Living with chronic obstructive pulmonary disease in Lebanon: a phenomenological study

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Abstract

Background: Lebanon has the fastest growing older adult population in the Arab region but few social resources to address their needs. No studies have explored the experience of patients with chronic obstructive pulmonary disease (COPD) in Lebanon.

Aims: Exploring the experiences of individuals living with COPD in Lebanon.

Method: Using a descriptive phenomenological research design, qualitative individual semi-structured interviews were conducted with COPD patients living in Lebanon, between May 2019 and September 2019.

Results: Fifty participants agreed to be interviewed. The majority were men (56%) and had moderate COPD (40%). Mean age was 71.5 (standard deviation 9.0) years. We found that COPD affects three dimensions of patients’ lives: educational, organizational and psychosocial.

Conclusion: The results highlight the need for multidisciplinary strategies to address the needs of people with COPD in Lebanon, including their caregivers. Strategies include patient education and the development of new methods to facilitate and promote partnership between health care professionals, COPD patients and their caregivers.

Keywords: COPD, elderly, qualitative study, phenomenology, Lebanon, morbidity, mortality

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Introduction

Chronic obstructive pulmonary disease (COPD) is one of the leading causes of morbidity and mortality worldwide. Studies estimate that by 2030, COPD would be the fifth leading cause of death in high-income countries and the third leading cause of death in middle-income countries (1). Since 2016, research has shown that COPD is already established as the third leading cause of death in the world with about 3 million deaths and 251 million cases, 90% of which occurred in low- and middle-income countries (2) such as Lebanon.

The first, and so far only, national study to determine the prevalence of COPD in the adult Lebanese population revealed a prevalence of 9.7% according to the GOLD definition, of which 80% of cases in the smoking population are unrecognized and undiagnosed (3). Moreover, a study conducted in 10 countries in the Middle East and North Africa (MENA) region showed that in Lebanon, 71.8% of COPD patients were still smoking and only 30.6% were receiving respiratory treatment (4).

Complete recovery from COPD is currently not achievable, and as the disease progresses patients experience worsening breathing difficulty and disruption that affect their quality of life (5,6). Therefore, COPD requires major changes in the daily lives of patients, who must adhere to treatment, change their lifestyle and monitor their signs and symptoms (7).

Essentially, COPD has been considered a condition of accelerated lung aging; consequently, the prevalence of COPD is three times higher in people over the age of 60 years (8). Lebanon has the fastest growing older adult population in the Arab region, but few social resources to address their needs (9). To date, no study has explored the experience of patients with COPD in Lebanon. Thus, the aim of our study was to describe the phenomenon of living with COPD from the Lebanese individuals’ perspective.

Methods

Design

A descriptive phenomenological research design was used, which is anchored in Husserl’s philosophical phenomenology, and developed and modified to be applicable as a scientific research method. The aim of phenomenology is to seek the essence of human phenomena as lived and experienced and present them and their meanings as faithfully as possible (10), free from preconceptions, beliefs and knowledge of the phenomenon.
Throughout this study, we followed the Standards for Reporting Qualitative Research guidelines (11).

Participants
The participants were recruited from four university hospitals in Beirut. Eligibility criterion was clinical diagnosis by pulmonologists, and participation was voluntary. First, the study was explained to the pulmonologists, then 135 patients on the lists they provided were contacted and asked if they were willing to be interviewed at their homes. The inclusion criteria were COPD diagnosis with over 40 years old, and being able and willing to consent to and engage in the research study.

Creswell states that “up to a maximum of 10 interviews” is sufficient for phenomenological research (12). However, because the size of a sample depends on the complexity of a phenomenon (13) and in order to create a rich understanding of the phenomenon of living with COPD in Lebanon, we included all participants who agreed to take part in the study, therefore covering and thus a broad range of people using characteristics such as sex, age and economic situation.

Data collection
Face-to-face, semi-structured interviews were conducted by the principal investigator between May 2019 and September 2019 at the participants’ homes. Interviews were systematically audio and video recorded after obtaining the participant’s consent. An interview guide with open questions was developed in advance by the principal investigator and tested with one participant. The interviews were relatively open and partly influenced by the participant’s concern. The investigator was well trained and careful to avoid leading questions. After some sociodemographic questions, the interview began with an opening question, “Tell me more about your experience with COPD”. Next, a series of prompts were used inviting the participants to describe their experiences and their visions for the future. The investigator then summarized the exchange so that the patient could validate whether they were in agreement with what had been said during the interview. The participants were visited a second time to present the results. All participants agreed with the results.

Data analysis
Analysis of the results started immediately after the interviews. The audio and video recordings were fully transcribed verbatim in Arabic and typed using Microsoft Word. Emotions, voice tone, or any changes in participants’ behaviours noticed in the videos were added to the observation notes. We used ATLAS.ti, version 7, a software for qualitative data analysis, to support the analysis.

The data analysis was inspired by Giorgi’s phenomenological method (10). Each interview was first read completely to gain a sense of the whole description. The interview was then reread from the start to identify key statements about the participants’ experiences, constituting “meaning units” expressed in the participants’ own everyday language. Once the meaning units were established, transformation of the participants’ everyday language was required (13). In this step, the statements of the participants were transformed by two investigators to express the insight contained in them more directly. Then the meaning units of the 50 interviews were combined inferring the general meaning structure of the experience from the transformed meaning units (14).

Ethical considerations
The study was authorized by the ethics committee of Hôtel-Dieu de France Hospital (CEHDF 1135). Oral and written informed consent was obtained from each participant after being provided with both written and oral information about the aim of the study. A number was randomly assigned to each patient (P1, P2, etc.) and a concordance table was kept to preserve anonymity.

Results
Demographics
A total of 135 patients were contacted, and 50 agreed to be interviewed, of whom 45 were filmed and 5 were audio-recorded. This number of participants was necessary to gain insight into the variety of meanings of the phenomena of living with COPD in Lebanon. The participants lived in different regions of Lebanon: 27 in Mount Lebanon, 21 in Beirut, and 2 in southern Lebanon.

The age of the respondents varied between 50 and 93 years, with a mean of 71.5 (standard deviation, 9) years and a sex ratio of 1.27 male/female (28 males and 22 females). The majority of participants had moderate COPD (40%), used support from a caregiver (84%) and were retired (64%) (Table 1).

Living with chronic obstructive pulmonary disease
The essence of living with COPD in Lebanon, from the participants’ perspective, is to be seen as living in a “lesser world”, striving for a breath of life. Living with COPD means living with vulnerability and uncertainty influenced by the body’s capacity and the environmental resources that impose the need for support to cope with daily life (Table 2).

Chronic obstructive pulmonary disease: vulnerability and uncertainty
Participants described living with COPD as being forced into sedentary and solitary lifestyle. The physical impact due to COPD expressed by participants ranged from fatigue and dyspnoea to inactivity and dependence, either because of the inability to move or the oxygen device that impede their movement, resulting in a negative experience. Living with this disease is living with dyspnoea that can worsen at any time. Difficulties in predicting the symptoms of COPD and their impact on daily life limit the participants’ ability to plan activities in advance.
Therefore, they avoid family gatherings and activities which they think they cannot cope with, e.g. walking alone in the garden, which in itself presents a risk of being sedentary. Moreover, they prefer to receive their care at home, either because it is difficult for them to move around or to preserve the same living conditions.

Participants described COPD as an anxiety-provoking disease that invades all aspects of their lives. They live in constant fear; fear of everything new, fear of long journeys, and fear that the disease will have an impact on their professional lives. An older woman described her fear as “suffocating is not funny”.

Concerns related to the Lebanese context, such as financial problems and the feeling of insecurity, make the participants to organize their daily lives according to their living context; which oxygen machine to use in case of a power outage; when to leave the house; what to do if the building does not have an elevator; and financial problems that prevent certain participants from visiting their doctors or even adhering to their medications.

Therefore, to save on expensive physiotherapy sessions, participants taught their caregivers how to help them with breathing exercises. Some participants used computerized consultation because of the difficulty in going to meet with their physician physically.

COPD has an impact on participants’ lives; it makes them feel resigned to their state of health as they struggle to maintain vital momentum. Some participants reported the impact of the disease on their physical image and some related these difficulties to the cultural context of the country.

Chronic obstructive pulmonary disease: striving for a breath of life

Some participants tried to continue their lives as normally as possible, taking precautions not to get sick, adhering to the recommendations of the pulmonologist, setting goals so as not to aggravate their symptoms and contacting their pulmonologist regularly. They wanted

<p>| Table 1 Demographic characteristics of the participants (n = 50) |
|-----------------|-----------------|-----------------|-----------------|</p>
<table>
<thead>
<tr>
<th>Characteristic</th>
<th>No.</th>
<th>Patient number</th>
<th>Frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&lt; 60</td>
<td>5</td>
<td>P8, P26, P27, P28, P34</td>
<td>10</td>
</tr>
<tr>
<td>60–69</td>
<td>15</td>
<td>P3, P4, P6, P15, P17, P18, P19, P21, P24, P25, P32, P36, P44, P46, P47</td>
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<tr>
<td>&gt; 80</td>
<td>7</td>
<td>P9, P11, P14, P39, P42, P45, P49</td>
<td>14</td>
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<tr>
<td>Sex</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Male</td>
<td>28</td>
<td>P1, P4, P5, P6, P7, P9, P13, P16, P18, P19, P20, P21, P22, P23, P24, P26, P27, P29, P30, P31, P32, P36, P37, P40, P41, P42, P43, P50</td>
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<tr>
<td>Female</td>
<td>22</td>
<td>P2, P3, P8, P10, P11, P12, P14, P15, P17, P25, P28, P33, P34, P35, P38, P39, P44, P45, P46, P47, P48, P49</td>
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<tr>
<td>Disease severity</td>
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<td></td>
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<tr>
<td>Mild</td>
<td>6</td>
<td>P13, P17, P18, P20, P22, P28</td>
<td>12</td>
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<tr>
<td>Severe</td>
<td>10</td>
<td>P7, P9, P10, P11, P27, P33, P37, P40, P42, P45</td>
<td>20</td>
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<tr>
<td>Very severe</td>
<td>14</td>
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<td>28</td>
</tr>
<tr>
<td>Age at diagnosis (years)</td>
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<td></td>
<td></td>
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<tr>
<td>&lt; 5</td>
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<tr>
<td>5–9</td>
<td>19</td>
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</tr>
<tr>
<td>&gt; 10</td>
<td>7</td>
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<td>14</td>
</tr>
<tr>
<td>Caregiver</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Family member</td>
<td>34</td>
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<td>68</td>
</tr>
<tr>
<td>Non-family member</td>
<td>8</td>
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<td>16</td>
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<tr>
<td>Without help</td>
<td>8</td>
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<tr>
<td>Current work</td>
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<td></td>
<td></td>
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<tr>
<td>Yes</td>
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<td>30</td>
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to know more about their illness and some became self- 
sufficient and active in their own care.

The experiences of participants with their disease was 
influenced by the support they receive, whether from 
family or medical personnel. They reported how helpful 
the motivation provided by their doctor was. For some, 
it was difficult to maintain compliance and set goals, 
so they reported their need for information and regular 
reassuring follow-ups.

Some participants expressed their need for help to 
quit smoking and remain so, but rejected the idea 
of attending specialized centers because of their 
constraining conditions and because they were convin-
ced that quitting is a matter of self-determination.

Monitoring, whether by the doctor or the family, is a key factor that helped participants in dealing with 
their disease. Participants’ families and friends helped 
them by securing suitable tobacco-free environments 
and a sense of safety. Participants clearly expressed the 
benefits of close monitoring to ensure that the right 
decisions are made, to detect any deterioration in health 
status or simply to feel that someone is interested in their 
condition.

Discussion

Participants in this study were held back by the symp-
toms of their disease, they struggled with physical move-
ments (e.g. walking and in some cases even speaking), 
which led to a more sedentary and solitary life than they 
had previously or wish to have. In line with our findings, 
previous studies have found that many COPD patients 
struggle with being held back by the symptoms of their 
disease and the challenges of breathing, which is associ-
ated with social isolation (15–20). Health care staff, par-
cifically nurses, play a major role in managing the health 
issues of patients with chronic illness (20,21). Nurses 
should be more aware that the sedentary lifestyle of 
COPD patients might be by necessity and not by choice, 
and therefore adopt a sensitive approach. Moreover, 
nurses are encouraged to develop strategies for patients 
to maintain physical and social activities and adhere to 
clinical recommendations.

It is important to recognize that respiratory symptoms 
are prevalent and distressing in pulmonary disease. All 
participants described anxiety, fear, panic or distress with 
shortness of breath with having to stop to get air in. They 
reported being stigmatized because of their symptoms, 
and this is consistent with the findings from other studies 
(22–25). Some even isolated themselves at home to avoid 
the embarrassment of showing their symptoms in public. 
From a phenomenological point of view, moving from 
avoidance to acceptance of one’s own body is a crucial 
shift since the body is understood to be a mode of access to 
the world (26). From this perspective, it is understandable 
that the negative feelings due to the symptoms of COPD 
lead to a distancing from one’s own body and certain 
dimensions of the everyday world. This contributes to 
the risk of living a sedentary lifestyle and reduced 
social network, resulting in further deterioration. Thus, 
it is important to adopt a multidisciplinary approach to 
addressing the psychosocial needs of COPD patients.

To overcome problems related to the unavailability of 
the doctor, the difficulty in moving around, long journeys 
to reach the clinic, or long waiting times at the doctor’s 
clinic, participants used the internet and TV shows to 
find out more about their disease, which exposed them 
to wrong information, unsuited to their state of health. 
Some participants reported the use of self-medication, 
consistent with other research (27). Even in emergency 
situations, participants declared the use of prescriptions 
collected during previous consultations. According to 
a previous survey, the vast majority of drugs used in 
self-medication came from a resumption of previously 
prescribed treatment, while 28% were purchased on the 
patient’s initiative (28); this could result in inappropriate 
or ineffective use of medication and consequently an 
unintentional non-adherence to recommendations. 
Thus, the importance of patient education cannot be 
over-emphasized. Moreover, to meet today’s demands for 
accessible and efficient care, it is important to develop 
new methods, such as eHealth strategies, to facilitate and 
promote partnership between health professionals and 
patients (29).

The majority of participants relied on family members 
to fill the gaps in carrying out tasks they once did alone; 
this is contrary to findings of other research showing that 
COPD participants tended to rely primarily on themselves 
to avoid burdening their relatives and close friends (22). 
This is explained by social relations and support from 
family as well as friends which is considered “the axis of 
Lebanese values, beliefs and culture” (30). The Lebanese 
population still has relatively strong family networks, 
and cultural ideals continue to support intergenerational 
co-residence (31). According to Lebanese cultural norms, 
older people generally have great expectations of help 
from their children and family members. Caregivers in 
this study had no previous knowledge or experience of 
disease management, decision-making during 
complications, or interpersonal challenges, which is in 
line with the findings of other studies that emphasize 
the importance of involving caregiver in any strategies 
tailored for COPD patients and as contributors partnering 
with health care professionals (32,33).

As in many other countries, Lebanon experiences 
economic inequalities in health; it offers few benefits 
in terms of welfare, health care and pensions. Economic 
problems were widely discussed during the interviews 
because they are barriers to therapeutic adherence 
and medical consultation. Although other American 
and British studies also described the financial worries 
of COPD participants concerning funding their care 
(34,35), policies to protect older adults in Lebanon are 
extremely weak in comparison (9). Considering the lack 
of formal government support, increasing attention by 
nongovernmental agencies is helping to address 
unmet needs for health care (9). Therefore, health care
professionals should help participants to identify available resources that are best for them.

**Methodological considerations**

Some authors claim that credibility, authenticity, criticality and integrity remain the main criteria of scientificity in qualitative research (36). Credibility is respected when the research results truly describe the phenomenon. Thus, it was our intention to recruit a large group of participants with diverse socioeconomic characteristics and to conduct interviews beyond data redundancy. Moreover, all prior assumptions of the authors about living with COPD were set aside throughout the process of data collection and analysis.

Authenticity is used to clarify whether the results correspond to the experience as described by each of the participants. Therefore, we read the transcripts of the interviews multiple times; we made audio and video recordings of the interviews and conducted a second visit to validate our interpretation of the transcripts. Audio recordings allowed us to gather information from patients intimidated by the presence of cameras, especially those who thought that the disease had an impact on their physical image.

Criticality refers to the constant critical attitude of the researcher in order to avoid bias. We followed the Giorgi methodological framework (10) throughout data collection and analysis.

Integrity illustrates the researcher’s concern to validate his or her interpretations based on the data, therefore we avoided prematurely formulating the essence of the phenomenon by achieving redundancy before ending data collection and making a second visit.

However, the interviews were conducted with patients living in only three governorates of Lebanon: Beirut, Mount Lebanon and South Lebanon. Excluding patients living in other more rural districts may have limited the findings of this study.

**Conclusion**

Living with COPD means living with vulnerability and uncertainty influenced by the body’s capacities and the economic challenges of a country with limited resources, which imposes on patients a need for support to cope with daily life. Therefore, participants turn to their families in a cultural context where the family continues to be seen as the primary provider of support. The results of our study show that COPD affected three dimensions of participants’ lives. These were the educational dimension, where the lack of information was a need and a factor that influenced the COPD patients’ experience with their disease; the coordination dimension as patients were obliged to reorganize their lives according to the Lebanese context (financial, social); and the psychosocial dimension given the impact of the disease on social life, professional life, physical image and emotional well-being as expressed by the participants. Therefore, therapeutic education has been recognized as a need for COPD patients in Lebanon. In addition, the development of interventions that include family caregivers and the cooperation of all health professionals is vital. Furthermore, to respond to patients’ demands for easy access to care, it is important to develop new methods of facilitating and promoting the partnership between health professionals and patients while preserving patient comfort. Thus governmental and health care efforts are needed to improve the experiences of patients with COPD in Lebanon and to improve their quality of life.

**Acknowledgement**

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**Competing interests:** None declared.

**Vivre avec une bronchopneumopathie obstructive chronique au Liban : une étude phénoménologique**

**Résumé**

**Contexte:** Le Liban est le pays de la région arabe qui connaît la croissance la plus rapide de sa population de personnes âgées, mais il dispose de peu de ressources sociales pour répondre à leurs besoins. Aucune étude n’a examiné l’expérience des patients atteints de bronchopneumopathie obstructive chronique au Liban.

**Objectifs:** Examiner les expériences des personnes vivant avec une bronchopneumopathie obstructive chronique au Liban.
المصابون بمرض الانسداد الرئوي المزمن في لبنان: دراسة الظواهر
رنا جورج نهرا، جان مانويل مورفييرز، هالة صقر، باسكال سلامة، مونيك روثان-توندور

الخلاصة
يسجل لبنان أسرع معدل نمو للسكان من كبار السن في المنطقة العربية، وذلك في ظل قلة موارده الاجتماعية لتنبيها هذه الفئة. ولم تتناول أي دراسات تجربة مرضى الانسداد الرئوي المزمن ومرضهم في لبنان.

الأهداف: هدفت هذه الدراسة إلى استكشاف تجارب الأفراد المصابين بمرض الانسداد الرئوي المزمن في لبنان.

مетодة: À l'aide d'un modèle de recherche phénoménologique descriptive, des entretiens qualitatifs individuels semi-structurés ont été menés auprès de personnes atteintes de bronchopneumopathie obstructive chronique vivant au Liban, entre mai et septembre 2019.

نتائج: Cinquante participants ont accepté d'être interrogés. La majorité étaient des hommes (56 %) et avaient une bronchopneumopathie obstructive chronique modérée (40 %). L'âge moyen était de 71,5 ans (écart type 9,0). Nous avons constaté que la bronchopneumopathie obstructive chronique a une incidence sur trois aspects de la vie des patients: éducatif, organisationnel et psychosocial.

استنتاجات: تؤكد النتائج على الحاجة إلى استراتيجيات متعددة التخصصات للمصابين بمرض الانسداد الرئوي المزمن في لبنان، مع تضمن القانونين على رعايتهم. وتشمل الاستراتيجيات تثقيف المرضى وتطوير أساليب جديدة لتسهيل الشراكة وتعزيزها بين مهني الرعاية الصحية ومرضى الانسداد الرئوي المزمن والقائمين على رعايتهم.

References


