Hospital acquired infections in COVID-19 patients in sub intensive care unit: analysis of two waves of admissions

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Abstract. Background and aim: The pandemic caused by SARS-COV-2 has increased Semi-Intensive Care Unit (SICU) admission, causing an increase in healthcare-associated infection (HAI). Mostly HAI reveals the same risk factors, but fewer studies have analyzed the possibility of multiple coinfections in these patients. The study aimed was to identify patterns of co-presence of different species describing at the same time the association between such patterns and patient demographics and, finally, comparing the patterns between the two cohorts of COVID-19 patients admitted at Policlinico during the first wave and the second one. Methods: All the patients admitted to SICUs during two COVID-19 waves, from March to June 2020 months and from October to December 2020, were screened following the local infection control surveillance program; whoever manifested fever has undergone on microbiological culture to detect bacterial species. Statistical analysis was performed to observe the existence of microbiological patterns through DBSCAN method. Results: 246 patients were investigated and 83 patients were considered in our study because they presented infection symptoms with a mean age of 67 years and 33.7% of female patients. During the first and second waves were found respectively 10 and 8 bacterial clusters with no difference regarding the most frequent species. Conclusions: The results show the importance of an analysis which considers the risk factors for the possibility of co- and superinfection (such as age and gender) to structure a good prognostic tool to predict which patients will encounter severe coinfections during hospitalization (www.actabiomedica.it)

Key words: COVID-19, Hospital Acquired Infection, Subintensive Unit, DBSCAN

Introduction

The pandemic of Severe Acute Respiratory Syndrome Coronavirus 2 (SARS-CoV-2) has and had caused an unprecedented medical crisis for all health services all over the world. In Italy, at the end of December 2020, there were 2.107.166 confirmed cases and 74.159 deaths overall; as of the 26th of June 2022, the confirmed cases are 18.071.634 and deaths 167.967. Almost 10% of patients with COVID-19 experimented hospital admission and 9% of them needed to stay in intensive care.
units but with substantial differences over the Italian
territory (1,2).

To face this huge request for intense use of hos-
pitals the health management boards had to organize
hospital areas devoted to the cure of COVID-19 pa-
tients (3). The COVID-19 patients admitted to in-
tensive or sub-intensive care units (ICUs/SICUs) to
the gravity of their conditions, like all the other pa-
tients in the hospital, faced also a considerable threat
to their safety caused by healthcare-associated infec-
tions (HAIs) which might determine adverse clinical
outcomes and aggressive antimicrobial therapies with
further resistance selection (4-10).

Both kinds of patients could present HAIs caused
by different bacteria, likely associated with ventila-
tors, invasive ventilation, and the usage of empirical
broad-spectrum antimicrobials (11). Many researchers
had investigated the problem of bacterial and fungal
infection in COVID-19 patients admitted to ICUs
but there are very few studies devoted to studying
COVID-19 patients admitted to SICUs (12).

In IRCCS Fondazione Ca' Granda Policlinico di
Milano, a research and teaching hospital located in the
center of Milan with active 716 beds, 84 of them were
turned up in sub-intensive care to admit patients who
needed less intensive care in both the two waves of the
epidemic. In total, during the first wave, from March
9th to June 6th, 2020, 246 patients were admitted to
these 84 beds for sub-intensive care and 80 of whom
perished (9). At the beginning of October started
the second period of very high pressure to admit
COVID-19 patients to the hospital, the second wave,
and the same SICUs were used with the same num-
ber of beds (84), and from October 15th to December
15th, 2020 were admitted 172 patients and 78 died
(administrative data from hospital records).

This study aims to evaluate, in two cohorts of
patients resulted positive to microbiological culture
examination (83 over 246 patients admitted during
the first wave, and 73 over 172 during the second
wave) in the IRCCS Fondazione Ca’ Granda Poli-
clinico di Milano, the prevalence of infections by
different species of bacteria, to identify patterns of
co-presence of different species, to describe the as-
association between such patterns and patient demo-
graphics (gender, age).

Data about the first and second waves were an-
alyzed separately and compared to consider the dif-
ferent features of the outbreak waves (13). For these
patients are also available the results of the local
infection control surveillance program to detect col-
onization by multidrug-resistant bacteria, namely
MRSA (Methicillin-resistant Staphylococcus aureus),
multidrug-resistant Gram-negative bacteria, and VRE
(Vancomycin-resistant enterococci).

**Patients and methods**

During 2020, 418 patients were admitted to our
SICUs, 246 during the first wave from March to June
and 172 during the second wave from October to De-
cember. All patients admitted to SICUs during the first
and second waves were routinely followed with the pro-
cedure of the local infection control surveillance program
to detect colonization by multidrug-resistant bacteria,
noticed MRSA (Methicillin-resistant Staphylococcus
aureus), multidrug-resistant Gram-negative bacteria
and VRE (Vancomycin-resistant enterococci) in add-
diction received all the microbiological investigation in
case of infectious symptoms. An infection control sur-
veillance program was performed through oral-nasal
and rectal swabs on all hospitalized patients respec-
tively to detect MRSA and multidrug-resistant Gram-
negative bacteria (also considering MDR enterococci).
Since fever could be considered one of the most im-
portant infection signs, in any patient manifesting fever
were performed microbiological cultures to detect spe-
cies of bacterium and antimicrobial resistance; depend-
ing on reported symptoms such as urgency or dysuria,
shortness of breath or chest pain, urine, sputum, and
blood culture were performed, through a non-selective
agar media for culture. No specific data about micro-
biological load were available. Microbiological cultures
were performed on 83 and 73 patients during the first
wave and the second one respectively.

**Statistical methods**

According to the main goals of the analysis, i.e.,
to evaluate the prevalence and co-presences of the
species of bacterium detected in the patients enrolled in the first wave (N=83) and the patients enrolled in the second wave (N=73), data were included in two distinct datasets and structured as follows. In the first wave, a total of 30 species of bacteria were detected (at least in one patient). Accordingly, 30 variables coded as presence/absence (0=absence, 1=presence) of the species above were included in the dataset. The same was done for the 23 species of bacteria detected in the second wave.

For each species of bacterium, the prevalence was defined as the percentage of infected over the total amount of patients. Since patients were submitted to several laboratory investigations, we also reported, for each species of bacterium, the percentage of positive diagnoses over the total number of investigations.

To evaluate the patterns of co-presence of different species, cluster analysis methods were used.

In presence of many species of bacteria with relatively very low prevalence (<2,5%), density-based clustering methods were considered. In particular the DBSCAN (14) was chosen, because of its appreciable performances in dealing with subjects with very uncommon coinfection patterns. Furthermore, for the reasons above, the clusters are not likely to show spherical shapes: therefore, DBSCAN should be preferred to other common clustering algorithms, such as K-means.

DBSCAN is based on the indexes of dissimilarity between subjects. For identifying subjects in the same cluster, the algorithm identifies, for each subject, the subjects who are more similar to him; this set is called “neighborhood”. To define the neighborhood two parameters are needed: the parameter $\varepsilon$ (called the “radius” of the neighborhood) and MinPoints (the minimum number of subjects in the neighborhood).

In this work, similarities were calculated according to two indexes specific for presence/absence data: namely, the Jaccard index and the Dice-Sorensen index. These indexes are based on the number of species of bacterium shared by two subjects. If two subjects have no species in common both the indexes are equal to 0, and when they share the same species, both indexes are equal to 1. In the remaining cases, the values of the two indices are different, with the Dice-Sorensen index giving more weight to the number of common species (15). It is worth noting that both these indices were considered in this work because there is no general agreement on which one performs better.

To choose the values of the parameters $\varepsilon$ and MinPoints, the procedure described by Schubert et al and the silhouette method were adopted (16,17). In particular, the chosen parameter values were those values that maximized the compactness of the clusters according to the silhouette.

The age of subjects within each cluster was described using a dotplot and gender by reporting the percentage of females within each cluster.

All the analyses were performed using the software R release 3.6.2 and package “dbscan” (18,19).

Results

Analysis of the first wave

Two hundred and forty-six patients were investigated during the first wave and 83 patients were considered in our study because they presented infection symptoms. The mean age of these patients was 67 years and ranged from 39 to 89 and female patients were 28 (33,7%).

The most frequent positive materials sent in microbiological laboratories were blood from peripheral blood vessels (107/284, 37,7%) and blood from venous catheters (80/284 – 28,2%). The most frequent investigation was blood culture (205/284 – 72,2%). According to the results of microbic cultures, 30 species of bacteria were found; the counts and proportions of patients affected by each species are reported in Table 1. The most frequent bacterium isolated was *Staphylococcus epidermidis* (39/284 – 13,7%) and *Staphylococcus aureus* (53/284 – 18,7%).

In cluster analysis, the results obtained adopting the Jaccard index and those obtained with the Dice-Sorensen index showed very few differences. Therefore, only the results obtained with the latter one are reported below. According to the silhouette method, the best values of $\varepsilon$ and MinPoints parameters were 0,20, and 3, respectively.

Ten clusters were identified, which altogether include 50 (60,2%) of the 83 subjects considered (Table 2). In six clusters, subjects were infected only by one species (cluster numbers: 1, 2, 5, 6, 8, and 9).
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Clusters 3 and 6 present narrower ranges: from 63 to 76 years for cluster 3, and 78 to 86 years for cluster 6. Cluster 10 is composed of relatively younger patients, whose age ranges from 43 to 69 years. The highest percentages (about 67%) were found in clusters 1, 3, and 9, whereas in clusters 7 and 10 all subjects were male.

In two clusters (clusters 3 and 7) subjects presented at most three species of bacterium. In cluster 4, at most 4 species can “co-exist”, and in cluster 10 at most 5 species can be found in a single patient.

Figure 1 presents the distribution of age and gender of patients within each cluster. Concerning age, clusters 3 and 6 present narrower ranges: from 63 to 76 years for cluster 3, and 78 to 86 years for cluster 6. Cluster 10 is composed of relatively younger patients, whose age ranges from 43 to 69 years. The highest percentages (about 67%) were found in clusters 1, 3, and 9, whereas in clusters 7 and 10 all subjects were male.

### Table 1. Infections of positive diagnoses detected during the first wave.

| Species of bacteria (abbreviation) | Infected patients: (N=83) | Positive diagnoses (N=284) |
|-----------------------------------|--------------------------|---------------------------|
| Staphylococcus epidermidis (S.ep) | 21 (25,3%)               | 39 (13,7%)                |
| Escherichia coli (E.co)           | 21 (25,3%)               | 28 (9,9%)                 |
| Enterococcus faecalis (E.fa)      | 20 (24,1%)               | 39 (13,7%)                |
| Staphylococcus aureus (S.au)      | 11 (13,2%)               | 53 (18,7%)                |
| Enterococcus faecium (E.fa1)      | 11 (13,2%)               | 18 (6,3%)                 |
| Staphylococcus hominis (S.ho)     | 10 (12,0%)               | 18 (6,3%)                 |
| Staphylococcus haemolyticus (S.ha)| 9 (10,8%)                | 11 (3,9%)                 |
| Klebsiella pneumoniae (K.pn)     | 8 (9,6%)                 | 18 (6,3%)                 |
| Proteus mirabilis (P.mi)         | 5 (6,0%)                 | 9 (3,2%)                  |
| Pseudomonas aeruginosa (P.ae)     | 4 (4,8%)                 | 10 (3,5%)                 |
| Bacillus clausii (B.cl)          | 4 (4,8%)                 | 5 (1,8%)                  |
| Enterobacter cloacae (E.cl)       | 2 (2,4%)                 | 2 (0,7%)                  |
| Morganella morganii (M.mo)        | 2 (2,4%)                 | 2 (0,7%)                  |
| Providencia stuartii (P.st)       | 2 (2,4%)                 | 2 (0,7%)                  |
| Stafilococco auro Meticillino Resitente (S.au.Res) | 1 (1,2%) | 9 (3,2%) |
| Serratia marcescens (S.ma)        | 1 (1,2%)                 | 3 (1,1%)                  |
| Corynebacterium urealyticum (C.ur)| 1 (1,2%) | 2 (0,7%) |
| Enterobacter aerogenes (E.ae)     | 1 (1,2%)                 | 2 (0,7%)                  |
| Staphylococcus capitis (S.ca)     | 1 (1,2%)                 | 2 (0,7%)                  |
| Staphylococcus cohnii (S.co)      | 1 (1,2%)                 | 2 (0,7%)                  |
| Aerococcus viridans (A.vi)        | 1 (1,2%)                 | 1 (0,3%)                  |
| coco-bacilli Gram positivi (Cc.pos) | 1 (1,2%) | 1 (0,3%) |
| Corynebacterium amycolatum (C.am) | 1 (1,2%) | 1 (0,3%) |
| Corynebacterium striatum (C.st)   | 1 (1,2%)                 | 1 (0,3%)                  |
| K. pneumoniae resistente ai carbapenemi (K.pn.res) | 1 (1,2%) | 1 (0,3%) |
| Stafilococco aures Meticillino Sensibile (S.au.sen) | 1 (1,2%) | 1 (0,3%) |
| Staphylococcus pettenkoferi (S.pe) | 1 (1,2%) | 1 (0,3%) |
| Stenotrophomonas maltophilia (S.ma1) | 1 (1,2%) | 1 (0,3%) |
| Streptococcus parasanguis (S.pa)  | 1 (1,2%)                 | 1 (0,3%)                  |
| Str. beta emol. Gr.F (Str.be)     | 1 (1,2%)                 | 1 (0,3%)                  |

Legend: For each species of bacteria: the prevalence (percentage of infection) was defined as the proportion of infected patients over the total number of patients (n=83); the percentage of positive diagnoses was calculated by the proportion of positive diagnoses over the total number of laboratory investigations (n=284).
### Table 2. Description of clusters of patients in the first wave.

| Cluster | Size | bacteria: species label (n) |
|---------|------|---------------------------|
| 1       | 6    | E.fa (6)                  |
| 2       | 14   | E.co (14)                 |
| 3       | 3    | E.fa (2), E.co (3), B.cl (2) |
| 4       | 3    | E.fa (3), S.ep (3), E.fa1(3), S.pa (1) |
| 5       | 6    | S.ho (6)                  |
| 6       | 3    | K.pn (3)                  |
| 7       | 3    | S.ep (3), S.au (3), S.au.res (1) |
| 8       | 3    | S.au (3)                  |
| 9       | 5    | S.ep (5)                  |
| 10      | 4    | E.fa (4), E.co (1), S.ep(4), S.ho (1), S.ha (4) |

Legend: size = cluster size, i.e. the number of subjects included in the cluster. In the third column, there is reported for each cluster, the description of species of bacterium: ‘species name’ is the abbreviation of the species, and n is the number of subjects within the cluster that present the species. E.fa= Enterococcus faecalis, E.co= Escherichia coli, B.cl= Bacillus clausii, S.ep= Staphylococcus epidermidis, E.fa1= Enterococcus faecium, S.pa= Streptococcus parasanguis, S.ho= Staphylococcus hominis, K.pn= Klebsiella pneumoniae, S.au= Staphylococcus aureus, S.au.res= Staphilococco aureo Meticillino Resistent, S.ha= Staphylococcus haemolyticus

Left panel: age. The blue line is the mean age in the total sample; the red triangles are the mean age for each cluster. Black dots: age of each subject. Right panel: gender. The blue line is the percentage of females in the total sample; the grey bars are the percentage of females for each cluster.

### Analysis of the second wave

All 172 patients were investigated during the second wave and 73 patients were considered in our study because they presented infection symptoms. Their age ranged between 42 and 96 years, with an average value of 73.4 years; female patients were 22 (30,1%).

The most frequent positive material sent in the microbiological laboratory was blood from peripheral veins (76/197 – 38.6%) and consequently, the most frequent investigation was blood culture (150/197 – 76.1%). The most frequent bacterium isolated was *Staphylococcus epidermidis* (42/197 – 21.3%).

Table 3 presents, for each species of bacteria, the number and percentage of infected subjects, and
clusters of the eight above were formed by subjects infected by only one species (cluster numbers: 1, 2, 3, 4, 6, 7, and 8). In one cluster (cluster 5) subjects presented at most four species of bacterium. Figure 2 presents the distribution of age and gender of patients within each cluster. Clusters 4 and 8 are formed by relatively older subjects: in fact, the age ranges from 75 to 88 in cluster 4, and from 73 to 84 years in cluster 8. The percentage of females in the sample was 69.9%. The highest percentage (100%) was

Concerning cluster analysis, according to the silhouette methods, the best values of $\varepsilon$ and MinPoints parameters were 0.20, and 3, respectively. Eight clusters were identified, which altogether include 39 of the 73 subjects considered (53.4%). Table 4 describes the species detected in the subject for each cluster. Seven

### Table 3. Amount of infections and of positive diagnoses detected during the second wave.

| Species of bacteria (abbreviation) | Infected patients: (n = 73) | Positive diagnoses (N=197) |
|-----------------------------------|-----------------------------|-----------------------------|
| Staphylococcus epidermidis (S.ep)  | 19 (26.1%)                  | 42 (21.3%)                  |
| Staphylococcus hominis (S.ho)     | 12 (16.4%)                  | 16 (8.1%)                   |
| Enterococcus faecalis (E.fa)      | 11 (15.1%)                  | 13 (6.6%)                   |
| Staphylococcus aureus (S.au)      | 10 (13.7%)                  | 23 (11.7%)                  |
| Enterococcus faecium (E.fa1)      | 9 (12.3%)                   | 23 (11.7%)                  |
| Escherichia coli (E.co)           | 9 (12.3%)                   | 13 (6.6%)                   |
| Pseudomonas aeruginosa (P.ae)     | 7 (9.6%)                    | 16 (8.1%)                   |
| Staphylococcus haemolyticus (S.ha)| 6 (8.2%)                    | 11 (5.6%)                   |
| Bacillus clausii (B.cl)           | 5 (6.8%)                    | 6 (3.0%)                    |
| Klebsiella pneumoniae (K.pn)      | 4 (5.5%)                    | 4 (2.0%)                    |
| Staphylococcus capitis (S.ca)     | 4 (5.5%)                    | 4 (2.0%)                    |
| Enterobacter hormaechei (E.ho)    | 2 (2.7%)                    | 8 (4.1%)                    |
| Enterobacter aerogenes (E.ae)     | 2 (2.7%)                    | 2 (1.0%)                    |
| Acinetobacter baumannii (A.ba)    | 2 (2.7%)                    | 2 (1.0%)                    |
| Actinomycetum haemolyticum (A.sc) | 1 (1.4%)                    | 2 (1.0%)                    |
| Proteus mirabilis (P.mi)          | 1 (1.4%)                    | 2 (1.0%)                    |
| Aerococcus urinae (A.ur)          | 1 (1.4%)                    | 1 (0.5%)                    |
| Candida albicans (C.al)           | 1 (1.4%)                    | 1 (0.5%)                    |
| Corynebacterium species (C.sx)    | 1 (1.4%)                    | 1 (0.5%)                    |
| Corynebacterium spp (C.spp)       | 1 (1.4%)                    | 1 (0.5%)                    |
| Corynebacterium striatum (C.st)   | 1 (1.4%)                    | 1 (0.5%)                    |
| Hafnia alvei (H.al)               | 1 (1.4%)                    | 1 (0.5%)                    |
| Staphilococcus aureus Meticillino Sensibile (S.au.sen) | 1 (1.4%) | 1 (0.5%) |
| Staphylococcus schleiferi (S.sc)  | 1 (1.4%)                    | 1 (0.5%)                    |
| Streptococcus mitis/oralis (S.m.o) | 1 (1.4%)                  | 1 (0.5%)                    |
| Streptococcus parasanguis (S.pa)  | 1 (1.4%)                    | 1 (0.5%)                    |

Legend: For each species of bacteria: the percentage of infection was defined as the proportion of infected patients over the total number of patients (n=73); the percentage of positive diagnoses was calculated by the proportion of positive diagnoses over the total number of laboratory investigations (N=197).
found in cluster 3, whereas the lowest one was found in cluster 6.

Left panel: age. The blue line is the mean age in the total sample; the red triangles are the mean age for each cluster. Black dots: age of each subject. Right panel: gender. The blue line is the percentage of females in the total sample; the grey bars are the percentage of females for each cluster.

Bacterial co-infections within the two waves

Table 5 presents the distribution of the patients according to the number of bacterial species found in the respective microbiologic cultures. Interestingly the majority of patients (respectively 57.8% and 61.6%) were affected by only one species of bacterium in both waves.

Conclusions

Since its first appearance, Sars-CoV-2 is considered a public health issue, not only due to its infectivity and mortality rate but also to the increase in hospital admission. Indeed, severe respiratory infections, such as COVID-19, have increased the risk of admission to semi-intensive and intensive care units, especially for immunocompromised patients and patients with co-pathologies.

At the same time, several studies have investigated the role of respiratory infections in hospital-acquired infection (HAI) showing the link between the

![Figure 2. Mean of age and percentage of females in clusters about the second wave.](image-url)
disruption of airways structure such as distortion of mucus secretion, cell death, lung edema, decreased mucosal clearance, reduced oxygen exchange through disrupted angiogenesis, and impaired surfactant secretion (20,21). These pathophysiological impairments and the increase in hospitalization, associated with combined therapy based on antibiotics and corticosteroids as suggested by several studies (22-24), affected the HAI rates with an increase not only in mortality and morbidity but also hospitalization, enhancing the length and the economic burden on the healthcare system (25).

In this setting, microbiological investigations assume a central role in the diagnostic workout of hospitalized patients. Microbiological investigations were performed on each patient who has been admitted to sub-intensive care wards for COVID-19 and who presented fever as an additional symptom. Among the 418 patients that were hospitalized in sub-intensive care wards during the year 2020, 156 were enrolled in the study due to at least one microbiological isolation.

To search for a specific pattern of the infection our team decided to describe coinfections clusters in semi-intensive care units during the first and second waves analyzing it also regarding patients’ demographic characteristics.

During the first wave of the pandemic, data from 83 patients were analyzed. The most represented microbiological investigation was blood culture, an inquiry routinely used in the diagnostic workup of persistent fever. S. aureus, a well-known hospital-acquired pathogen, was the commonest bacteria, along with coagulase-negative staphylococci (CoNS), species often considered contaminants. E. faecium and E. coli were among the most frequent, representing 13.7% and 9.8% respectively. Almost 60% of the patients studied had only one species of bacteria isolated from the samples collected.

Analyzing demographic characteristics of different clusters, it’s interesting to note as can’t be observed a specific link between the number of isolated species and cluster mean age or gender prevalence, setting out that the increase of species for a cluster could be linked to the specificity of a single patient rather than a demographic characteristic.

Considering the second wave, data collected, from 73 patients, confirmed that blood culture was the most frequent microbiological investigation performed. S. epidermidis was the commonest bacteria isolated, with S. aureus and E. faecium sharing the second place. Interestingly, P. aeruginosa, another known hospital-acquired pathogen, was the third most represented bacteria isolated accounting for 8% of all the isolated germs. Similarly, to the first wave, a little more than 60% of the patients showed only one detectable species of bacteria. As previously view for the first wave, the analysis of demographic characteristics for the second wave doesn’t show a specific link between the number of isolated species and demographic characteristics.

It’s interesting to note regarding isolate species as, considering the six most common isolates species they are the same in both waves, and S. epidermidis represents the commonest one, likely due to several invasive devices in sub-intensive care units (26).

These results emphasize the importance of an overall analysis of a single patient to avoid antimicrobial therapy mistakes based on non-specific characteristics that could invalidate antimicrobial-stewardship programs implemented in high-risk HAI areas, such as the emergency department or infectious disease ward.

As previously described, several studies have analyzed co- and superinfection in COVID-19 hospitalized patients and it’s interesting to note as none of the studies analyzed has considered the possibility of co-infection clusters simply describing the coinfection rates regardless of the number of different microbiological species isolate (27,28,29,30,31).

**Table 5. Number of species of bacteria found in patients.**

| Number of species | First wave: frequency (%) | Second wave: frequency (%) |
|-------------------|---------------------------|---------------------------|
| 1                 | 48 (57,8%)                | 45 (61,6%)                |
| 2                 | 18 (21,7%)                | 19 (26,0%)                |
| 3                 | 9 (10,8%)                 | 7 (9,6%)                  |
| 4                 | 6 (7,2%)                  | 1 (1,4%)                  |
| 5                 | 1 (1,2%)                  | 0 (0,0%)                  |
| 6                 | 1 (1,2%)                  | 1 (1,4%)                  |
| TOT               | 83                        | 73                        |
It’s evident from several studies that COVID-19 patients are an HAI high-susceptibility patient category during hospitalization and in HAI patients the number of microbiological species is directly linked to the severity of prognosis. The lack of studies that analyze co-infection clusters highlights the importance of our study as the first one exploring this topic. This study has several limitations; firstly, the retrospective design reduces control over especially data collection, and secondly, the study was limited to a single hospital despite it being carried out through two COVID-19 waves amounting to 150 days. In conclusion, the importance of HAI prevention, especially during the COVID-19 pandemic, highlights the importance of a tool to detect which colonized patients may develop infections to improve a specific prophylaxis procedure.

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