# RESEARCH



# Epidemiology, ventilation management and outcomes of COVID–19 ARDS patients versus patients with ARDS due to pneumonia in the Pre–COVID era

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# Abstract

**Background** Ventilation management may differ between COVID–19 ARDS (COVID–ARDS) patients and patients with pre–COVID ARDS (CLASSIC–ARDS); it is uncertain whether associations of ventilation management with outcomes for CLASSIC–ARDS also exist in COVID–ARDS.

**Methods** Individual patient data analysis of COVID–ARDS and CLASSIC–ARDS patients in six observational studies of ventilation, four in the COVID–19 pandemic and two pre–pandemic. Descriptive statistics were used to compare epidemiology and ventilation characteristics. The primary endpoint were key ventilation parameters; other outcomes included mortality and ventilator–free days and alive (VFD–60) at day 60.

**Results** This analysis included 6702 COVID–ARDS patients and 1415 CLASSIC–ARDS patients. COVID–ARDS patients received lower median V<sub>T</sub> (6.6 [6.0 to 7.4] vs 7.3 [6.4 to 8.5] ml/kg PBW; p < 0.001) and higher median PEEP (12.0 [10.0 to 14.0] vs 8.0 [6.0 to 10.0] cm H<sub>2</sub>O; p < 0.001), at lower median  $\Delta P$  (13.0 [10.0 to 15.0] vs 16.0 [IQR 12.0 to 20.0] cm H<sub>2</sub>O; p < 0.001) and higher median Crs (33.5 [26.6 to 42.1] vs 28.1 [21.6 to 38.4] mL/cm H<sub>2</sub>O; p < 0.001). Following multivariable adjustment, higher  $\Delta P$  had an independent association with higher 60–day mortality and less VFD–60 in both groups. Higher PEEP had an association with less VFD–60, but only in COVID–ARDS patients.

**Conclusions** Our findings show important differences in key ventilation parameters and associations thereof with outcomes between COVID–ARDS and CLASSIC–ARDS.

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# Background

The high numbers of patients who needed invasive ventilation early in the unprecedented pandemic of coronavirus disease 2019 (COVID–19) has led to numerous studies of epidemiology, ventilation management and outcomes in patients with acute respiratory distress syndrome (ARDS) related to an infection with SARS–CoV– 2. COVID–19 ARDS would differ from ARDS before the pandemic (CLASSIC–ARDS) in several aspects [1, 2], and different phenotypes have even been suggested [3, 4].

The number of studies that directly compared ventilation management of COVID–ARDS with CLASSIC– ARDS is limited [5, 6]. It remains uncertain whether practice of invasive ventilation in COVID–ARDS patients really differed from that in CLASSIC–ARDS patients. It is also unknown whether associations of certain aspects of ventilation with outcomes found in CLAS-SIC–ARDS also exist in COVID–ARDS. This would have serious implications on how to set the ventilator in the two patient groups, as then certain recommendations in guidelines for ventilation in CLASSIC–ARDS may not apply in COVID–ARDS [7].

We performed an analysis of a conveniently-sized database that pooled the data of individual patients of six observational ventilation studies, four of which were conducted in the COVID-19 pandemic and two pre-pandemic, to compare epidemiology, ventilator management and associations of ventilation characteristics and outcome between COVID-ARDS and CLASSIC-ARDS patients. To have comparable patient groups, we only selected patients with ARDS from a respiratory infection from the two pre-pandemic studies. We hypothesized that key ventilator parameters would be different between the two groups, and used multivariable analyses to determine associations with outcomes.

# Methods

## Study design and participants

This is a meta–analysis using the individual patient data of patients in six preselected large observational studies focusing on a diverse representation of epidemiological features and ventilation management in both COVID–19 and pre–pandemic ARDS. The six studies were selected because they all contained detailed data on epidemiological features, ventilation data, and outcomes, originating from various regions worldwide, both in resource–limited and resource–rich settings. The corresponding authors of the original studies accepted the invitation, after which the data dictionaries of the studies were compared to check whether the data could be harmonized. Then, the databases were transferred after local approval and agreement on the analysis plan of the current investigation.

The two pre-pandemic studies were the national 'Epidemiology of Respiratory Insufficiency in Critical Care' study (ERICC) conducted in 2011 in Brazil [8], and the international 'Large Observational Study to UNderstand the Global Impact of Severe Acute Respiratory FailurE' study (LUNG SAFE) conducted in 2014 in 50 countries worldwide [9]. All four studies were conducted during the COVID-19 pandemic, ranging from March 2020 to 2021 and included: the national 'Practice of Ventilation in COVID-19 patients' study (PRoVENT-COVID) from The Netherlands [10], the national 'EPIdemiology of Critical COVID-19' study (EPICCoV) from Brazil [11, 12], the national 'Centro de Investigación Biomédica en Red Enfermedades Respiratorias COVID-19 study' (CIBERESUCICOVID) from Spain [13], and the national 'Sociedad Argentina de Terapia Intensiva-COVID-19 study' (SATI-COVID-19) from Argentina [14].

The study protocols of the original studies were approved by Institutional Review Boards if applicable, and need for individual patient informed consent was waived for all studies due to their observational designs. Details of all studies can be found in the original publications [8-10, 12-14]. We invited the corresponding investigators of the original studies to provide us the case report forms and data dictionaries, and the data of all patients. The creation of the pooled database did not require additional ethical approval. The databases of the original studies were harmonised using the case report forms and data dictionaries, and finally merged. This current analysis is registered at clinicaltrials.gov (study identifier NCT05650957), and its statistical analysis plan was finalized before cleaning and closing of the database.

Patients in the merged database were eligible for participation in this current analysis if: (1) aged 18 years or higher; (2) having received invasive ventilation within the first 48 h of ICU admission, regardless of its duration; and (3) fulfilling the Berlin definition of ARDS. We excluded CLASSIC-ARDS patients when ARDS was reported not to be caused by a respiratory infection.

## Data available for merging

The following baseline and demographic variables were available for merging into the new database—sex, age, body weight and height, comorbidities including hypertension and cardiac failure, chronic obstructive pulmonary disease (COPD), diabetes mellitus, kidney failure, liver failure, and cancer, date of hospital and intensive care unit (ICU) admission, and disease severity scores, including the Simplified Acute Physiology Score (SAPS) II at ICU admission and a daily Sequential Organ Failure Assessment (SOFA) scores.

Collected ventilation variables were—mode of ventilation, tidal volume ( $V_T$ ), positive end—expiratory pressure (PEEP), fraction of inspired oxygen (FiO<sub>2</sub>), respiratory rate (RR), peak pressure (Ppeak) in volume–controlled ventilation and plateau pressure (Pplat) in pressure– controlled ventilation, blood gas analyses results, and adjunctive therapies to improve oxygenation in case of refractory hypoxaemia. The first available measurement of the day was used. If multiple measurements were taken on the same day, we selected earliest one.

The dynamic driving pressure ( $\Delta P$ ) was calculated by subtracting PEEP from the maximum airway pressure [15, 16]. Respiratory system compliance (Crs) was calculated by dividing  $V_T$  by  $\Delta P$ . MP was calculated using the power Eq. (17), wherein MP  $(J/min) = 0.098 * V_T *$ RR \* (Ppeak – 0.5 \*  $\Delta P$ ) [17]; a modified power equation was used if no Ppeak was available 0.098 \*  $V_T$  \* RR \* (Pplat – 0.5 \*  $\Delta P$ ) [16]. The ventilatory ratio was calculated as (minute ventilation \* PaCO<sub>2</sub>)/(predicted bodyweight \* 100 \* 37.5) [18]. The number of ventilator-free days at day 60 (VFD-60) was calculated by subtracting the number of calendar days a patient received invasive ventilation up to the day of successful extubation from 60, similar to the method used for calculating VFD-28. Patients that died before or at day 60 received zero VFD-60 [19, 20].

The following follow-up data were available for merging—last day of ventilation, tracheostomy use, last day in ICU and hospital, and life status at day 60.

#### Endpoints

The primary endpoint of this analysis was a combination of the following key ventilation characteristics as done before [10]— $V_T$ , PEEP,  $\Delta P$ , and Crs. Secondary outcomes were other ventilator parameters, the use of prone positioning, muscle paralysis or extracorporeal membrane oxygenation, and 60–day mortality and the number of VFD–60.

#### **Power analysis**

We did not perform a formal power analysis; instead, the number of available patients served as the sample size.

#### Statistical analysis

Baseline demographics were compared using Fisher's exact tests for categorical variables and Wilcoxon rank–sum tests for continuous variables. Continuous distributed variables are presented as medians and interquartile ranges, categorical variables are presented as frequencies and proportions.

The first day a patient received invasive ventilation and the first full calendar day were combined into 'day 1', the next day was designated as 'day 2'. Information on missing values for each ventilation parameters and other variable can be found in the Supplementary Material (eTable 1). Only SOFA scores were available for all patients, therefore, we chose to only report these instead of other severity scores.

To compare ventilation characteristics between COVID–ARDS and CLASSIC–ARDS patients, a Wilcoxon rank–sum test was used. Cumulative distribution frequencies of each ventilation variable or parameter, wherein vertical dotted lines represent broadly accepted safety cutoffs for each variable, and horizontal dotted lines show the respective proportion of patients reaching that cutoff.

As a post-hoc analysis to identify whether  $V_T$ , PEEP and  $\Delta P$  have independent associations with 60-day mortality and the number of VFD-60, a multivariable mixed-effects model with centre as random effect was performed. A linear mixed-effects model was used for the number of VFD-60 and a logistic mixed-effects model for 60-day mortality.

The following covariates, with a known or suspected association with these two outcomes were included in the model, based on clinical relevance: (1)  $PaO_2/FiO_2$ ; and (2) demographic variables, including sex, age, BMI, history of heart failure, COPD, diabetes mellitus, kidney failure, liver failure and cancer.

In this mixed model analysis, when a covariate exhibited more than 10% missing data, we utilized multiple imputation techniques implemented through the MICE package in R. The model was checked for collinearity using variance–inflation factors, wherein a variance– inflation factor <5 was deemed acceptable. The variance– inflation factor was <2 for all included variables in our model.

The estimate refers to the average effect of the ventilation parameter, i.e.,  $V_T$ , PEEP or  $\Delta P$  on the outcome of interest, i.e., 60–day mortality and VFD–60 while controlling for the other variables in the model. A positive estimate indicates that an increase in the predictor variable tends to lead to a corresponding increase in the response variable, indicating a proportional relationship between them. Conversely, a negative estimate suggests that an increase in the predictor variable tends to result in a decrease in the response variable, indicating an inverse proportional relationship between them.

All analyses were conducted in R v.4.0.3 (R Foundation for Statistical Computing, Vienna, Austria). A p value < 0.05 was considered statistically significant.

# Results

We received the individual data of a total of 8374 COVID–ARDS patients and 3795 CLASSIC–ARDS patients (Fig. 1). After exclusion of patients that did not fulfil the Berlin definition of ARDS, patients that did not receive invasive ventilation on the first and second day in the study, and patients included in the two pre–pandemic studies who did not have a respiratory infection as the cause for ARDS, we had 6702 fully–analysable COVID– ARDS patients and 1415 fully–analysable CLASSIC– ARDS. COVID–ARDS patients were more often male, had higher median BMI, a history of diabetes more often, and a history of COPD or chronic kidney disease less often (Table 1). COVID–ARDS patients had lower median SOFA scores, and ARDS severity was more often classified as moderate or severe.

COVID–ARDS patients were ventilated with volume– controlled ventilation more often than CLASSIC–ARDS patients (Table 2) and received ventilation with lower V<sub>T</sub> (6.6 [6.0 to 7.4] vs 7.3 [6.4 to 8.5] ml/kg PBW; p < 0.001), higher PEEP (12.0 [10.0 to 14.0] vs 8.0 [6.0 to 10.0] cm H<sub>2</sub>O; p < 0.001), at lower  $\Delta P$  (13.0 [10.0 to 15.0] vs 16.0 [IQR 12.0 to 20.0] cm H<sub>2</sub>O; p < 0.001) and higher Crs (33.5 [26.6 to 42.1] vs 28.1 [21.6 to 38.4] mL/cm H<sub>2</sub>O; p < 0.001) (Fig. 2) COVID–ARDS patients received higher PEEP than CLASSIC–ARDS patients at any FiO<sub>2</sub> level (eFigure 2). Within each group, the ventilation characteristics were not different between day 1 and 2 (eTable 2 and eFigure 1 and 2).

Prone positioning and neuromuscular blocking agents were more often used in COVID–ARDS patients than in CLASSIC–ARDS patients (Table 2). COVID–ARDS patients received a tracheostomy more often than CLAS-SIC–ARDS patients.

Mortality at day 60 was higher in COVID–ARDS patients compared to CLASSIC–ARDS patients (Table 2 and Fig. 3), and COVID–ARDS patients had significantly less VFD–60. Following multivariable adjustment, higher  $\Delta P$  had an association with higher 60–day mortality and less VFD–60 in both groups. Higher PEEP also had an association with less VFD–60, but only in COVID–ARDS patients and not in CLASSIC–ARDS patients. In both groups, V<sub>T</sub> neither had an association with 60–day mortality nor with VFD–60 (eFigure 3 and eFigure 4).

# Discussion

We pooled the individual data of patients from six observational studies of ventilation and compared ventilation characteristics and associations with outcomes between COVID–ARDS with CLASSIC–ARDS. The main findings were: (1) compared to CLASSIC–ARDS patients, COVID–ARDS patients were ventilated with lower V<sub>T</sub> and higher PEEP, at lower  $\Delta P$  and higher Crs, however with a higher MP; (2) 60–day mortality was not different between COVID–ARDS and CLASSIC–ARDS, but COVID–ARDS patients had less VFD–60; (3) higher  $\Delta P$  had an association with higher 60–day mortality and less VFD–60 in COVID–ARDS and CLASSIC–ARDS; and



Fig. 1 Flowchart of included studies. Abbreviations: ARDS = acute respiratory distress syndrome; COVID-19 = coronavirus disease 2019

	COVID-ARDS	CLASSIC-ARDS	р
	N=0.702	N=1.415	
Demographics			
age, years, median [IQR]	64.0 [55.0 to 71.0]	63.0 [50.0 to 74.0]	0.361
height, cm, median [IQR]	170.0 [163.0 to 175.0]	168.0 [160.0 to 175.0]	< 0.001
weight, kg, median [IQR]	82.0 [74.0 to 95.0]	74.0 [62.0 to 86.0]	< 0.001
male gender, n/N (%)	4.655 (69.5)	859 (60.7)	< 0.001
BMI, kg/m <sup>2</sup> , median [IQR]	28.5 [25.6 to 32.4]	25.8 [22.5 to 30.1]	< 0.001
SOFA score, median [IQR]	6.0 [4.0 to 8.0]	10.0 [7.0 to 12.0]	< 0.001
Comorbidities			
heart failure, N (%)	633 (9.4)	143 (10.1)	0.472
COPD, N (%)	688 (10.3)	323 (22.8)	< 0.001
diabetes, N (%)	1.906 (28.4)	322 (22.8)	< 0.001
chronic kidney disease, N (%)	402 (6.0)	140 (9.9)	< 0.001
liver failure, N (%)	122 (1.8)	53 (3.7)	< 0.001
active neoplasm, N (%)	253 (3.8)	119 (8.4)	< 0.001
ARDS severity categories			< 0.001
mild, N (%)	1.937 (28.9)	362 (25.6)	
moderate, N (%)	3.477 (51.8)	673 (47.6)	
severe, N (%)	1.288 (19.2)	380 (26.9)	

**Table 1** Patient demographics, Baseline Characteristics and ARDS Severity

Abbreviations: ARDS Acute respiratory distress syndrome, IQR Interquartile range, N Number, BMI Body mass Index, SOFA Sequential organ failure assessment, COPD Chronic obstructive pulmonary disease

(4) higher PEEP also had an association with less VFD– 60, but only in COVID–ARDS.

Our findings add to the current understanding of differences and similarities between COVID-19 ARDS patients and pre-COVID ARDS patients. The international design of our study increases the generalizability of the findings across diverse healthcare systems, both in ARDS patients caused by COVID-19 and in patients with ARDS due to pneumonia from before the pandemic. The large sample size and high quality of the collected data allowed for sophisticated analyses of epidemiology, respiratory support strategies, and outcomes. Additionally, we found associations between key ventilator settings and patient outcomes.

Several studies have compared COVID–19 ARDS with pre–COVID ARDS. The epidemiological differences between COVID–19 ARDS and pre–COVID ARDS patients in our study align with previous findings [21]. As with other studies [22, 23], we also found significant differences in ventilator variables like  $V_T$ , PEEP, and  $\Delta P$ , and in the use of adjunctive therapies. Our study contributes by demonstrating these differences specifically among ARDS patients and comparing COVID–19 ARDS to pre–COVID ARDS due to respiratory infections. Differences in outcomes found in our study are, at least in part, in line with prior research findings [21, 23]. Our findings confirm that there are differences in mortality and the number of VFD–60 between COVID–19 ARDS and

pre–COVID ARDS patients. However, these difference disappeared after propensity matching. This is important as it shows that, at least when comparing outcomes in ARDS patients from an infectious cause, outcomes are not different, opposite to what was thought at the start of the pandemic.

We observed more frequent use of lower  $V_T$  in COVID-ARDS compared to CLASSIC-ARDS. Indeed, proportions of COVID-ARDS patients that received ventilation with a  $V_T < 6$  or between 6 and 8 ml/kg PBW was higher than in CLASSIC-ARDS patients. This finding can be explained in several ways--e.g., it could be that the use of lung-protective ventilation with a lower  $V_T$  has improved in the last decade [15]. It is also conceivable that, at least early in the pandemic care for COVID-ARDS patients was provided by inexperienced ICU staff which could have been more adherent to existing guidelines for management of patients with ARDS [10, 24]. It is also possible that use of low  $V_T$  in COVID–ARDS is easier to control--these patients were often deeply sedated and paralyzed allowing a stricter adherence to lower  $V_{T}$ . Of note, especially in those patients, ventilation with a lower  $V_T$  might be more beneficial than in spontaneous breathing patients [25].

Higher PEEP was more often used in COVID–ARDS patients than in CLASSIC–ARDS patients, at any  $FiO_2$  level. Indeed, proportions of COVID–ARDS patients that received ventilation with a PEEP between 8 and 12

	$\frac{\text{COVID}-\text{ARDS}}{N=6.702}$	CLASSIC-ARDS N=1.415	p
Ventilation characteristics			
mode of ventilation, N (%)			< 0.001
volume-controlled ventilation	4.689 (70.3)	521 (36.8)	
pressure-controlled ventilation	1.403 (21.0)	493 (34.8)	
pressure-support ventilation	336 (5.0)	143 (10.1)	
other	242 (3.6)	258 (18.2)	
V <sub>T</sub> , mL/kg PBW, median [IQR]	6.6 [6.0 to 7.4]	7.3 [6.4 to 8.5]	< 0.001
<6 ml/kg PBW	1.464 (24.8)	211 (16.3)	< 0.001
6–8 ml/kg PBW	3.601 (61.2)	645 (50.0)	< 0.001
8–10 ml/kg PBW	698(11.9)	324 (25.1)	< 0.001
> 10 ml/kg PBW	125 (2.1)	102 (7.9)	< 0.001
PEEP, cmH <sub>2</sub> O, median [IQR]	12.0 [10.0 to 14.0]	8.0 [6.0 to 10.0]	< 0.001
$< 8 \text{ cmH}_2\text{O}$	325 (4.8)	530 (37.5)	< 0.001
8–12 cmH <sub>2</sub> O	4.075 (60.8)	713 (50.4)	< 0.001
$12-16 \text{ cmH}_{2}\text{O}$	2.101 (31.3)	140 (9.9)	< 0.001
> 16 cmH <sub>2</sub> O	201 (2.9)	32 (2.3)	0.37
$P_{MAX}$ , cmH <sub>2</sub> O, median [IQR]	25.5 [21.0 to 30.0]	25.0 [22.0 to 28.0]	0.001
<b>dynamic</b> $\Delta P$ , cmH <sub>2</sub> O, median [IQR]	13.0 [10.0 to 15.0]	16.0 [12.0 to 20.0]	< 0.001
C <sub>Rs</sub> , mL/cmH <sub>2</sub> O, median [IQR]	33.5 [26.6 to 42.1]	28.1 [21.6 to 38.4]	< 0.001
MP, J/min, median [IQR]	16.6 [13.4 to 20.6]	15.2 [11.3 to 19.2]	< 0.001
FiO <sub>2</sub> median [IQR]	0.6 [0.5 to 0.9]	0.6 [0.5 to 0.9]	0.017
total RR, breaths per min, median [IQR]	22.0 [20.0 to 25.0]	20.0 [16.0 to 25.0]	< 0.001
ventilatory ratio, median [IQR]	1.75 [1.43 to 2.20]	1.76 [1.35 to 2.27]	0.835
Adjunctive therapies			
prone positioning, N (%)	4.615 (69.2)	144 (10.2)	< 0.001
recruitment manoeuvres, N (%)	1.924 (39.4)	313 (22.1)	< 0.001
ECMO, N (%)	142 (2.5)	56 (4.0)	0.005
tracheostomy, N (%)	2.033 (30.5)	214 (15.1)	< 0.001
neuromuscular blocking agents, N (%)	3.838 (73.9)	350 (24.7)	< 0.001
continuous sedation, N (%)	1.188 (84.0)	1.785 (98.1)	< 0.001
vasopressor use, N (%)	4440 (85.5)	1025 (72.5)	< 0.001
Arterial blood gas analysis			
pH, median [IQR]	7.35 [7.28 to 7.41]	7.34 [7.26 to 7.41]	< 0.001
PaO <sub>2</sub> , mmHq, median [IQR]	80.1 [67.5 to 98.5]	83.6 [68.0 to 105.8]	< 0.001
PaCO <sub>2</sub> , mmHg, median [IQR]	44.25 [38.0 to 52.0]	44.0 [37.0 to 54.0]	0.320
PaO <sub>2</sub> /FiO <sub>2</sub> , median [IQR]	135.0 [94.0 to 184.9]	146.0 [99.4 to 207.3]	< 0.001
Outcomes		- •	
60-day mortality, N (%)	2963 (44.2)	515 (36.4)	< 0.001
VFD–60, median [IQR]	11.0 [0.0 to 47.0]	41.0 [0.0 to 54.0]	< 0.001

## Table 2 Ventilation Characteristics, Adjunctive Therapies, Arterial Blood Gas Analysis and Outcomes

Abbreviations: ARDS Acute respiratory distress syndrome,  $V_T$  Tidal volume; PBW Predicted bodyweight, IQR Interquartile range, N Number, PEEP Positive end–expiratory pressure,  $P_{max}$  Maximum airway pressure,  $\Delta P$  Driving pressure, MP Mechanical power,  $C_{RS}$  Respiratory system compliance, FiO<sub>2</sub> Fraction of inspired oxygen, RR Respiratory rate, PaO<sub>2</sub> Partial pressure of arterial oxygen, PaCO<sub>2</sub> Partial pressure of arterial carbon dioxide, VFD Ventilator–free days and alive

 $\rm cmH_2O$  and even between 12 and 16  $\rm cmH_2O$  was higher than in CLASSIC–ARDS patients. This finding can also be explained in several ways––e.g., a preference for use of higher PEEP in COVID–ARDS patients may have been triggered by the severity of ARDS, as COVID–ARDS was more often classified as moderate or severe, and more severe hypoxaemia naturally triggers the use of higher PEEP if PEEP/FiO<sub>2</sub> tables are used. It is also possible that higher PEEP was used in the assumption that lung lesions with COVID–ARDS are more recruitable than in CLAS-SIC–ARDS. This may at least explain the lower  $\Delta P$  and higher Crs in COVID–ARDS patients.



**Fig. 2** Key ventilation parameters. Cumulative frequency distribution of  $V_T$ , PEEP,  $\Delta P$ , and respiratory system compliance on the first calendar day for each variable. Vertical dotted lines represent broadly accepted safety cutoffs for each variable, and horizontal dotted lines show the respective proportion of patients reaching that cutoff. Abbreviations:  $V_T$ =tidal volume; PBW=predicted bodyweight; PEEP=positive end-expiratory pressure;  $\Delta P$ =driving pressure;  $C_{PS}$ =respiratory system compliance

In COVID–ARDS patients, mechanical power exceeded that of CLASSIC–ARDS, even though the driving pressure was lower. This observation marks the significance of considering factors beyond driving pressure, such as respiratory rate and PEEP, when evaluating the protective nature of invasive ventilation. These findings emphasize the complexity of respiratory management in COVID–ARDS and the need for a comprehensive approach to optimize lungprotective ventilation strategies.

COVID-ARDS patients received prone positioning more often than CLASSIC-ARDS patients. Before the

pandemic, prone positioning remained underused, probably because it was more considered a rescue therapy for refractory hypoxaemia [26]. While we cannot rule out that use of prone positioning increased already before the pandemic, we favour the idea that the higher use of prone positioning in COVID–ARDS patients was triggered by the more severe hypoxaemia in COVID–ARDS patients.

Our analysis found several associations between ventilation parameters and outcome. The association of higher  $\Delta P$  with higher 60–day mortality and less VFD– 60 is in line with previous studies [27–29]. The association of higher PEEP with worse outcome confirms the

Parameter	Estimate (95%–CI)	р	Favours Lower	Favours Higher	
VFD-60			VFD	VFD-60	
V⊤, mL/kg PBW				1	
COVID-ARDS	0.1124 (-0.3773 – 0.6034)	0.63		• • •	
Classic-ARDS	0.2409 (-0.4789 – 1.0879)	0.45			
PEEP, cm H <sub>2</sub> O				1	
COVID-ARDS	-0.2613 (-0.4638 – -0.0575)	0.01		i	
Classic-ARDS	-0.0082 (-0.4417 –0.4244)	0.97		•	
Dynamic ∆P, cm H	<sub>2</sub> O				
COVID-ARDS	-0.3985 (-0.5516 – -0.2454)	< 0.001		i l	
Classic-ARDS	-0.5910 (-0.90510.2767)	0.002		1	
			-1.0 -0.5	0.0 0.5 1.0 days	
60–day mortality			60–day	mortality	
V⊤, mL/kg PBW				1	
COVID-ARDS	0.0008 (-0.0096 - 0.0112)	0.87	_	<b>←</b>	
Classic–ARDS	0.0004 (-0.1362 - 0.0145)	0.95		• I	

Classic–ARDS	0.0004 (-0.1362 - 0.0145)	0.95
PEEP, cm H <sub>2</sub> O		
COVID-ARDS	0.0017 (-0.0027 – 0.0059)	0.45
Classic–ARDS	-0.0061 (-0.0138 – 0.0015)	0.12
Dynamic ΔP, cm I	H <sub>2</sub> O	
COVID-ARDS	0.0077 (0.0044 - 0.0110)	<0.001
Classic–ARDS	0.0119 (0.0061 – 0.0176)	0.002



Fig. 3 Mortality and ventilator-free days and Alive at day-60, and associations with ventilator parameters. The estimate is the average effect of the predictor variable on the response variable, while controlling for the other variables in the model. A positive estimate suggests a proportional effect, whereas a negative estimate suggests an inversely proportional effect. Abbreviations: ARDS = acute respiratory distress syndrome; VFD = ventilator-free days and alive; IQR = interguartile range; N = number; CI = confidence interval;  $V_{\tau}$  = tidal volume; PBW = predicted bodyweight; PEEP = positive end-expiratory pressure;  $\Delta P$  = driving pressure

findings of earlier studies [30, 31]. Of note, this association was only found for COVID-ARDS. This may have been caused by the more frequent use of higher PEEP in COVID-ARDS than in CLASSIC-ARDS. One reason for the association between higher PEEP and worse outcome may be that sicker patients, with a higher chance of dying and prolonged ventilation, received higher PEEP than patients that were less sick. Nonetheless, a high PEEP is suggested to have detrimental effects [32], emphasizing the need to determine the optimal PEEP level based on lung recruitability rather than hypoxemia alone. Actually, one analysis of PRoVENT-COVID suggested worse outcomes if patients received ventilation according to a higher PEEP/lower FiO<sub>2</sub> table as compared to ventilation according to a lower PEEP/higher FiO<sub>2</sub> [30]. A post-hoc Bayesian analysis of a randomised clinical study, named the 'Alveolar Recruitment for ARDS Trial' (ART), wherein patients were randomized to receive ventilation with PEEP titrated to the best Crs and aggressive recruitment manoeuvres versus ventilation with a low PEEP strategy, suggested that higher PEEP with recruitment manoeuvres worsens the outcome of ARDS from pneumonia, while it may be beneficial in ARDS from another cause [33]. A posthoc analysis of a randomised clinical study named 'Lung Imaging for Ventilator Setting in ARDS trial' (LIFE), suggest that higher PEEP worsens outcomes in patients with ARDS with lesions that may not be recruitable with higher PEEP [34].

The findings of this pooled analysis extend the existing knowledge of the epidemiology, management of invasive ventilation and outcomes in COVID-ARDS. Our study shows that lung-protective ventilation was applied well in COVID-ARDS, and was comparable to best practice used in management for patients with CLASSIC-ARDS. Additionally, the effect of PEEP on major outcomes may have implications for care. At least it should trigger new studies that directly compare different PEEP strategies. Meanwhile, it could be more attractive to not use higher PEEP by default.

Our study has several strengths. We managed to receive and merge the datasets of four large observational studies of ventilation conducted in the COVID-19 pandemic with two well-performed pre-pandemic observational studies of ventilation--these six studies all focused on ventilation management and reported outcomes of invasively ventilated ARDS patients, allowing a robust

analysis of ventilation management and the impact of certain ventilation parameters on outcome. While the COVID-19 studies were all national investigations, they are from different regions worldwide and were conducted in different types of hospitals, which increases the generalizability of our findings. The datasets from the original studies were rich and comprehensive, encompassing baseline and demographic data, granular ventilator settings and ventilation variables, and key clinical outcomes. All data could be harmonized and merged into one database.

We had an analysis plan in place before cleaning and closing of the new database, and this plan was strictly followed. The large numbers of patients allowed us to perform sophisticated statistical analyses of associations with outcomes.

This study has limitations. First, individual data was obtained from observational studies, which limits the ability to establish causality. Additionally, the willingness of data sharing could have led to selection bias towards the inclusion of ICUs with an interest in invasive ventilation and management of ARDS in the original studies. Second, studies in COVID-ARDS were conducted early in the COVID-19 pandemic, during which inexperienced staff and resource limitations could have influenced clinical decision making. Third, data was collected early in the pandemic when patient care took priority over data collection, resulting in more missing data than in previous studies. This affects the completeness and may impact the accuracy of our analysis. Fourth, we only reported on ventilation characteristics on day 1 and 2, because not all studies collected ventilation data beyond this timepoint. Therefore we were not able to compare ventilation management beyond day 2. Nevertheless, previous studies have shown ventilation characteristics don't significantly change in the first four days after initiation of invasive ventilation [10]. Fifth, it is imperative to acknowledge the temporal distance between comparator cohorts. For the pre-COVID ARDS group we used patients of which data was collected between seven to nine years before the pandemic. We cannot exclude temporal differences, for instance due to studies that showed the importance of limiting liberal use of oxygen, and reducing the intensity of ventilation, e.g., by targeting a low driving pressure or a low mechanical power of ventilation, as well as the importance of early use of prone positioning. Sixth, is the lack of detailed subgroup analyses, particularly in patients with chronic respiratory comorbidities such as COPD. Although recent findings from a post-hoc analysis of the PRoVENT-COVID study by Tripipitsiriwat et al. [35] indicated that ventilation parameters did not show significant differences between COPD and non-COPD patients, it could be interesting to explore these subgroups. However, it was beyond the scope of our primary endpoint. Conducting such detailed subgroup investigations would require careful consideration to ensure the data from all included studies are appropriate for this type of analysis.

Finally, all COVID–19 ARDS patients, by definition, had a viral pneumonia, while patients in the classic ARDS group had respiratory infections of which the pathogen was not collected. This is an important limitation, as ARDS from a viral respiratory infection may differ from ARDS due to bacterial pneumonia. Consistent with other studies comparing COVID–19 ARDS to ARDS caused by other viruses, we found that the duration of ventilation was longer, and mortality was higher [21, 36, 37].

# Conclusions

Epidemiology and key ventilation characteristics were different in patients with COVID–ARDS compared to CLASSIC–ARDS, also  $\Delta P$  was lower in COVID–ARDS patients.  $\Delta P$  had an independent association with outcome in both groups, whereas PEEP had an independent association with outcome only in COVID–ARDS patients.

#### Supplementary Information

The online version contains supplementary material available at https://doi. org/10.1186/s12931-024-02910-2.

Supplementary Material 1

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for the ERICC<sup>3</sup>-,LUNG SAFE<sup>b</sup>-, PROVENT-COVID<sup>c</sup>-, EPICCoV<sup>d</sup>-, CIBERESUCICOVID<sup>e</sup>- and SATI-COVID-19<sup>f</sup>-investigators <sup>a</sup>ERICC, 'Epidemiology of Respiratory Insufficiency in Critical Care' <sup>b</sup>LUNG SAFE, 'Large Observational Study to UNderstand the Global Impact of Severe Acute Respiratory FailurE' <sup>c</sup>PROVENT-COVID, 'Practice of Ventilation in COVID-19 patients' <sup>d</sup>EPICCoV, EPIdemiology of Critical COVID-19 <sup>e</sup>CIBERESUCICOVID, 'Centro de Investigación Biomédica en Red Enfermedades Respiratorias COVID-19' <sup>f</sup>SATI-COVID-19, 'Sociedad Argentina de Terapia Intensiva-COVID-19' **Authors' contributions** 

Author contribution: FV, SB, MS, FP and DM had full access to all the data and take responsibility for the integrity of the data and the accuracy of the data analysis. Concept and design: All authors Acquisition, analysis, or interpretation of data: FV, SB, MS, FP and DM Drafting of the manuscript: FV, SB, MS, FP and DM Critical revision of the manuscript for important intellectual content: All authors Statistical analysis and data verification: FV, SB, and DM Obtained funding: Not applicable; the original studies were performed with funding as stated in the original reports. Administrative, technical, or material support: LA, GB, MB, EE, JF, JL, TP, AT Supervision: MS, FP and DM.

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#### Availability of data and materials

Data sharing: A de-identified dataset can be made available upon request to the corresponding authors one year after publication of this study, but

only after permission of the principal investigators of all original studies. The request must include a statistical analysis plan.

#### Declarations

#### Ethics approval and consent to participate

The creation of the pooled database did not require additional ethical approval.

#### **Consent for publication**

Not applicable.

#### **Competing interests**

The authors declare no competing interests.

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