ABSTRACT

Objective: The present study was conducted with the objective of analyzing the efficacy of triple-drug combination therapy (formoterol, ciclesonide, tiotropium) by comparing it with double drug combination therapy (formoterol, budesonide).

Methods: A prospective observational study was conducted. Sixty patients were enrolled, and divided into two groups of thirty each; one group was treated with the double-drug and the other with a triple drug combination. FEV1 and FVC pre-and post-treatment in either group were assessed spirometrically. Score ranges of 0-10, 11-20 and 21-30 were allotted to mild, moderate and severe categories and results were analyzed statistically.

Results: Of the 60 patients recruited, 61-70 yrs olds constituted the majority (35%) of the population. Males (63.3%) were more in number compared to females (36.6%). Twenty-three of thirty-eight men smoked (60.5%); there were no female smokers. Common symptoms included cough (93.3%), dyspnoea (85%), fever (45%) and haemoptysis (15%). Hypertension accounted for 70% of patient comorbidities, followed by diabetes (60%) and cardiovascular diseases (40%). Three months after treatment with triple therapy, a significant increase in the differences of means of both FEV1 (14.27) and FVC (14.90) values was observed. Further analysis based on score ranges demonstrated that triple therapy administration markedly reduced the number of patients suffering from severe COPD.

Conclusion: Our comparative analysis indicated that triple therapy was more effective in improving lung function, enhancing patients’ quality of life (evidenced from score ranges) thereby reducing mortality. While much is known about the greater effectiveness of triple over dual therapy, researchers to formulate the most effective triple therapy are in progress.

Keywords: Chronic obstructive pulmonary disease (COPD), Inhaled corticosteroids, Long-acting β2 agonist, Long-acting muscarinic antagonist, Spirometry

INTRODUCTION

The history of COPD can be traced back to Dr. Rene Laennec, who was the first to precisely establish a relationship between chronic bronchitis and emphysema [1]. Over the years, advancements in the understanding of the disease, its explicit etiology, detailed pathology and distinct pharmacotherapy have aided in improving patient outcomes [1, 2]. In 1962, the American Thoracic Society Committee defined emphysema as “anatomic alteration of the lung characterized by an abnormal enlargement of the airspace distal to the terminal, non- respiratory bronchiole accompanied by destruction of the alveolar walls” [3]. Chronic bronchitis on the other hand, was defined in 1965 as “a cough with expectoration that has occurred on most days during at least three consecutive months for more than two successive years” [4]. As of now, the Global Initiative for Chronic Obstructive Lung Disease (GOLD) defines COPD as “common, preventable and treatable disease that is characterized by persistent respiratory symptoms and airflow limitation that is due to airway and/or alveolar abnormalities usually caused by significant exposure to noxious particles or gases” [5].

Analogous to the discovery of COPD which coursed gradually through decades, the establishment of a peculiar treatment therapy for the disease took years to develop. Until the 1950s, no specific treatment was authorized, excepting a few sedatives and antibiotics. Corticosteroids were almost never used. During the 1960s, mechanical ventilators and long-term oxygen therapy measures began to be extensively studied and trialled and were shown to improve outcomes, hence used in patients with longstanding COPD [1]. Smoking as a cause of respiratory derangements was authenticated only in the late 1990s [6]. By the end of the twentieth century, available treatment options included smoking cessation, antibiotics, bronchoactive agents and steroids [7].

 Ranked as the fifth leading cause of death in 2002 [8], COPD accounted for six percent of the world’s mortality in 2012 [5]. By 2030, it is expected to secure the fourth position and may account for 7.8% of total deaths globally [8]. The disease is more probable in high-income countries [9]. The development of COPD is multifactorial, risk factors implicated in its etiology include: cigarette smoking, indoor (biomass) or outdoor environmental pollutants, genetic factors and recurrent juvenile infections of the lower respiratory tract [5, 10]. Spirometric irregularity, flawed arterial blood gas tests and other findings on physical examination hint to the presence of COPD [5, 10, 11].

Present day medications used to treat COPD include beta-agonists, parasympatholytics, methylxanthines, corticosteroids, phosphodiesterase inhibitors, and combination therapies incorporating two or more drugs [12]. There has been some controversy regarding the effectiveness of one combination regimen over the other. Therefore, the current research was conducted with the objective of comparing dual drug with triple-drug therapy and evaluating the efficacy of either regimen. Spirometric analysis was utilized for obtaining reliable results.

MATERIALS AND METHODS

The present prospective observational study was carried out from September 2014 to February 2015 in the Department of Pulmonology, Krishna Institute of Medical Sciences (KIMS) Hospital, Secunderabad, Telengana, India. Data for this study was collected prospectively from case sheets and from consent-obtained questionnaires distributed to patients in both inpatient (IP) and
outpatient (OP) departments. The study was approved by the KIMS Foundation and Research Centre Ethics Committee (KFRC/EC/APR/024/2015, dated 14.02.15). All patients diagnosed with chronic obstructive pulmonary disease and aged 40 y or above were the study population. Pregnant or lactating women, patients who portrayed reluctance for enrollment, and those who failed to show up for subsequent review were excluded from the study.

**Study procedure**

The study was conducted on 60 patients suffering from COPD. The participants were divided into two groups of 30 each. One group received a dual combination inhaler of formoterol 6 µg and budesonide 100 µg, while the other group was given a triple drug combination inhaler of formoterol 6 µg, ciclesonide 200 µg and tiotropium 9 µg. Before the commencement of treatment, patients’ FEV1 and FVC values were recorded, while ‘symptoms encountered’ were assessed with the aid of a questionnaire validated by a registered clinician. Patients were required to appear for a follow-up visit to the clinic three months succeeding initiation of therapy. During follow-up visits, post-treatment FEV1 and FVC values were charted, while the distribution of questionnaires analogous to those previously distributed, enabled reassessment of any persisting symptoms. Data collected was then subjected to statistical analysis.

**Statistical analysis**

We used Microsoft Excel 2007 and mean of samples to compute the observed values of pre-and post-treatment.

The following parameters were analysed statistically:

- **FEV1 and FVC values pre-and post-treatment in dual therapy.**
- **FEV1 and FVC values pre-and post-treatment in triple therapy.**

A comparative pre-and post-treatment analysis in either group was then done to assess any improvement in symptoms.

**RESULTS**

**Demographic distribution of patients**

In the present study, a total of sixty patients were enrolled and distributed according to their age (<50 y or >50 y) and sex. Fifteen patients were aged below 50 y (9 males and 6 females), while the remaining 45 were above 50 y of age (29 males and 16 females).

Demographic distribution of patients is shown in Fig. 1.

**Age distribution of patients**

Age distribution of patients was enumerated and it was found that 35% of the diseased population was aged between 61 and 70 y. This was followed by the age group of 51-60 y who constituted another 25% of the total. Age groups of 41-50 and 71-80 y olds constituted 15% each. The least prone was the age range of 81-90 y (10%).

**Gender wise distribution of patients**

The sample consisted of 38 men (majority of the sample size) and 22 women suffering from the chronic illness. The percentages in men and women were reported as 63.33% and 36.67% respectively.

**Smokers and non-smokers having COPD**

From the 60 patients enrolled in the study, 38 were males and 22 were females. No female smokers were documented in this prospective study. However, among the 38 men, there were 23 smokers and 15 non-smokers. Smokers accounted for about 38.33% while non-smokers, from the total, made up the remaining 61.67%.

**Presence of symptoms in COPD patients**

We categorised the manifestations of COPD as cough, shortness of breath, fever and haemoptysis. From a sum of 60 patients, 56 experienced cough, accounting for 93.33%, 51 suffered from shortness of breath, accounting for 85%, 27 presented with pyrexia, constituting 45%, and 9 patients experienced haemoptysis, accounting for 15% of the total subjects.

**Comorbidities in COPD**

Comorbidities such as hypertension, heart diseases, diabetes and others such as psychological disorders are usual among patients with the chronic respiratory disease. Data obtained from the present study demonstrated that majority of patients were hypertensive (70%); the second most common comorbidity was diabetes mellitus (60%). Cardiovascular and psychological disorders accounted for 40 and 10% of the sample size respectively.

**Statistical analysis of FEV1 and FVC values**

Statistical analysis was carried out for comparing FEV1 and FVC values pre-and post-treatment in dual and triple therapies, shown in table 1 (a) and (b).

<table>
<thead>
<tr>
<th>Statistical value</th>
<th>Dual therapy</th>
<th>FEV1</th>
<th>Triple therapy</th>
<th>FEV1</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>FVC</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Pre-treatment</td>
<td>Post-treatment</td>
<td>Pre-treatment</td>
<td>Post-treatment</td>
</tr>
<tr>
<td>Mean</td>
<td>46.70</td>
<td>55.30</td>
<td>46.97</td>
<td>55.83</td>
</tr>
</tbody>
</table>

On thoroughly inspecting the data, the difference between the mean of FVC pre-and post-treatment in dual therapy was found to be 8.60 and the difference between the mean of FEV1 pre-and post-treatment was calculated as 8.87. In triple therapy, the difference between the mean of FVC pre-and post-treatment was 14.90 and the difference between the mean of FEV1 pre-and post-treatment was 14.27.
Distribution of patients based on disease severity

For judging the comparative superiority of one drug regimen over the other, we graded the disease severity based on specific score ranges. Scores of 0-10, 11-20 and 21-30 represented mild, moderate or severe disease states respectively. We then compared the score ranges pre-and post-treatments in either therapy to make a conclusion regarding the efficacy of one over the other.

Comparative distribution of patient’s pre-and post-treatment with dual therapy

The former group of subjects undergoing dual therapy were categorized as mild, moderate and severe, with respect to their scores before medication administration. Out of 30 patients, it was observed that 36.67% patients suffered from moderate COPD, while a majority 63.34% suffered from severe COPD. No mild cases were documented.

Three months post-treatment, the highest cases observed were those of moderate COPD (66.67%), followed by mild (20%) and severe (13.34%) disease. Table 2 compares the scores of patients before and after treatment with dual therapy.

Comparative distribution of patient’s pre-and post-treatment with triple therapy

The latter group of subjects undergoing triple therapy were classified as mild, moderate and severe, based on their scores before the onset of therapy. From a total of 30 patients, severe cases took the lead (66.67%), the successors being moderate (30%) and mild cases (3.34%).

After three months, it was found that 46.67% were moderate, 43.34% were mild and the least documented were severe cases (10%). Table 3 compares the scores of patients before and after treatment with triple therapy.
aged between 51 and 70 y constituted the majority (60%) of the population. This was probably due to long-term occupational exposure and indoor and outdoor pollution [5]. Other studies report a higher prevalence in patients aged 70 y and above [15–18], attributing the cause to tobacco smoking and advancing age [15]. The least prone was the age group of 50-59 y, credited to the healthy survivor effect [15, 18], in which individuals succumb to death before 80 y of age owing to chronicity of the illness and associated comorbidities.

Gender wise analysis reports a higher preponderance in males (63.3%) compared to females (36.7%). This was consistent with the results of other researchers [9, 15-18]. A contributing element in this regard is habitual smoking in men.

The current research consisted of 23 male smokers, accounting for 65.5% of the male population and 38.3% of the total. No female smokers were documented. The two most essential etiologic agents implicated in COPD are smoking and environmental pollutants [19, 20]. As men exhibit considerable exposure to either of these, their dominance in this respect is clearly comprehensible.

The GOLD guidelines, however, have stated increasing age and female sex as risk factors for COPD [5]. Over the past few decades, cigarette smoking by women in developed countries has become ‘trendy’ [9], leading to more rapidly deteriorating lung function and a greater annual decline in FEV1 [21]. This has contributed to a notable increase in the incidence of the disease among young females aged 50-59 y and above, while prevalence is still dominated by males [9, 15-18, 21]. Additionally, recent researches have evinced a possible link between COPD and autoimmunity. Consequently, as women are more prone to autoimmune disorders, they would unambiguously be inclined to develop COPD [21].

Symptom evaluation revealed that almost all the patients experienced cough (93.3%) and dyspnoea (85%). The successors were fever (45%) and haemoptysis (15%). While the former two are the result of airway obstruction and excessive mucus secretion [22], latter manifestations may indicate oncoming respiratory infections. However, diagnosis of COPD cannot rely solely on clinical manifestations, as a small percentage of patients present with no respiratory symptoms at all [15]. The guidelines, therefore, portray an organized approach to assessing affected patients [5].

Association between COPD and comorbidities is complex. Our study showed that hypertensive patients constituted 70% of the sample total. Diabetes mellitus was the second most prevalent comorbidity. This was proceeded by cardiovascular diseases (40%), pneumonia (25%) and psychological abnormalities (10%). The co-existence of a disease alongside COPD is multifactorial and is ultimately specific to the patient [16]. However, several mechanisms have been proposed to explain the relation between the two. Impairment of lung function due to the chronic illness builds pressure in pulmonary vessels, causing pulmonary hypertension. This further increases the workload on the heart and may result in the development of cardiovascular disorders such as ischemic heart disease, heart failure and arrhythmias [23]. Right ventricular failure, if present, may remain unaltered despite ongoing treatment [24]. The relation between diabetes and COPD is intricate, with each pre-existing condition worsening the prognosis of the other. Pre-existing diabetes aggravates COPD through direct harmful effects of hyperglycaemia on lung function, while pre-existing COPD augments diabetes via its deteriorating effects on insulin resistance [25].

After gauging FEV1 and FVC values before and after treatments, we allotted score ranges of 0-10, 11-20 and 21-30 to mild, moderate and severe categories and assessed the superiority of one regimen over the other. From our observations, we concluded that the number of patients suffering from severe COPD decreased significantly in triple-drug therapy trial, indicative of its increased efficacy over dual-drug regimen. Our findings coincided with those of others [26-28].

The current research involved administration of triple therapy to patients independent of disease severity, in contrast to GOLD guidelines which recommend reservation of the triple drug regime for severe cases with frequent exacerbations [14]. Singh et al. [26] and Cazzola et al. [27] conducted analogous researches by comparing triple therapy involving LABA, LAMA and ICS with LABA+ICS or LAMA alone. Both the authors reported better outcomes with a triple-drug regimen, evident from improvement in pulmonary tests. Singh et al. [26] Additionally inferred that triple therapy is more effective with a single dose and that it elicits its therapeutic response gradually over a period of two weeks [26]. Anticholinergics exert their effects by blocking parasympathetically induced bronchoconstriction [12]. They also aid in enhancing FEV1, decrease inflation of lungs [26] while minimizing exacerbations [29]. While studies claim the efficacy of tiotropium, salmeterol and fluticasone to be equal to glycopyrromium, salmeterol and fluticasone, there exists minor superiority of the latter in improving FEV1 [30] and eliciting peak response with only a single dose [31].

The use of corticosteroids over longer durations is raising concerns owing to its resistance and related adverse effects [29]. Researches in this field are ongoing, with trials to replace ICS with the better phosphodiesterase-4 inhibitors or other alternatives, where possible [12, 14].

Limitation of the present study was its small sample size. The study was also confined to three months duration and therefore, could not address maintenance therapy and associated adverse effects. The study also excluded the exacerbations of COPD.

CONCLUSION

In conclusion, our data suggested that the triple drug combination of formoterol, iciclesonide and tiotropium showed more efficacy in patients suffering from COPD than the dual drug combination of formoterol and budesonide. The former group was shown to elicit its therapeutic response by producing a greater reduction in FVC and FEV1 post-treatment. The use of triple therapy also extends in reducing the number of patients suffering from severe COPD. It demonstrates benefits on lung function and improvement of symptoms, thereby enhancing patients' quality of life. Additionally, patients with advanced COPD reported that triple therapy combined with pulmonary rehabilitation provided substantial benefit in terms of lung function.

With the present available literature, the question “Is three better than two?” seems to be answered with conclusive evidence. The new question that arises is “Which three is better?” The search for the most effective triple therapy portraying maximal benefits and with least documented adverse effects is still ongoing.

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AUTHORS CONTRIBUTIONS

All authors have contributed equally to the research work.

CONFLICT OF INTERESTS

All authors have none to declare.

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