Severe Asthma in adults does not significantly affect the outcome of COVID-19 disease: results from the Italian Severe Asthma Registry

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June 28, 2020

To the Editor,

Severe asthma (SA) is a chronic disease affecting around 3-8% of adult asthma population in Europe, with the refractory form estimated to occur in 0.1% of the general population (1,2). SA is characterized by increased use of healthcare resources (i.e. emergency room/hospital admissions, access to intensive care units (ICU), use of biologics) due to exacerbations compared to the less severe form. In the current SARS-CoV-2 pandemic, there is an ongoing debate on the role of asthma and use of immunomodulating drugs, like corticosteroids and biologics, on COVID-19 outcomes. According to available data on COVID-19 hospitalizations, asthma seems to play little role on the clinical severity or access to health resources, unlike other chronic conditions such as hypertension, obesity and chronic obstructive pulmonary disease (3). However, to date, no information is available on the burden of SA on COVID-19 severity and hospitalization rates.

A questionnaire was submitted to the Italian Registry of Severe Asthma (IRSA) network (4), assessing the prevalence and clinical characteristics of patients with SA who contracted COVID-19 during the outbreak in Italy (February 24th - May 18th 2020), and 41 out of 78 centers distributed evenly among different Italian regions participated to the survey (Figure 1a).

Among the 558 subjects surveyed, 7 subjects contracted COVID-19 (1.25% of the national sample), with an average age of 54.5 years: 5 isolated at home/received home care (71.5%), while 2 subjects were admitted to the hospital (28.5%), none required accessed to ICU and no deaths were reported. All COVID-19 subjects with SA came from 2 regions of Northern Italy (6 Lombardy, 1 Emilia-Romagna, 3.7% of the regional population), all showing one or more comorbidities, and were treated with high-dose inhaled corticosteroids plus long-acting beta-2 agonists (ICS-LABA) and biologics (see Table 1).
We then compared our results with data provided by the Italian Department for Civil Protection in the same time period from the affected geographic areas (5), and we observed that the frequency of COVID-19 among subjects referred to IRSA centers strongly correlated with the prevalence of SARS-CoV-2 infection in the corresponding province (Figure 1b). Furthermore, the hospitalization rate in COVID-19-SA subjects was not significantly different from the general population (24.1%, 23.6-24.6 95% C.I.; p=0.25, Chi-squared test). Lastly, we could not observe a significantly increased COVID-19 frequency in subjects undergoing high-dose ICS-LABA and biologics compared to SA treated with ICS-LABA alone (p=0.09, Fisher exact test).

These findings from the IRSA registry offer some insights on the susceptibility to SARS-CoV-2 infection, access to healthcare resources and mortality by SA patients.

Given the low prevalence of SA in Italy (2), we expected less COVID-19-SA cases per region than what reported by the IRSA survey. However, we observed that the geographic location of COVID-19-SA patients mostly reflected the bimodal distribution of the COVID-19 outbreak in Italy, mainly clustered in Lombardy and neighboring regions, where the highest cumulative COVID-19 cases were recorded (>500/100000 cases per inhabitants) (5). In these areas, the prevalence of positive cases by province also strongly correlated with the frequency of COVID-19-SA patients observed in each IRSA center (Figure 1b), suggesting that patients with SA most likely contract the infection when high circulation of the virus within the area of residence is present. The lack of positive cases reported in Southern regions further proves this hypothesis, and demonstrates the efficacy of the lockdown measures adopted to contain the further spread of the virus.

Our results also suggest no increased risk of contracting COVID-19 in SA treated with biologics compared to ICS-LABA alone. Although there is currently no strong evidence that biologics used in asthma might affect the risk of contracting COVID-19, new evidence suggests a protective effect of inhaled corticosteroids against viral entry by ACE2 receptor downregulation, that are usually prescribed at a high dose in SA (6), thus a possible explanation to the lack of observed differences in our cohort.

Despite the severity of asthma and reported comorbidities, no ICU admissions were reported, and hospital admissions in COVID-19-SA subjects did not differ from the median rate observed in the same geographic areas (5). Furthermore, we could observe no difference in the median monthly hospitalization rate of SA patients in 2019 compared to 2020 in Lombardy region where both hospital-admitted subjects reside (0.97 vs 0.9%, IRSA data).

Our result is consistent with recent literature, showing that asthma in Western countries was not associated with an increased hospitalization rate and ICU admissions due to COVID-19 (3,8). It is still debated if a protective effect of Th2-inflammation in a significant proportion of asthma sufferers (7), or concomitant anti-inflammatory therapy could be the reasons for such outcomes (6). However, if asthma patients with COVID-19 require intubation, the duration of hospitalization was shown to be longer than average (8).

As for the role of biologics in COVID-19 disease progression, we could not observe an increase in hospital admissions in patients with SA treated with biologics compared to the general population, with the majority isolating at home and requiring no additional treatment. Considering that, in areas with high prevalence of SARS-CoV-2 infection, 68.2% of SA subjects were treated with either omalizumab or mepolizumab, our observations further prove the safety of biologics during the COVID-19 pandemic.

Lastly, we did not observe any deaths in our cohort, but we speculate that this outcome is most likely due to the small sample size and younger average age. In fact, advanced age seems to be the most determining risk factor on mortality due to COVID-19 compared to other causes. (9)

Taken together, our results point at a neutral role of SA in the COVID-19 disease course and hospital admissions. One major strengths of our study is that, by using a fast and inexpensive tool, we could outline the salient features of severe asthma and COVID-19 at a national level, while the major weakness is the limited number of SA subjects diagnosed with COVID-19, that could lead to sampling bias and low accuracy. Further confirmation of these results with an increased sample size is therefore warranted.

Conclusion paragraph
Despite the clinical burden of severe asthma is substantial, there is evidence of a neutral effect of severe asthma in the clinical progression and hospitalization due to COVID-19 in a cohort of Italian severe asthma patients. Treatment with biologics for severe asthma also seems to have no significant effect on the outcome of COVID-19.

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Acknowledgements

The authors wish to thank Centro Studi AIPO-ITS for its invaluable support in the generation of the data.

Funding sources

The Italian Registry for Severe Asthma (IRSA) is co-funded by the Italian Thoracic Society (ITS - AIPO) and the Italian Association of Territorial and Hospital Allergists and Immunologists (AAIITO).

Conflict of interests statement

LA was an invited speaker by Novartis, GlaxoSmithKline, Astra Zeneca, Sanofi Genzyme, Chiesi Group and Menarini.

LR reports fees as invited speaker by Chiesi Group and GlaxoSmithKline.

FB was an invited speaker by Chiesi Group, Astra Zeneca, Novartis, Menarini, Roche and Boehringer Ingelheim.

FDM declares fees as speaker/lecturer and Advisor Board by AstraZeneca, GlaxoSmithKline.

AM declares Advisory Board fees from AstraZeneca, GlaxoSmithKline, Mylan, Sanofi Genzyme.

CM was an invited speaker by Novartis, GlaxoSmithKline, AstraZeneca, Sanofi, Chiesi Group, Menarini, Guidotti, Zambon and Berlin Chemie.

MBB declares receiving fees as invited speaker/lecturer by GlaxoSmithKline.

CT, GM, AV, AS, FM have no conflict of interest to disclose.

Author contribution statement

LA and CM designed the study; FDM, AM, CM, MBB (IRSA scientific board) approved the study and MBB was appointed supervisor; GM, LR, AV, FB, AS and FM collected information on COVID-19 patients; LA and CT performed the statistical analyses and wrote the manuscript; all authors revised and contributed to the final version of the manuscript.

Data availability statement

The data that support the findings of this study are available from the corresponding author upon reasonable request.

Table 1 – Clinical characteristics of subjects with severe asthma and COVID-19 (n = 7)

| Demographic data | Demographic data | Demographic data | Demographic data | Demographic data | Demographic data | Demographic data |
|------------------|------------------|------------------|------------------|------------------|------------------|------------------|
| IRSA center id.  | Region           | Province         | Pt. ID           | Sex             | Age             |
| 25               | E-R              | R. Emilia        | 1                | F               | 48               |
| 40               | Lombardy         | Bergamo          | 2                | F               | 65               |
| 46               | Lombardy         | Milan            | 3                | F               | 40               |
| 102              | Lombardy         | Cremona          | 4                | F               | 51               |
| 114              | Lombardy         | Brescia          | 5                | F               | 57               |
| 114              | Lombardy         | Brescia          | 6                | M               | 65               |
| 114              | Lombardy         | Brescia          | 7                | F               | 56               |
| Average          | -                | -                | -                | -               | 54.6             |
| SD               | -                | -                | -                | -               | 8.4              |

Abbreviations: BMI, body mass index; E-R, Emilia-Romagna; GERD, gastroesophageal reflux disease; ICS,
inhaled corticosteroids; ICU, intensive care unit; IRSA, Italian Registry of Severe Asthma; LABA, long-acting beta-2 agonists; LAMA, long-acting muscarinic antagonists; OCS, oral corticosteroids; SD, standard deviation.

FIGURE CAPTION

Figure 1. (a) Distribution of IRSA Centers with number of surveyed subjects per region (left) and prevalence of COVID-19 cases per inhabitants per region (adapted from data of the Department of Civil Protection, 18/05/2020 report) (right); (b) Correlation between frequency of COVID-19 cases per IRSA Center and COVID-19 prevalence per province.