# Non-interventional Study Protocol

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<tr>
<td>BI Study Number:</td>
<td>1237.43</td>
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<tr>
<td>BI Investigational Product(s):</td>
<td>Spiolto® Respimat®</td>
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<tr>
<td>Title:</td>
<td>Assessment of physical functioning and handling of Spiolto® Respimat® in patients with chronic obstructive pulmonary disease (COPD) requiring long-acting dual bronchodilation in routine clinical practice.</td>
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<td>Protocol version identifier:</td>
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<td>Date of last version of protocol:</td>
<td>13 July 2016</td>
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<tr>
<td>PASS:</td>
<td>No</td>
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</table>
| EU PAS register number: | Not yet available  
(study will be registered in EU PAS Register (ENCePP) and clinicaltrials.gov) |
| Active substance: | R03AL06  
Tiotropium bromide + Olodaterol |
| Medicinal product: | Spiolto® Respimat® 2.5 microgram/2.5 microgram, inhalation solution; tiotropium/olodaterol |
| Product reference: | NL/H/3157/001/DC |
| Procedure number: | n.a. |
| Marketing authorisation holder(s): | MAH:  
Boehringer Ingelheim International GmbH  
Binger Straße 173  
55216 Ingelheim am Rhein  

This study is initiated, managed and sponsored by: |
| Joint PASS: | No |
### Research question and objectives:
The primary objective of this NIS study is to measure changes in physical functioning - serving as a surrogate for physical activity and exercise capacity - in COPD patients being treated with Spiolto® Respimat® after approximately 6 weeks.

The secondary objective is to evaluate the patient’s general condition (physician’s evaluation) at visit 1 (= baseline visit at the start of the study) and at visit 2 (= final visit approx. 6 weeks after visit 1), as well as patient satisfaction with Spiolto® Respimat® at visit 2.

### Country(-ies) of study:
Italy

### Author:
Tel:  
Fax:  
Mobile:

### Marketing authorisation holder(s):
Boehringer Ingelheim International GmbH  
Binger Str. 173  
D-55216 Ingelheim am Rhein  
Germany

### In case of PASS, add:
**MAH contact person:** Not applicable

**In case of PASS, add:**  
**<EU-QPPV>:** Not applicable

**In case of PASS, add:**  
**<Signature of EU-QPPV>:** Not applicable

### Date:
13 July 2016
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2. LIST OF ABBREVIATIONS

ADR  Adverse Drug Reaction
AE  Adverse Event
AESI  Adverse Event of Special Interest
CML  Local Clinical Monitor
COPD  Chronic Obstructive Pulmonary Disease
CRA  Clinical Research Associate
eCRF  Electronic Case Report Form
DMP  Data Management Plan
EU  European Union
FDC  Fix Dose Combination
FEV1  Forced expiratory volume in one second
GCP  Good Clinical Practice
GEP  Good Epidemiological Practice
GPP  Good Pharmacoepidemiology Practice
GOLD  Global Initiative for Chronic Obstructive Lung Disease
ICH  International Conference on Harmonisation
ICS  Inhalative Corticosteroids
IEC  Independent Ethics Committee
ISF  Investigator Site File
LABA  Long-acting beta2 adrenoceptor agonist
LAMA  Long-acting muscarinic antagonist
mMRC  Modified Medical Research Council Scale
NIS  Non-Interventional Study
PF-10  Physical Functioning patient questionnaire
PGE  Physician’s Global Evaluation
SAE  Serious Adverse Event
SAP  Statistical Analysis Plan
SF-36  36-Item Short Form Health Survey
SGRQ  St George’s Respiratory Questionnaire
SmPC  Summary of Product Characteristics
### 3. RESPONSIBLE PARTIES

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<td>Trial Data Manager (TDM)</td>
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4. ABSTRACT

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<thead>
<tr>
<th>Name of company:</th>
<th>Boehringer Ingelheim</th>
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<tr>
<td>Name of finished medicinal product:</td>
<td>Spiolto® Respimat®.</td>
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<tr>
<td>Name of active ingredient:</td>
<td>R03AL06 Tiotropium bromide + Olodaterol</td>
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<tr>
<td>Protocol date:</td>
<td>13 July 2016</td>
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<tr>
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<tr>
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<td>1.0</td>
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<tr>
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<td>Title of study:</td>
<td>Assessment of physical functioning and handling of Spiolto® Respimat® in patients with chronic obstructive pulmonary disease (COPD) requiring long-acting dual bronchodilation in routine clinical practice.</td>
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<tr>
<td>Rationale and background:</td>
<td>Reduced physical activity resulting in deconditioning and restricted physical functioning is a common constraint of patients with moderate to very severe COPD. Clinical studies investigating treatment with Spiolto® Respimat® and its single components have shown significant improvements in exercise capacity in patients with COPD. Real-world data on the effects of a fixed-dose combination (LABA+LAMA) therapy with tiotropium and olodaterol administered in a single device, in COPD patients who need treatment with two long-acting bronchodilators, is not available.</td>
</tr>
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<td>Research question and objectives:</td>
<td>The primary objective of the study is to measure changes in physical functioning - serving as a surrogate for physical activity and exercise capacity - in COPD patients being treated with Spiolto® Respimat® after approximately 6 weeks. The secondary objective is to evaluate the patient’s general condition (physician’s evaluation) at visit 1 (= baseline visit at the start of the study) and at visit 2 (= final visit approx. 6 weeks after visit 1), as well as patient satisfaction with Spiolto® Respimat® at visit 2.</td>
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<td>Study design:</td>
<td>Open-label observational study, including COPD patients receiving treatment with Spiolto® Respimat® for approximately 6 weeks, which is the average time between two medical consultations.</td>
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<td>Population:</td>
<td>COPD patients requiring a fixed combination therapy of two long-acting bronchodilators (LAMA + LABA) according to approved SmPC and GOLD guidelines.</td>
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<td>Name of company:</td>
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<td>Variables:</td>
<td>- Patient demographics (age, gender, height &amp; weight)</td>
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<td>- Smoking status (current smokers, former smokers, and never smokers) and pack-years</td>
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<td></td>
<td>- Concomitant diseases / Comorbidities such as cardiovascular disease, diabetes mellitus, musculoskeletal impairment, renal diseases, liver diseases, osteoporosis, gastroesophageal reflux (GERD), or lung cancer - COPD related and other relevant concomitant medication such as beta-blockers, beta-agonists, corticosteroids, or proton pump inhibitors</td>
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<td>- Reported exacerbations based on medical history in the last 12 months and exacerbations leading to hospitalization in the last 12 months</td>
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<td>- Assessment of the severity of breathlessness based on the Modified Medical Research Council Questionnaire (mMRC)</td>
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<td>- Physical Functioning based on PF-10 scores (sub-domain of SF-36)</td>
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<td>- Patient satisfaction with Spiolto® Respimat® assessing the overall satisfaction with Spiolto® Respimat®, the satisfaction of inhalation, as well as the device handling satisfaction</td>
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<td>- General condition of patient based on Physician’s Global Evaluation (PGE) assessing the general condition of the patient at visit 1 and at visit 2</td>
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<td>- Safety Reporting; Adverse Drug Reactions (serious and non-serious), fatal AEs, and pregnancies</td>
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<td></td>
<td>- GOLD spirometric classifications (1, 2, 3, 4) and GOLD patient groups (A, B, C, D) based on GOLD guidelines at visit 1</td>
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Name of company: Boehringer Ingelheim

Name of finished medicinal product: Spiolto® Respimat®.

Name of active ingredient: R03AL06 Tiotropium bromide + Olodaterol

Protocol date: 13 July 2016

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<td>1.0</td>
<td>n.a.</td>
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Data sources:
- To be completed by the physician:
  - Patient demographics
  - Patient medical files
  - Physician’s Global Evaluation (PGE) at visit 1 and visit 2
- To be completed by the patient at visit 1:
  - mMRC breathlessness scale
- To be completed by the patient at visit 1 and at visit 2:
  - Physical Functioning Questionnaire (PF-10)
- To be completed by the patient at visit 2 only:
  - Patient satisfaction survey

Study size: It is planned that data of approximately 400 patients in 40 Italian centres will be collected

Data analysis:

Primary outcomes:
“Therapeutic success” at visit 2 (10-point increase in the PF-10 score between visit 1 and visit 2).

Secondary outcomes:
- Changes in the PF-10 score from visit 1 to visit 2
- General condition of the patient, evaluated by the physician (PGE score) at visit 1 and visit 2.
- Patient satisfaction with Spiolto® Respimat® at visit 2.

Milestones:
- Planned start of data collection: January-2017
- Planned end of data collection: January-2018
- Planned final study report: April-2018
5. AMENDMENTS AND UPDATES

None.
6. **MILESTONES**

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<td>Start of data collection</td>
<td>31 January 2017</td>
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<tr>
<td>End of data collection</td>
<td>31 January 2018</td>
</tr>
<tr>
<td>Final report of study results:</td>
<td>30 April 2018</td>
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7. RATIONALE AND BACKGROUND

7.1 MEDICAL BACKGROUND

COPD is defined as a preventable and treatable disease of the airways, with significant systemic consequences. Inactivity is believed to be crucial to the development of the extrapulmonary effects of the disease like skeletal muscle weakness, osteoporosis and cardiovascular disease. Recent data suggest that patients suffering from COPD with low levels of physical activity have increased risk for hospital admission and have significantly enhanced mortality. Epidemiological data suggest that this may directly or indirectly lead to more rapid decline in lung function [R13-3633]. Physical activity is reduced early in the disease progression, as of GOLD Stage 2 [R13-3633]. More recent evidence from large placebo controlled clinical trials indicates that COPD patients are experiencing a steeper absolute decline in lung function with GOLD 2 airflow limitation than with GOLD 3 and 4 [R15-5015]. All these observations suggest the importance of early optimal treatment of the disease. Long-acting bronchodilators, such as long-acting muscarinic antagonists (LAMAs), are the cornerstone of maintenance therapy for patients with moderate to very severe chronic obstructive pulmonary disease (COPD) whose symptoms are not adequately controlled by short-acting bronchodilators alone [P13-05794, P14-01052].

An option recommended by GOLD guideline for patients not adequately controlled on a single long-acting bronchodilator is to combine a LAMA with a long-acting β2-agonist (LABA) [P14-01052]. This has prompted the development of combining LAMA+LABA as fixed-dose combinations [P13-05794]. The rationale for combining bronchodilators with different mechanisms is based on the notion of additive relaxation of airway smooth muscle by direct inhibition of cholinergic activity and functional antagonism of bronchoconstriction through β2-adrenergic pathways, with the expectation of an increase in the degree of bronchodilation for equivalent or lesser side effects. Several recent studies have provided evidence in support of this increased bronchodilatory effect when LABAs are added to LAMAs.

When long acting beta-agonists (LABA) and long acting muscarinic antagonists (LAMA) with similar or equivalent posologies are combined, the opportunity exists for offering a simpler and more convenient administration regimen with the development of fixed combinations within the same inhaler device. Fixed-dose combinations of a short-acting β2--agonist and a short-acting anticholinergic have been developed and have been shown to be safe, efficacious and convenient for the patient (e.g., Berodual®: Fenoterolhydrobromid + Ipratropiumbromid, Combivent®: salbutamol + ipratropium bromide; [P94-1346]). Olodaterol is a highly selective and nearly full β2 agonist [P10-07776, P11-07720] that provides 24-h bronchodilation in patients with COPD [P13-11467, P13-14112, P13-11346, P13-11345]. Olodaterol is also associated with symptomatic benefit [P13-11341] and enhanced exercise capacity [P13-14109].

The complementary modes of action of tiotropium and olodaterol have previously been demonstrated in animal models, phase II clinical trials and during the Phase IIIa programs [P10-09337, P14-12073, P10-14042, P13-02357]. In the Phase III program the additional benefits of the tiotropium + olodaterol fixed-dose combination (FDC) over its mono-components has been assessed on lung function, quality of life (St. George's Respiratory Questionnaire -SGRQ), dyspnea...
(Transition Dyspnea Index-TDI) and exercise endurance time. Another clinically important potential benefit of the tiotropium + olodaterol FDC over the mono components, the impact on exacerbations of COPD, is currently evaluated in a Phase IIIb trial.

7.2 DRUG PROFILE

Tiotropium + olodaterol FDC is an aqueous solution of tiotropium and olodaterol contained in a cartridge. It is administered by using the Respimat® Inhaler. One cartridge is used per inhaler, which is inserted into the device prior to first use. In pivotal clinical trials and for the intended marketed product, the clinical dose consists of two puffs once daily. The Respimat® Inhaler uses mechanical energy to create a soft mist which is released over a period of approximately 1.5 seconds.

Tiotropium + olodaterol FDC was shown to be safe and well tolerated over 1 year in a moderate to very severe COPD population. The overall incidences of adverse events (AEs), serious adverse event (SAEs), fatal AEs, frequencies for cardiac events and major adverse cardiovascular event in the tiotropium + olodaterol FDC treatment group were similar to the mono-components. The nature and frequency of AEs in general was consistent with the disease under study. There were no results in the clinical development program suggesting the need for absolute contraindications for the combination product [P15-03349].

In conclusion, the clinical trials conducted to date have shown tiotropium + olodaterol FDC to be a safe, well tolerated and efficacious combination therapy according to treatment guidelines in a moderate to very severe COPD patient population [P15-04531, P15-03349]. The observed incremental bronchodilator response for the combination compared to the individual components translated into benefits that were meaningful to the patient, with improvements in several patient centered outcomes. For further information please refer to the SmPC of Spiolto® Respimat®.
8. **RESEARCH QUESTION AND OBJECTIVES**

8.1 **RATIONALE FOR PERFORMING THE STUDY**

The contribution of physical inactivity to disability in COPD is difficult to distinguish from disease progression; however, it is clear that physical activity is significantly lower in patients with COPD than in healthy controls [R13-3633]. COPD prevents patients from carrying out daily activities due to exercise intolerance, which is often attributed to limited pulmonary ventilation. Physical inactivity may be related to avoidance of exertion as a result of fear of dyspnea. Furthermore, physical inactivity has been associated with skeletal muscle weakness and exercise intolerance [R15-4559, R15-4561].

The loss of physical activity in COPD is also associated with increased mortality. Data from a study of 2386 patients with COPD demonstrated that, following adjustment for relevant confounders, subjects who reported low, moderate or high physical activity had a significantly lower risk of all-cause mortality than those with very low physical activity (p = 0.001) [R15-4564]. Clinical studies of both Spiriva® [P13-04267, P05-09483, P13-14109] and Striverdi® Respimat® in COPD patients have demonstrated significant improvement in exercise capacity [P13-14109].

The benefits of tiotropium + olodaterol FDC have been studied in controlled Phase III programs on exercise endurance, however, data regarding physical activity when treated with Spiolto® Respimat® is not available from a real life setting.

8.2 **STUDY OBJECTIVES**

The primary objective of the study is to measure changes in physical functioning - serving as a surrogate for physical activity and exercise capacity - in COPD patients being treated with Spiolto® Respimat® after approximately 6 weeks in routine clinical practice.

The secondary objectives are to evaluate the patient’s general condition (physician’s evaluation) at visit 1 and at visit 2, as well as patient satisfaction with Spiolto® Respimat® at visit 2.
9. RESEARCH METHODS

9.1 STUDY DESIGN

This is a self-controlled study design enrolling consented COPD patients who will be treated with Spiolto® Respimat® according to the approved SmPC. Patients will be enrolled consecutively and will be followed over an observational period of approx. 6 weeks. Data as listed in table 9.1:1 will be collected.

Table 9.1:1: Visit flow chart and data collection parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Visit 1; baseline visit</th>
<th>Visit 2; approx. 6 weeks after baseline visit</th>
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<tbody>
<tr>
<td>Informed Consent</td>
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<td></td>
</tr>
<tr>
<td>Inclusion / Exclusion Criteria</td>
<td>X</td>
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<tr>
<td>Patient demographics (age, gender, height, and weight)</td>
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<td></td>
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<tr>
<td>Start date of COPD</td>
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<tr>
<td>Number of exacerbations in the last 12 months</td>
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</tr>
<tr>
<td>Number of exacerbations leading to hospitalization in the last 12 months</td>
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<tr>
<td>mMRC breathlessness scale, completed by the patient</td>
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<td>Past COPD therapies (6 months before visit 1)</td>
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<td>Respimat® training (yes/no)</td>
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<td>COPD severity based on GOLD assessment¹</td>
<td>X</td>
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<td>Smoking status/history</td>
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<td>Concomitant diseases / Comorbidities</td>
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<tr>
<td>COPD related and other relevant concomitant medication</td>
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<td>X</td>
</tr>
<tr>
<td>Physical functioning (PF-10) questionnaire, completed by the patient</td>
<td>X</td>
<td>X</td>
</tr>
<tr>
<td>General condition of patient evaluated by Physician’s Global Evaluation (PGE)</td>
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<td>X</td>
</tr>
<tr>
<td>Safety: Adverse Drug Reactions (serious and non-serious), fatal AE, pregnancy</td>
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<td>X</td>
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<tr>
<td>Patient satisfaction with Spiolto® Respimat®, completed by the patient</td>
<td>X</td>
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</tr>
<tr>
<td>Rational for Spiolto® Respimat® treatment discontinuation (if applicable)</td>
<td>X</td>
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</tr>
<tr>
<td>Continuation or discontinuation of treatment with Spiolto® Respimat® after the study (yes/no)</td>
<td>X</td>
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</table>

¹ GOLD assessment: Global Initiative for Chronic Obstructive Lung Disease.
GOLD patient group (A, B, C or D) will be automatically calculated within the eCRF based on available exacerbation history, mMRC, and GOLD spirometric classification of airflow limitation based on post-bronchodilator FEV₁ if available.

Baseline characteristics of eligible patients who gave Informed Consent, but were not treated in the study will also be collected.

9.1.1 Outcomes

Primary outcomes:
“Therapeutic success” at visit 2 (10-point increase in the PF-10 score between visit 1 and visit 2).

Secondary outcomes:
- Changes in the PF-10 score from visit 1 to visit 2
- General condition of the patient, evaluated by the physician (PGE score) at visit 1 and visit 2.
- Patient satisfaction with Spiolto® Respinmat® at visit 2.

9.2 SETTING

It is planned that data of approximately 400 patients from approximately 40 sites in Italy will be collected. Site selection will be performed to reflect routine COPD care in order to secure representativeness of the COPD population.

A log of all patients included in the study (i.e. who have given informed consent) will be maintained in the Investigator Site File (ISF) irrespective of whether they have been treated or not.

Boehringer Ingelheim reserves the right to discontinue the study or to remove a patient at a particular study site for the following reasons:
1. Failure to meet expected enrolment goals overall or at a particular study site.
2. Emergence of any efficacy/safety information that could significantly affect continuation of the study, or any other administrative reasons, i.e. lack of recruitment.
3. Violation of the protocol, the contract, or applicable laws and regulations for non-interventional studies, which could disturb the appropriate conduct of the NIS.

9.2.1 Study sites

It is planned that data of approximately 400 patients from approximately 40 sites in Italy will be collected. Site selection will be performed to reflect routine COPD care in order to secure representativeness of the COPD population.

9.2.2 Study population

- Inclusion Criteria:
  Patients can be included if all of the following criteria are met:
  1. Written informed consent prior to participation
  2. Female and male patients ≥ 40 years of age
3. Patients diagnosed with COPD and requiring long-acting dual bronchodilation (LAMA + LABA) treatment according to approved Spiolto® Respimat® SmPC.

- **Exclusion Criteria:**
  1. Patients with contraindications according to Spiolto® Respimat® SmPC
  2. Patients who have been treated with a LABA/LAMA combination (free and fixed dose) in the previous 6 months
  3. Patients continuing LABA-ICS treatment should not be additionally treated with Spiolto® Respimat® in order to avoid a double dosing of long-acting beta-agonists
  4. Patients for whom further follow-up is not possible at the enrolling site during the planned study period of approx. 6 weeks
  5. Pregnancy and lactation
  6. Patients currently listed for lung transplantation
  7. Current participation in any clinical trial or any other non-interventional study of a drug or device.

### 9.3 VARIABLES

The following parameters will be collected and assessed at visit 1 and/or visit 2:

- Patient demographics (age, gender, height & weight)
- Smoking status (current smokers, former smokers, and never smokers) and pack-years
- Concomitant diseases / Comorbidities such as cardiovascular disease, diabetes mellitus, musculoskeletal impairment, renal diseases, liver diseases, osteoporosis, gastroesophageal reflux (GERD), or lung cancer
- COPD related and other relevant concomitant medication such as beta-blockers, beta-agonists, corticosteroids, or proton pump inhibitors
- Reported exacerbations based on medical history in the last 12 months and exacerbations leading to hospitalization in the last 12 months
- Assessment of the severity of breathlessness based on the Modified Medical Research Council Questionnaire (mMRC)
- Physical Functioning based on PF-10 scores*
- Patient satisfaction with Spiolto® Respimat® assessing the overall satisfaction with Spiolto® Respimat®, the inhalation satisfaction as well as the device handling
- General condition of patient based on Physician’s Global Evaluation (PGE) assessing the general condition of the patient at the beginning and at the end of the study
- Safety Reporting; Adverse Drug Reactions (serious and non-serious), fatal AEs, and pregnancies at the beginning and at the end of the study
- GOLD spirometric classifications (1, 2, 3, 4) and GOLD patient groups (A, B, C, D) based on GOLD guidelines

*The PF-10 is a sub-domain of the SF-36 and consists of 10 questions evaluating the extent of experienced restrictions while conducting usual activities. Each
question of the PF-10 can be answered with “yes, limited a lot”, “yes, limited a little”, or “No, not limited at all”, with a score of 1, 2, or 3. The scores over the 10 questions will be summed, resulting in a value between 10 (a patient answering all questions with “yes, limited a lot”) and 30 (a patient answering all questions with “No, not limited at all”). The final sum of the individual scores will be standardized to a range of 0 to 100 using the following formula: 

\[ 100 \times \frac{\text{sum} - 10}{20} \]

9.4 DATA SOURCES

Medical records collected through routine clinical care will be used to assess the inclusion/exclusion criteria of patients. Such medical records will be used for patient demographics, smoking history, collection of previous COPD medication, concomitant diseases, and concomitant medication. All patients will be enrolled consecutively. The treating physician will use the Physician’s Global Evaluation (PGE) to evaluate the general condition of the patient on an 8-point ordinal scale from 1 (very poor) to 8 (excellent). PGE will be completed before and approx. 6 weeks after treatment initiation.

The modified Medical Research Council (mMRC) scale will be used to assess the breathlessness state of the patient before the treatment. The mMRC stage (0 to 4) collected from the patient, as well as the exacerbation history and the post-bronchodilator FEV1, will be used to automatically calculate the GOLD stage and patient group in the eCRF.

The physical functioning (PF-10) questionnaire is a subscale of the validated 36-Item Short Form Health Survey and contains 10 questions about everyday physical activity and functioning. Patients will be asked to complete the PF-10, in order to evaluate their physical functioning before and after treatment with Spiolto® Respimat®. A patient satisfaction survey will also be completed at the end of the study, using a 7-point ordinal scale with divisions from very dissatisfied to very satisfy.

9.5 STUDY SIZE

In a previous study, 205.426 with over 1000 patients treated with Spiriva® Respimat®, therapeutic success (i.e., 10-point increase in the PF-10 score between visit 1 and visit 2) was achieved in 61% of the patients. In this study, a lower therapeutic success rate is expected as patients may be already on maintenance treatment at baseline. Assuming a 50% therapeutic success rate and 372 patients, the 95% confidence interval for the therapeutic success rate would be between 44.9% (lower limit) and 55.1% (upper limit). To account for a 7% drop-out rate, the sample size becomes 400 patients.

9.6 DATA MANAGEMENT

A data management plan (DMP) will be created to describe all functions, processes, and specifications for data collection, cleaning and validation. The electronic Case Report Forms (eCRFs) will include programmable edits to obtain immediate feedback if data are missing (also negative answers, unknown), out of range, illogical or potentially erroneous. These rules may encompass simple checks such as range
validation or presence/absence of data. Concurrent manual data review may be performed based on parameters dictated by the DMP. Ad hoc queries to the sites may be generated and followed up for resolution. A source data quality audit may be initiated to ensure that the data in the database is accurate. Source data verification (SDV) will be performed at sites identified by a risk-based approach as needed. The database will be housed in a physically and logically secure computer system maintained in accordance with a written security policy. The system will meet the standards of the International Committee on Harmonization guideline E6R1 regarding electronic study data handling. Patient confidentiality will be strictly maintained.

9.7 DATA ANALYSIS

9.7.1 Statistical design – Model

Details will be described in the statistical epidemiological analysis plan (SEAP).

9.7.2 Further analysis

No formal hypothesis testing will be performed since this is a self-controlled study.

9.7.3 Planned analyses

All patients who have received at least one dose of Spiolto® Respimat® will be included in the analysis; this is the treated set. All analyses will be performed on the treated set (as-treated analysis). If patients have missing values for an outcome, those patients will be excluded for that outcome’s analysis.

The assessment will be carried out using SAS® software. The statistical characteristics presented in the end-of-text tables will be N / mean / SD / min / median / max for continuous variables. Tabulations of relative and absolute frequencies will be presented for categorical variables. Incidence rates and 95% CI will be given when appropriate.

The analyses will relate to the following data as recorded in the eCRF:

- Patient demographics (age, gender, height, and weight)
- Comorbidities
- COPD related and other concomitant medication
- History of smoking
- Reported exacerbations
- Breathlessness based on mMRC score
- Physical Functioning based on PF-10 scores (therapeutic success at visit 2); primary outcome
- Changes from visit 1 to visit 2 in the PF-10 score; secondary outcome
- Patient satisfaction with Spiolto® Respimat® at visit 2 only; secondary outcome
- General condition of the patient: evaluated by the physician (Physician’s Global Evaluation (PGE)); secondary outcome
- Adverse Drug Reactions (ADR & SADR), fatal AEs, pregnancies
- GOLD spirometric classification (1,2, 3, 4)
- GOLD patient groups (A, B, C, D)
- Details of treatment with inhaled respiratory agents before the study
- Details of treatment with respiratory agents during the study
- Reasons for ending treatment during the observation period
- Details of treatment continuation / discontinuation

Main analysis
For the primary outcome, the proportion of patients with therapeutic success will be presented together with the 95% confidence interval.

Further analyses
The patient’s general condition (PGE) at visit 1 and visit 2, mMRC at visit 1 and patient satisfaction at visit 2 are categorical variables so they will be analyzed as tabulations of frequencies. Change from visit 1 to visit 2 in the PF-10 score is a continuous outcome, so it will be analyzed with N / mean / SD / min / median / max.

The safety data will be reported according to local requirements.
As similar studies will also be performed in other European countries, data pooling might be considered at the end.

9.7.4 Handling of missing data
If less than half of the PF-10 questions are missing for a patient, the missing values will be replaced with the mean of the other values and the PF-10 score will be calculated. If half or more than half of the PF-10 questions are missing, no score will be calculated and the PF-10 score will be marked as missing. No other missing data will be imputed. Every effort will be made to collect complete data at the specified time points. Any removal from the analysis will be documented, stating the site and patient number as well as the reason for removal.

9.8 QUALITY CONTROL
To improve and secure data quality, automatic data checks upon data entry will be done within the eCRF. In the eCRF, plausible ranges of values for numeric data entries as well as logical data entries and listings will be provided for each entry field.
Based on this, checks on completeness and plausibility will be performed upon data entry in the eCRF. Validity of data entry thus is ensured by integrated validation checks performed by the system, indicating missing or implausible entries to the document list or investigator. All corrections will be visible from the systems audit trail. No regular source data verification is planned in this study. However, in case of decreasing compliance (i.e. of missing data, data discrepancies, protocol violations, etc.) a for-cause audit or risk-based monitoring visit will be performed.

9.9 LIMITATIONS OF THE RESEARCH METHODS

The intention of this NIS is to collect new data on the physical functioning and exercise capacity of COPD patients on treatment with Spiolto® Respimat® in a real life setting. A NIS is the most suitable instrument for obtaining information about the use of medicines in everyday therapeutic practice and thus for investigating prospectively questions in everyday therapeutic practice. Consecutive enrolment will be employed to minimize selection bias. The entry criteria are non-restrictive which will permit the enrolment of a broad patient population. The choice of treatment is at the discretion of the investigator. Selection bias could occur at the site level and the patient level. To minimize the site level selection bias, the goal is to have participating centers that have access to all available treatment options which are approved for use in that country for the targeted COPD patients. To minimize selection bias at the patient level, consecutive enrolment is performed. Information bias will be minimized by the use of standard eCRF, questionnaire and physicians’ training on the study protocol. An additional limitation is that the physical functioning questionnaire (PF-10) is assessing 10 items of the full 36-Item Short Form Health Survey (SF-36). The 7-item satisfaction scale, which is to be completed by the patient in order to measure satisfaction with Spiolto® Respimat® use, is a self-designed Boehringer-Ingelheim scale, without a public source or validation status.

9.10 OTHER ASPECTS

9.10.1 Informed Consent, Data Protection, Study Records

The study will be carried out in compliance with the protocol, the principles laid down in the Declaration of Helsinki, and as close as possible to the standards of the International Conference of Harmonisation (ICH) Tripartite Guideline for Good Clinical Practice (GCP), Guidelines for Good Epidemiological Practice (GEP) [R10-4560], Good Pharmacoepidemiology Practice (GPP) [R09-0182] and relevant BI Standard Operating Procedures (SOPs). Standard medical care (prophylactic, diagnostic and therapeutic procedures) remains in the responsibility of the treating physician. The investigator should inform the sponsor immediately of any urgent safety measures taken to protect the study subjects against any immediate hazard, and also of any serious breaches of the protocol.
The rights of the investigator and of the sponsor with regard to publication of the results of this study are described in the investigator contract. As a general rule, no study results should be published prior to finalization of the Study Report.

Insurance Cover: The requirements for insurance depend on local law and legislations in the participating country. If required, the terms and conditions of the insurance cover are made available to the investigator and the patients, and the documentation must be archived in the Investigator Site File (ISF).

9.10.1.1 Study Approval, Patient Information, and Informed Consent

This study will be initiated only after all required legal documentation has been reviewed and approved by the respective Independent Ethics Committee (IEC) and competent authority (CA) according to national and international regulations. The same applies for the implementation of changes introduced by amendments.

Prior to patient participation in the study, written informed consent must be obtained from each patient (or the patient’s legally accepted representative). Each signature must be personally dated by each signatory and the informed consent and any additional patient-information form retained by the investigator as part of the study records. A signed copy of the informed consent and any additional patient information must be given to each patient or the patient’s legally accepted representative.

9.10.1.2 Data Quality Assurance

A quality assurance audit/inspection of this study may be conducted by the sponsor or sponsor’s designees or by IECs or by regulatory authorities. The quality assurance auditor will have access to all medical records, the investigator’s study-related files and correspondence, and the informed consent documentation of this study.

9.10.1.3 Records

Case Report Forms (CRFs) for individual patients will be provided by the sponsor via remote data capture. All of the clinical data and site/investigator characteristics will be captured via a web-based Electronic Data Capturing system. The site staff will enter and edit the data via a secure network, with secure access features (username, password and secure identification – an electronic password system). A complete electronic audit trail will be maintained.

Patients must not be identified on the eCRF by name. Appropriately coded identification (i.e. Patient numbers) must be used.

All patient questionnaires will be paper-based and will be left at the site upon completion by the patient.

9.10.1.3.1 Source documents

Source documents provide evidence for the existence of the patient and substantiate the integrity of the data collected. Source documents are filed at the investigator’s site. Data entered in the eCRFs that are transcribed from source documents must be consistent with the source documents or the discrepancies must be explained. The investigator may need to request previous medical records or transfer records; also current medical records must be available.
For the eCRF, the following data must be derived from source documents:

- Patient demographics (age, gender, height, and weight);
- Smoking status / history
- Reported exacerbations
- Past COPD therapies (6 months before visit 1)
- Respimat® training (Yes/No)
- GOLD spirometric classification (1,2, 3, 4)
- GOLD patient groups (A, B, C, D)
- Concomitant diseases / Comorbidities
- Concomitant COPD and other relevant medication
- Breathlessness based on mMRC score
- Physical Functioning (PF-10) Questionnaire
- Patient Satisfaction Questionnaire
- Adverse Drug Reactions (ADR & SADR), fatal AEs, pregnancies
- Rational for Spiolto® Respimat® treatment discontinuation (if applicable)
- Details of treatment continuation / discontinuation

9.10.1.3.2 Details of treatment continuation / discontinuation Direct access to source data and documents

The investigator / institution will permit study-related monitoring, audits, IEC review and regulatory inspection, providing direct access to all related source data / documents. eCRFs and all source documents, including progress notes and copies of laboratory and medical test results must be available at all times for review by the sponsor’s clinical study monitor, auditor and inspection by health authorities. The Clinical Research Associate (CRA) / on site monitor and auditor may review all eCRFs, and written informed consents. The accuracy of the data will be verified by reviewing the documents described in Section 9.10.1.3.1.

9.10.1.4 Statement of Confidentiality

Individual patient medical information obtained as a result of this study is considered confidential and disclosure to third parties is prohibited. Patient confidentiality will be ensured by using patient identification code numbers.

Data generated as a result of the study need to be available for inspection on request by the participating physicians, the sponsor’s representatives, by the IEC and the regulatory authorities, i.e. the competent authority (CA).

9.10.1.5 Completion of Study

The EC/competent authority in each participating EU member state needs to be notified about the end of the study (last patient out) or early termination of the study.
9.10.1.6 Protocol Violations

There are no protocol waivers. All protocol violations must be reported to the sponsor immediately.
10. PROTECTION OF HUMAN SUBJECTS

Please refer to section 9.10.1
11. MANAGEMENT AND REPORTING OF ADVERSE EVENTS/ADVERSE REACTIONS

11.1 DEFINITIONS OF ADVERSE EVENTS

Adverse event
An adverse event (AE) is defined as any untoward medical occurrence in a patient or clinical investigation subject administered a medicinal product and which does not necessarily have a causal relationship with this treatment. An adverse event can therefore be any unfavourable and unintended sign (e.g. an abnormal laboratory finding), symptom, or disease temporally associated with the use of a medicinal product, whether or not considered related to the medicinal product.

Adverse reaction
An adverse reaction is defined as a response to a medicinal product which is noxious and unintended. Response in this context means that a causal relationship between a medicinal product and an adverse event is at least a reasonable possibility. Adverse reactions may arise from use of the product within or outside the terms of the marketing authorisation or from occupational exposure. Conditions of use outside the marketing authorization include off-label use, overdose, misuse, abuse and medication errors.

Serious adverse event
A serious adverse event is defined as any AE which
- results in death,
- is life-threatening,
- requires in-patient hospitalization, or
- prolongation of existing hospitalisation,
- results in persistent or significant disability or incapacity, or
- is a congenital anomaly/birth defect

Life-threatening in this context refers to a reaction in which the patient was at risk of death at the time of the reaction; it does not refer to a reaction that hypothetically might have caused death if more severe.

Medical and scientific judgement should be exercised in deciding whether other situations should be considered serious reactions, such as important medical events that might not be immediately life threatening or result in death or hospitalisation but might jeopardise the patient or might require intervention to prevent one of the other outcomes listed above. Examples of such events are intensive treatment in an emergency room or at home for allergic bronchospasm, blood dyscrasias or convulsions that do not result in hospitalisation or development of dependency or abuse. Any suspected transmission via a medicinal product of an infectious agent is also considered a serious adverse reaction.

Adverse Event of Special Interest (AESI)
The term Adverse Event of Special Interest (AESI) relates to any specific AE that has been identified at the project level as being of particular concern for prospective safety monitoring and safety assessment within this study, e.g. the potential for AEs based on knowledge from other compounds in the same class.

No AESIs have been defined for this study.

11.2 ADVERSE EVENT AND SERIOUS ADVERSE EVENT COLLECTION AND REPORTING

The investigator shall maintain and keep detailed records of all AEs in their patient files.

Collection of AEs

The study design is of non-interventional nature and the study is conducted within the conditions of the approved marketing authorisation. Sufficient data from controlled interventional trials are available to support the evidence on the safety and efficacy of the studied BI drug. For this reason the following AE collection and reporting requirements have been defined.

The following must be collected by the investigator in the (e)CRF from signing the informed consent onwards until the end of the study:

- all adverse drug reaction (ADRs) (serious and non-serious),
- all AEs with fatal outcome,

Note*: For all patients on these data must be recorded on the AE pages in the eCRF. The separate NIS (S)AE form must in addition be used and forwarded to the local Pharmacovigilance as indicated in the form.

All ADRs, including those persisting after study completion must be followed up until they are resolved, have been sufficiently characterized, or no further information can be obtained.

The investigator carefully assesses whether an AE constitutes an ADR using the information below.

Causal relationship of adverse event

The definition of an adverse reaction implies at least a reasonable possibility of a causal relationship between a suspected medicinal product and an adverse event. An adverse reaction, in contrast to an adverse event, is characterised by the fact that a causal relationship between a medicinal product and an occurrence is suspected.

Medical judgment should be used to determine the relationship, considering all relevant factors, including pattern of reaction, temporal relationship, de-challenge or re-challenge, confounding factors such as concomitant medication, concomitant diseases and relevant history.
Arguments that may suggest a **reasonable causal relationship** could be:
- The event is **consistent with the known pharmacology** of the drug
- The event is known to be caused by or **attributed to the drug class**.
- A **plausible time to onset of the event** relative to the time of drug exposure.
- Evidence that the **event is reproducible** when the drug is re-introduced
- **No medically sound alternative etiologies** that could explain the event (e.g. preexisting or concomitant diseases, or co-medications).
- The event is typically **drug-related and infrequent in the general population** not exposed to drugs (e.g. Stevens-Johnson syndrome).
- An indication of dose-response (i.e. greater effect size if the dose is increased, smaller effect size if dose is diminished).

Arguments that may suggest that there is **no reasonable possibility of a causal relationship** could be:
- No plausible time to onset of the event relative to the time of drug exposure is evident (e.g. pre-treatment cases, diagnosis of cancer or chronic disease within days/weeks of drug administration; an allergic reaction weeks after discontinuation of the drug concerned)
- Continuation of the event despite the withdrawal of the medication, taking into account the pharmacological properties of the compound (e.g. after 5 half-lives).
  Of note, this criterion may not be applicable to events whose time course is prolonged despite removing the original trigger.
- Additional arguments amongst those stated before, like alternative explanation (e.g. situations where other drugs or underlying diseases appear to provide a more likely explanation for the observed event than the drug concerned).
- Disappearance of the event even though the study drug treatment continues or remains unchanged.

**Intensity of adverse event**
The intensity of the AE should be judged based on the following:
- **Mild**: Awareness of sign(s) or symptom(s) which is/are easily tolerated
- **Moderate**: Enough discomfort to cause interference with usual activity
- **Severe**: Incapacitating or causing inability to work or to perform usual activities

**Pregnancy**:
In rare cases, pregnancy might occur in this study. Once a subject has been enrolled into the study with Spiolto® Respimat®, the investigator must report any drug exposure during pregnancy, which occurred in a female subject or in a partner to a male subject, to the Sponsor by means of Part A of the Pregnancy Monitoring Form. The outcome of the pregnancy associated with the drug exposure during pregnancy must be followed up and reported by means of Part B of the Pregnancy Monitoring Form.

In the absence of a reportable AE, only the Pregnancy Monitoring Form must be completed, otherwise the NIS AE form is to be completed and forwarded to the local Pharmacovigilance point of contact for each country within respective timelines.

**Expedited Reporting of AEs and Drug Exposure During Pregnancy**
The following must be reported by the investigator on the NIS AE form from signing the informed consent onwards until the end of the study:

Table 11.2:1 Reporting types and timelines

<table>
<thead>
<tr>
<th>Type of Report</th>
<th>Timeline</th>
</tr>
</thead>
<tbody>
<tr>
<td>All serious ADRs associated with Spiolto® Respimat®</td>
<td>immediately within 24 hours</td>
</tr>
<tr>
<td>All AEs with fatal outcome in patients exposed to</td>
<td>immediately within 24 hours</td>
</tr>
<tr>
<td>All non-serious ADRs associated with Spiolto® Respimat®</td>
<td>7 calendar days</td>
</tr>
<tr>
<td>All pregnancy monitoring forms associated with Spiolto® Respimat®</td>
<td>7 calendar days</td>
</tr>
</tbody>
</table>

The same timelines apply if follow-up information becomes available for the respective events. In specific occasions the Investigator could inform the Sponsor upfront via telephone. This does not replace the requirement to complete and fax the NIS AE form.

Information required

For each reportable adverse event, the investigator should provide the information requested on the appropriate (e)CRF pages and the NIS AE form.

Reporting of related Adverse Events associated with any other BI drug

The investigator is encouraged to report all adverse events related to any BI drug other than the associated with Spiolto® Respimat® according to the local regulatory requirements for spontaneous AE reporting at the investigator’s discretion by using the locally established routes and AE report forms. The term AE includes drug exposure during pregnancy, and, regardless of whether an AE occurred or not, any abuse, off-label use, misuse, medication error, occupational exposure, lack of effect, and unexpected benefit.

11.3 REPORTING TO HEALTH AUTHORITIES

Adverse event reporting to regulatory agencies will be done by the MAH according to local and international regulatory requirements. Also the investigator is encouraged to report all adverse event related to any drug to LHA according to local regulatory requirements.
12. PLANS FOR DISSEMINATING AND COMMUNICATING STUDY RESULTS

Results of this non-interventional study will be disclosed on encepp.eu and clinicaltrials.gov and a study specific publication plan will be developed to describe planned publications.
13. REFERENCES

13.1 PUBLISHED REFERENCES


P10-09337 Bouyssou T, Schnapp A, Casarosa P, Pieper MP. Addition of the new once-daily LABA BI 1744 to tiotropium results in superior bronchoprotection in pre-clinical models. ATS 2010, 106th Int Conf of


13.2 UNPUBLISHED REFERENCES
n.a.
ANNEX 1. LIST OF STAND-ALONE DOCUMENTS
ANNEX 1. LIST OF STAND-ALONE DOCUMENTS

The stand-alone documents for this non-interventional study are:

- Informed Consent Form
- PF-10 Questionnaire based on SF-36
- Breathlessness Scale (mMRC)
- Patient Satisfaction Survey
- Statistical Epidemiological Analysis Plan (SEAP)
- Data Management Plan (DMP)
- Serious Adverse Event Report in Non-Interventional Studies - (S)AE NIS Form
- Pregnancy Monitoring Form
- Publication Plan

All of the above documents will be archived in the Trial Master File in its original English master version.
ANNEX 2. ENCEPP CHECKLIST FOR STUDY PROTOCOLS

Not applicable.
ANNEX 3. ADDITIONAL INFORMATION

1. Physical Functioning Questionnaire (PF-10) based on the 36-Item Short Form Survey.

The following questions are about activities you might do during a typical day. Does your health now limit you in these activities? If so, how much?

<table>
<thead>
<tr>
<th></th>
<th>Yes, limited a lot</th>
<th>Yes, limited a little</th>
<th>No, not limited at all</th>
</tr>
</thead>
<tbody>
<tr>
<td>a) Vigorous activities,</td>
<td>☐ 1</td>
<td>☐ 2</td>
<td>☐ 3</td>
</tr>
<tr>
<td>b) Moderate activities,</td>
<td>☐ 1</td>
<td>☐ 2</td>
<td>☐ 3</td>
</tr>
<tr>
<td>c) Lifting or carrying</td>
<td>☐ 1</td>
<td>☐ 2</td>
<td>☐ 3</td>
</tr>
<tr>
<td>d) Climbing several</td>
<td>☐ 1</td>
<td>☐ 2</td>
<td>☐ 3</td>
</tr>
<tr>
<td>e) Climbing one flight</td>
<td>☐ 1</td>
<td>☐ 2</td>
<td>☐ 3</td>
</tr>
<tr>
<td>f) Bending, kneeling, or</td>
<td>☐ 1</td>
<td>☐ 2</td>
<td>☐ 3</td>
</tr>
<tr>
<td>g) Walking more than a</td>
<td>☐ 1</td>
<td>☐ 2</td>
<td>☐ 3</td>
</tr>
<tr>
<td>h) Walking several hundred yards</td>
<td>☐ 1</td>
<td>☐ 2</td>
<td>☐ 3</td>
</tr>
<tr>
<td>i) Walking one hundred</td>
<td>☐ 1</td>
<td>☐ 2</td>
<td>☐ 3</td>
</tr>
<tr>
<td>j) Bathing or dressing</td>
<td>☐ 1</td>
<td>☐ 2</td>
<td>☐ 3</td>
</tr>
</tbody>
</table>
2. Modified Medical Research Council (mMRC) Questionnaire for Assessing the Severity of Breathlessness

Please circle the number which best describes your grade of breathlessness.

I only get breathless with strenuous exercise. 0

I get short of breath when hurrying on the level or walking up a slight hill. 1

I walk slower than people of the same age on the level because of breathlessness, or have to stop for breath when walking at my own pace on the level. 2

I stop for breath after walking about 100 meters or after a few minutes on level. 3

I am too breathless to leave the house or I am breathless when dressing or undressing. 4

3. Patient Satisfaction Questionnaire

Please choose the number which best describes your satisfaction with Spiolto® Respimat®.

What is your overall satisfaction with the Spiolto® Respimat® treatment?

1 very dissatisfied 2 dissatisfied 3 rather dissatisfied 4 neither satisfied nor dissatisfied 5 rather satisfied 6 satisfied 7 very satisfied

How satisfied are you with inhaling from the Respimat® device?

1 very dissatisfied 2 dissatisfied 3 rather dissatisfied 4 neither satisfied nor dissatisfied 5 rather satisfied 6 satisfied 7 very satisfied
How satisfied are you with the handling of the Respimat® inhalation device?

<table>
<thead>
<tr>
<th></th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 very</td>
<td></td>
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<td>dissatisfied</td>
<td></td>
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<tr>
<td>2 dissatisfied</td>
<td></td>
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<td></td>
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<tr>
<td>3 rather dissatisfied</td>
<td></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4 neither satisfied nor dissatisfied</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td>5 rather satisfied</td>
<td></td>
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<td>6 satisfied</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>7 very satisfied</td>
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<td></td>
</tr>
</tbody>
</table>

4. Physicians’ Global Evaluation (PGE) to be used directly within the eCRF

General condition of the patient at the initial examination (Visit 1)
Please mark with a cross as applicable

<table>
<thead>
<tr>
<th>Poor</th>
<th>Satisfactory</th>
<th>Good</th>
<th>Excellent</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>5</td>
<td>6</td>
<td>7</td>
<td>8</td>
</tr>
</tbody>
</table>

General condition of the patient after approximately 6 weeks of treatment (Visit 2)
Please mark with a cross as applicable

<table>
<thead>
<tr>
<th>Poor</th>
<th>Satisfactory</th>
<th>Good</th>
<th>Excellent</th>
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