5 years, standard deviation [SD]: 8.2 years) with a BMI between 19.3 and 29.7 kg/m² (mean: 24.57 kg/m², SD: 2.66 kg/m²). No relevant medical history or baseline conditions were reported for any of the participating subjects. Of the 95 entered subjects, 94 completed both treatment periods. One subject completed the first treatment period (single tablets) but withdrew her consent before the start of the second treatment period (FDC). There was no important protocol violation reported in this trial.

For both linagliptin and metformin, gMean plasma concentration-time profiles were similar for the FDC and single tablet treatments.

Geometric mean AUC₀₋₇₂ of linagliptin was 181 nmol·h/L for both the FDC and the single tablets (the inter-subject geometric coefficient of variation [gCV] was 27.6% and 26.4%, respectively). Geometric mean C_{max} of linagliptin was 5.33 nmol/L (gCV 27.3%) for the FDC and 5.45 nmol/L (gCV 26.9%) for the single tablets. Median t_{max} was 3.00 h for both the FDC and the single tablets.

Geometric mean AUC_{0-∞} of metformin was 7430 ng·h/mL (gCV 23.7%) for the FDC and 7480 ng·h/mL (gCV 24.5%) for the single tablets. Geometric mean AUC_{0-tz} of metformin was 7320 ng·h/mL (gCV 24.3%) for the FDC and 7360 ng·h/mL (gCV 25.9%) for the single tablets. Geometric mean C_{max} of metformin was 1130 ng/mL (gCV 27.4%) for the FDC and 1160 ng/mL (gCV 28.4%) for the single tablets. Median t_{max} was 2.02 h for the FDC and 3.00 h for the single tablets.


The adjusted gMean ratios (FDC to single tablets), 90% CIs, and intra-subject gCVs of AUC₀₋₇₂ (linagliptin only), AUC_{0-∞} and AUC_{0-tz} (metformin only), and C_{max} (both analytes) are summarised in Table 1.

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| Clinical pharmacology results (continued): | Table 1: | Adjusted gMean ratio, 90% confidence interval and intra-subject gCV for the key parameters of linagliptin and metformin | | | |
| | | Adjusted gMean ratio (FDC/single tablets ¹) [%] | 2-sided 90% confidence interval | | Intra-individual gCV [%] |
| | | | Lower limit [%] | Upper limit [%] | |
| | Linagliptin 2.5 mg | | | | |
| | AUC ₀₋₇₂ | 99.9 | 96.6 | 103.3 | 13.9 |
| | C _{max} | 98.1 | 94.4 | 101.9 | 16.0 |
| | Metformin 500 mg | | | | |
| | AUC _{0-∞} | 99.1 | 96.4 | 102.0 | 11.6 |
| | AUC _{0-tz} | 99.4 | 96.5 | 102.3 | 12.3 |
| | C _{max} | 97.9 | 94.4 | 101.5 | 14.9 |
| | ¹ FDC N=94, single tablets N=95 | | | | |
| | For both linagliptin and metformin, all 90% CIs for AUC and C _{max} were contained in the bioequivalence acceptance range of 80 to 125%. Therefore, bioequivalence of the FDC compared to the single tablets can be concluded. | | | | |
| Safety results: | Ninety-four subjects received a total dose of 5 mg linagliptin and a total dose of 1000 mg metformin during the trial as planned. The subject who withdrew after the first treatment period received 2.5 mg linagliptin and 500 mg metformin as single tablets only. | | | | |
| | Forty subjects (42.1%) reported at least 1 AE during the 2 treatment periods. Twenty-seven subjects (28.4%) experienced AEs during the treatment period with single tablets of linagliptin and metformin and 20 subjects (21.3%) experienced AEs during the treatment period with the FDC. The most frequently reported AEs by system organ class were nervous system disorders (20 subjects, 21.1%), infections and infestations (16 subjects, 16.8%), and gastrointestinal disorders (10 subjects, 10.5%). On the preferred term level, the most frequently reported AEs were headache (20 subjects, 21.1%), nasopharyngitis (13 subjects, 13.7%), and diarrhoea (6 subjects, 6.3%). | | | | |
| | All AEs were of mild or moderate intensity, no AE was of severe intensity. No AE was drug-related as assessed by the investigator. All reported AE episodes | | | | |

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| <p>had resolved by the end of the trial with the exception of nasopharyngitis in 1 subject. No subject was discontinued due to an AE. There was no serious AE (SAE) reported in this trial.</p> <p>Clinical laboratory tests, vital signs and ECG recordings revealed no safety concerns in this study. Global tolerability was assessed as good for all exposed subjects in both treatment periods.</p> <p>Conclusions: The fixed dose combination tablet of linagliptin 2.5 mg and metformin 500 mg was bioequivalent to single tablets of 2.5 mg linagliptin and 500 mg metformin administered together. Both the fixed dose combination and combined administration as single tablets were well tolerated and safe in healthy male and female volunteers.</p> | | | | |