Late mortality after acute hypoxic respiratory failure


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Late mortality after acute hypoxic respiratory failure

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ABSTRACT
Background Acute hypoxic respiratory failure (AHRF) is associated with significant acute mortality. It is unclear whether later mortality is predominantly driven by pre-existing comorbid disease, the acute inciting event or is the result of AHRF itself.

Methods Observational cohort study of elderly US Health and Retirement Study (HRS) participants in fee-for-service Medicare (1998–2012). Patients hospitalised with AHRF were matched 1:1:1 to otherwise similar adults who were not currently hospitalised and separately to patients hospitalised with acute inciting events (pneumonia, non-pulmonary infection, aspiration, trauma, pancreatitis) that may result in AHRF; here termed at-risk hospitalisations. The primary outcome was late mortality—death in the 31 days to 2 years following hospital admission.

Results Among 15075 HRS participants, we identified 1268 AHRF and 13117 at-risk hospitalisations. AHRF hospitalisations were matched to 1157 non-hospitalised adults and 1017 at-risk hospitalisations. Among patients who survived at least 30 days, AHRF was associated with a 24.4% (95% CI 19.9% to 28.9%, p<0.001) absolute increase in late mortality relative to adults not currently hospitalised and a 6.7% (95% CI 1.7% to 11.7%, p=0.01) increase relative to adults hospitalised with acute inciting event(s) alone. At-risk hospitalisation explained 71.2% of the increased odds of late mortality, whereas the development of AHRF itself explained 28.8%. Risk for death was equivalent to at-risk hospitalisation beyond 90 days, but remained elevated for more than 1 year compared with non-hospitalised controls.

Conclusions In this national sample of older Americans, approximately one in four survivors with AHRF had a late death not explained by pre-AHRF health status. More than 70% of this increased risk was associated with hospitalisation for acute inciting events, while 30% was associated with hypoxemic respiratory failure.

INTRODUCTION
Acute hypoxic respiratory failure (AHRF) is a leading cause of intensive care unit admission1,2 with high acute mortality.1 However, the impact of AHRF on late mortality (death in the 31 days to 2 years after hospital admission) is unclear. Studies with longer-term follow-up indicate that patients continue to die at a high rates after surviving hospitalisation for acute respiratory distress syndrome (ARDS, a severe form of AHRF),3,4 but without a comparison cohort, we cannot determine whether this late mortality is the result of pre-existing comorbid disease versus a direct effect of AHRF. An older study suggests that patients surviving ARDS do not have increased risk of late mortality relative to patients surviving hospitalisation for inciting events alone—but was not powered to detect effects smaller than a 14% absolute difference.5 With declining in-hospital mortality,6-8 AHRF-related mortality may now be shifted to the posthospital period. Other critical illnesses, such as sepsis9 and delirium,10 are associated with increased late mortality. Thus, we hypothesised that AHRF may also increase patients’ risk for late mortality.

Understanding whether patients who survive AHRF acutely face increased risk for late death is important for (1) measuring the overall burden of AHRF-related mortality, (2) determining whether AHRF is associated with prolonged pathobiological derangements that may be amenable to treatment, (3) informing the timing of interventions to improve long-term survivorship after AHRF, (4) selecting clinical trial end-points and (5) understanding the prognosis of patient surviving AHRF.

To address these questions, we compared patients hospitalised with AHRF to two controls: adults not currently hospitalised and patients hospitalised with acute inciting events that may result in AHRF—pneumonia, non-pulmonary infection, aspiration, trauma, pancreatitis.
aspiration, trauma or pancreatitis. These comparisons allow us to determine both (1) the excess late mortality of an episode of AHRF and (2) the incremental excess late mortality of AHFR, above and beyond what is associated with hospitalisation for acute inciting events.

METHODS

Study population

We studied participants in the US Health and Retirement Study (HRS), a longitudinal cohort of 37,000 adults aged 51 and above in 23,000 households, ongoing since 1992.11 The study uses a multistage probability sample to identify participants.11 The sociodemographic and racial distribution is broadly representative of the older US population.12–14 The cohort is interviewed every 2 years with a follow-up rate consistently over 90%.33 Survey questions focus on wealth, health, cognition and employment.34 Data are linked to federal health insurance (Medicare) claims.11 Participants provide informed consent on enrolment to HRS and again for linkage to Medicare insurance claims. We considered fee-for-service Medicare beneficiaries aged ≥65 who took part in at least one survey during 1998–2010 for inclusion in the study.

Study cohorts

We identified three study cohorts. The primary cohort was patients hospitalised with AHFR. The comparison cohorts were (1) adults not currently in the hospital (non-hospitalised adults) and (2) patients hospitalised with an acute inciting event that may cause AHFR (at-risk hospitalisations).

We defined AHFR hospitalisations pragmatically in administrative claims as (1) diagnostic codes for one or more acute inciting events that may result in AHFR—pneumonia, non-pulmonary infection, aspiration, trauma or pancreatitis and (2) respiratory failure—a procedural code for invasive mechanical ventilation and/or an explicit diagnosis code of acute respiratory failure (online supplementary table S1). We validated this claims-based AHFR definition in the University of Michigan Health System. It had a positive predictive value of 81.4% (95% CI 72.3% to 88.6%) and a negative predictive value of 91.9% (95% CI 90.0% to 93.4%) when compared with a clinical AHFR definition of PaO₂/FIO₂ < 300 while on ≥40% fraction of inspired oxygen (online supplementary appendix 1).

We defined at-risk hospitalisations as those with diagnostic codes for one or more acute inciting events that may result in AHFR, but no diagnostic code for either respiratory failure or mechanical ventilation. We excluded hospitalisations with diagnostic codes for chronic or sporadic respiratory failure from both hospitalised cohorts, as these diagnoses suggest a cause for respiratory failure other than an acute pulmonary process.

The cohort of non-hospitalised adults was not in the hospital at the time point that they were matched, but no exclusions were placed on this cohort being admitted to the hospitai either before or after the date of match. (Further details of this matching approach are described in a previous publication.29)

Matching

At the time of each HRS survey, we estimated each participant’s risk of having an AHFR hospitalisation in the next 2 years using multiple logistic regression. We selected the following predictor variables based on previous research and/or clinical experience suggesting it is an important risk factor for AHFR, or acute hospitalisation15

► Demographics: age, self-reported race and ethnicity, gender, partnership status.
► Economic status: wealth (sum of all assets and debts) standardised to 2013 $ using the annual gross domestic product price index,16 previous or current use of food stamps (a government assistance programme for low-income families and individuals).
► Healthcare utilisation: number of hospitalisations in the prior year, AHFR hospitalisation in the prior year, residence in a nursing home.
► Health status: limitations of five activities and six instrumental activities of daily living (IADLs/ADLs), self-rating of health, body mass index (BMI).
► Comorbidity burden: 17 Charlson comorbidities.17,18

Limitations of ADLs, IADLs, self-reported health, government assistance and BMI were missing in 5.6%, 0.1%, 0.1%, 0.5% and 1.4%, respectively, and these values were imputed using multiple imputation with chained equations and five imputations.19 Other covariates were present for the entire population. We considered select interactions (gender and partnership) and non-linear forms (age) based on a priori knowledge of the relations between the predictors and the likelihood of hospitalisation.20

We matched patients in the AHFR cohort 1:1 to patients in the at-risk hospitalisation cohort by age, percentile risk for AHFR hospitalisation, number of hospitalisations in the year prior to index admission, AHFR hospitalisation in the prior year, acute precipitating events during index hospitalisation (pneumonia, non-pulmonary infection, aspiration, trauma and pancreatitis) and other conditions during the index hospitalisation that may contribute to respiratory failure (congestive heart failure and asthma/chronic obstructive pulmonary disease) using coarsened exact matching.21

We examined the balance of each covariate (those directly matched, as well as those included within the multivariable risk for AHFR hospitalisation) between the hospitalised cohorts using X² and t tests, as appropriate, without consideration of any outcome.22 We then re-matched the hospitalised cohorts including any variables that were unbalanced on the initial match. We then re-examined and confirmed covariate balance before examining any results. We matched patients in the AHFR cohort 1:1 in adults in the non-hospitalised cohort again using an iterative process to ensure covariate balance. Further details are provided in the online supplementary appendix 2.

Outcomes and statistical analysis

Our primary outcome of interest was late mortality, defined as death in the 31 days to 2 years following AHFR hospital admission, at-risk hospital admission or the date of match, depending on the cohort. All patients were followed to death, or for 2 years. We used multivariable Cox proportional hazards models to calculate the HR for late mortality, adjusted for age, gender and propensity of AHFR. In addition, we used multivariable logistic regression models in order to calculate adjusted proportions of mortality. The technique of matching patients, then also adjusting for potential confounders using regression, is known as ‘doubly-robust’ estimation because it combines two methods for reducing bias and is less sensitive to misspecification.23,24 In the primary analysis, we accounted for the clustering of hospitalisations within people using clustered robust standard errors.25 This approach corrects standard errors so that significance tests, which assume independence of observations, are still valid.

Beyond measuring risk of late mortality overall, we also measured risk of death at multiple time intervals (0–30 days,
31–90 days, 181 days–1 year and 1–2 years) to examine how long excess mortality persists after AHRF. In addition, we present the risk of mortality over time in a hazard plot. To estimate the proportion of late mortality due to hospitalisation for the acute inciting event(s) versus the development of AHRF, we examined the ratio of logistic regression coefficients among AHRF hospitalisations that were matched to both controls (as described in the online supplementary appendix 2).

We performed several exploratory analyses. First, we examined whether excess late mortality after AHRF differed by type of acute inciting event, age, sex, self-rating of health, comorbidity burden, pre-AHRF functional limitations and nursing home residence. Second, we examined risk of late death separately for earlier (1998–2005) versus later hospitalisations (2006–2012), since management of AHRF has changed over time. Third, we examined risk of late death among patients with no hospitalisation for AHRF in the prior year. Fourth, we examined the proportion of patients surviving to 31 days who had a subsequent hospitalisation, as well as the subset who survived to 2 years. Fifth, we examined the principal diagnosis category of the final hospitalisations (a proxy for cause of death) among patients with AHRF with a late death.

We performed five sensitivity analyses, described in the online supplementary appendix 2. Conceptually, sensitivity analyses 1–3 used alternative approaches to account for clustering of observations within people, since each person may have multiple hospitalisations and/or times of non-hospitalisation. Sensitivity analyses 4 and 5 used alternate approaches to matching. We conducted all analyses with Stata MP V. 14 (StataCorp, College Station, TX). We used two-sided hypothesis testing and set significance at *p*<0.05.

**RESULTS**

From 15075 HRS participants with Medicare linkage and at least one survey completed in 1998–2010, we identified 1268 AHRF hospitalisations, 13117 at-risk hospitalisations and 15033 non-hospitalised adults (figure 1).

For the 1238 AHRF hospitalisations, patients were elderly (median age 79), 54.5% were female, 80.2% were white, 11.2% were nursing home residents, with a median of one functional limitation and two medical comorbidities (table 1). Over 60% of patients rated their health as fair or poor. Risk factors for AHRF were non-pulmonary infection (65.4%), pneumonia (45.0%), aspiration (18.0%), trauma (5.6%) and pancreatitis (1.3%). 33.5% had multiple risk factors.

Mortality was high: 42.7% at 30 days, 65.5% at 1 year and 73.3% at 2 years. Among the 727 patients who survived to day 31, 131 (18.0%) died by 90 days and 389 (53.5%) died by 2 years. Of the 389 patients who died during day 31 to 2 years, all but 23 (5.9%) were discharged from the hospital prior to death.

We were able to match 1157 AHRF hospitalisations (91.5% of all AHRF hospitalisations) to non-hospitalised adults, and 1017 AHRF hospitalisations (80.2%) to at-risk hospitalisations (figure 1, online supplementary tables S2 and S3). Nine hundred and fifty-two AHRF hospitalisations (75.1%) were matched to both an at-risk hospitalisation and a non-hospitalised adult; mortality for these double-matched cohorts is presented in online supplementary table S4. AHRF was associated with markedly increased acute mortality relative to matched non-hospitalised adults (adjusted HR (aHR) for 30-day mortality: 30.5; 95% CI 19.8 to 47.1) and matched at-risk hospitalisations (aHR for 30-day mortality: 4.0; 95% CI 3.3 to 4.8). The risk of mortality waned with time (figure 2), becoming equivalent to at-risk hospitalisation around day 90. However, risk remained slightly elevated compared with non-hospitalised controls to beyond 1 year.

Compared with adults not in the hospital matched on all baseline characteristics, patients with AHRF experienced a 24.4% (95% CI 19.9% to 28.9%) absolute increase (or 1.9-fold
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Table 1  Baseline characteristics of AHRF cohort (n=1268)

<table>
<thead>
<tr>
<th>Demographics</th>
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<tbody>
<tr>
<td></td>
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<tr>
<td>Age, years, median (IQR)</td>
<td>79 (72–85)</td>
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<tr>
<td>Male, n (%)</td>
<td>589 (46.5%)</td>
</tr>
<tr>
<td>Race</td>
<td></td>
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<tr>
<td>White/Caucasian</td>
<td>1017 (80.2%)</td>
</tr>
<tr>
<td>Black/African American</td>
<td>220 (17.4%)</td>
</tr>
<tr>
<td>Other</td>
<td>31 (2.4%)</td>
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<tr>
<td>Hispanic, n (%)</td>
<td>96 (8.4%)</td>
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<tr>
<td>Married or partnered, n (%)</td>
<td>591 (46.6%)</td>
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</table>

<table>
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<tr>
<th>Economic status</th>
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<tbody>
<tr>
<td>Total wealth</td>
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<tr>
<td>Quintile 5 positive assets</td>
<td>172 (13.7%)</td>
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<tr>
<td>Quintile 4 positive assets</td>
<td>186 (14.7%)</td>
</tr>
<tr>
<td>Quintile 3 positive assets</td>
<td>186 (14.7%)</td>
</tr>
<tr>
<td>Quintile 2 positive assets</td>
<td>242 (19.2%)</td>
</tr>
<tr>
<td>Quintile 1 positive assets</td>
<td>324 (25.7%)</td>
</tr>
<tr>
<td>Net negative or zero assets</td>
<td>150 (11.9%)</td>
</tr>
<tr>
<td>Government assistance</td>
<td>120 (9.6%)</td>
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<tr>
<th>Pre-AHRF health status</th>
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<tbody>
<tr>
<td>Charlson Comorbidity Index, median (IQR)</td>
<td>2 (0–4)</td>
</tr>
<tr>
<td>Congestive heart failure</td>
<td>387 (30.5%)</td>
</tr>
<tr>
<td>Dementia</td>
<td>97 (7.7%)</td>
</tr>
<tr>
<td>IADL/ADL limitations, median (IQR)</td>
<td>1 (0–5)</td>
</tr>
<tr>
<td>Self-rating of health, n (%)</td>
<td></td>
</tr>
<tr>
<td>Excellent</td>
<td>31 (2.5%)</td>
</tr>
<tr>
<td>Very good</td>
<td>150 (11.9%)</td>
</tr>
<tr>
<td>Good</td>
<td>300 (23.7%)</td>
</tr>
<tr>
<td>Fair</td>
<td>391 (30.9%)</td>
</tr>
<tr>
<td>Poor</td>
<td>394 (31.1%)</td>
</tr>
<tr>
<td>Body mass index, n (%)</td>
<td></td>
</tr>
<tr>
<td>Very severely obese</td>
<td>57 (4.6%)</td>
</tr>
<tr>
<td>Severely obese</td>
<td>73 (5.9%)</td>
</tr>
<tr>
<td>Obese</td>
<td>164 (13.2%)</td>
</tr>
<tr>
<td>Overweight</td>
<td>355 (28.6%)</td>
</tr>
<tr>
<td>Normal</td>
<td>499 (40.2%)</td>
</tr>
<tr>
<td>Underweight</td>
<td>94 (7.6%)</td>
</tr>
<tr>
<td>Multivariable risk for AHRF</td>
<td>5.9% (3.4%–10.7%)</td>
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</tbody>
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<table>
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<tr>
<th>Pre-AHRF healthcare use</th>
<th></th>
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<tbody>
<tr>
<td>Hospitalisations in prior year, median (IQR)</td>
<td>1 (0–2)</td>
</tr>
<tr>
<td>AHRF in prior year, n (%)</td>
<td>151 (11.9%)</td>
</tr>
<tr>
<td>Residence in a nursing home, n (%)</td>
<td>142 (11.2%)</td>
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<tr>
<th>Hospitalisation diagnoses</th>
<th></th>
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<tbody>
<tr>
<td>Risk factor for direct AHRF</td>
<td>766 (60.4%)</td>
</tr>
<tr>
<td>Pneumonia</td>
<td>570 (45.0%)</td>
</tr>
<tr>
<td>Aspiration</td>
<td>228 (18.0%)</td>
</tr>
<tr>
<td>Risk factor for indirect AHRF</td>
<td>890 (70.2%)</td>
</tr>
<tr>
<td>Non-pulmonary infection</td>
<td>834 (65.8%)</td>
</tr>
<tr>
<td>Trauma</td>
<td>71 (5.6%)</td>
</tr>
<tr>
<td>Pancreatitis</td>
<td>16 (1.3%)</td>
</tr>
</tbody>
</table>

AHRF, acute hypoxic respiratory failure; IADL, instrumental activities of daily living. relative increase) in late mortality. Adjusted 31-day to 2-year mortality was 52.4% (95% CI 48.5% to 56.2%) in patients with AHRF surviving to at least 31 days versus 28.0% (95% CI 25.4% to 30.6%) among matched non-hospitalised patients who survived to at least 31 days (aHR for late mortality: 2.5, p<0.001) (figure 3a, online supplementary table S5). Adjusted 91-day to 2-year mortality was 42.0% (95% CI 37.9% to 46.2%) in survivors with AHRF versus 25.9% (95% CI 23.3% to 28.6%) in matched non-hospitalised patients, p<0.001. The higher mortality in the AHRF cohort persisted for the full 2-year follow-up period. Among patients with AHRF who survived to 1 year, the adjusted 2-year mortality was 21.0% (95% CI 17.0% to 25.0%) versus 14.9% (95% CI 12.6% to 17.2%) in their controls who were not in the hospital at the time of matching (aHR for 2-year mortality: 1.4, p=0.006).

Compared with matched patients hospitalised with the same acute inciting event, patients with AHRF experienced a 6.7% (95% CI 1.7% to 11.7%) absolute increase (1.15-fold relative increase) in late mortality (aHR for late mortality: 1.2, p=0.01) (figure 3b, online supplementary table S5). The increased risk for late mortality did not persist beyond 90 days. Among those who survived to at least 91 days, risk mortality at 2 years was indistinguishable from patients surviving hospitalisation for AHRF risk factors alone (aHR for 2-year mortality=1.1, p=0.23).

Among patients with AHRF who survived to day 31 and were matched to both an at-risk hospitalisation and a non-hospitalised adult, at-risk hospitalisation accounted for 71.2% (95% CI 52.7% to 89.6%) of the increased odds of late mortality, whereas the development of AHRF itself accounted for 28.8% (95% CI 10.4% to 47.3%) of the increased odds of late mortality.

In all five sensitivity analyses, the point estimates of ORs for late mortality relative to non-hospitalised adults and relative to patients with an at-risk hospitalisation were similar to the ORs in the primary analyses (online Supplementary table S6). Statistical significance was lost, however, in two instances due to the smaller sample sizes in the sensitivity analysis. Results were also similar for earlier versus later hospitalisations.

In stratified analysis, the excess late (31 days to 2 years) mortality after AHRF was relatively constant across patient subgroups defined by type of acute inciting event, sex, comorbidity burden, functional limitations, self-rating of health and nursing home residence (Figure 4 online supplementary figure S1). However, the incremental effect of AHRF versus at-risk hospitalisation on late mortality was greater in patients ≥80 years versus patients<80 years (figure 4).

Risk for late death was similar when limiting the analysis to earlier hospitalisations (1998–2005), later hospitalisation (2006–2012) and patients without a hospitalisation for AHRF in the prior year (online supplementary table S7 and S8).

Re-hospitalisation was more common in AHRF versus comparison cohorts (online supplementary table S9 and figure S2). Of the 398 patients with AHRF who experienced a late death following AHRF, 290 (72.9%) were re-hospitalised prior to their death. The most common hospitalisation diagnoses prior to death were sepsis (16.6% of terminal hospitalisations), respiratory failure (15.2%), congestive heart failure (9.0%), aspiration pneumonia (6.6%) and pneumonia (5.2%).

DISCUSSION

In this national cohort of older Americans, late mortality was substantially increased after AHRF relative to non-hospitalised adults. Among patients who survived to at least day 31 after hospitalisation...
admission for AHRF, more than half died within the next 2 years. This high rate of late mortality was not explained by age, sociodemographics or health status prior to AHRF. Compared with adults not in the hospital who were indistinguishable on a variety of potential confounders prior to acute illness, patients surviving AHRF have a nearly 25% absolute increase in late mortality.

The majority (70%) of the late mortality after AHRF was explained by hospitalisation for the acute inciting event(s), while a minority (30%) was explained by the development of AHRF. Compared with patients hospitalised with acute inciting event(s) but without AHRF, those with AHRF had a 7% absolute increase in late mortality. Among hospitalised patients who survived for at least 90 days; however, the risk for subsequent mortality was indistinguishable. Subgroup analysis suggested that the incremental effect of AHRF on late mortality may be concentrated in the oldest patients.

Prior studies have shown that ARDS is associated with significantly increased early (30-day) mortality compared with hospitalisation for inciting events that may lead to ARDS, such as hospitalisation for sepsis and trauma. Our findings on early mortality are similar. Compared with patients who were hospitalised for acute inciting events alone, AHRF was associated with a fivefold relative increase, or approximately 30% absolute increase in 30-day mortality in our study. This can be considered a form of ‘positive control’, in that it confirms that our AHRF population is distinct from the non-AHRF hospitalised patients.

We were able to identify just one prior study that examined late mortality of survivors with ARDS or AHRF relative to matched controls. This study of patients treated in the mid-1990s found no difference in late mortality between 127 survivors with ARDS and 127 controls matched by acute inciting event (sepsis or trauma), illness severity and admission half-year. However, the study had limited power, and so an important difference in late mortality (as large as 13%) could not be ruled out. With a larger sample, our study was able to detect a small increase in risk of late mortality after AHRF relative to hospitalisations for acute inciting events alone, but approximately 70% the excess risk for late mortality after AHRF was explained by at-risk hospitalisation.

While it may seem counterintuitive for survivors with AHRF to experience similar late mortality as survivors of less severe illness, this phenomenon has precedent. Elderly survivors of in-hospital cardiac arrest have only modestly lower 2-year mortality risk compared with non-AHRF hospitalised controls.
survival rates compared with patients who survive a hospitalisation for heart failure, and the survival curves converge at 3 years.\(^3\)

The contrast between the present results showing a small degree of excess late mortality after AHRF relative to matched hospitalised controls and previous work using a similar methodology for late mortality after sepsis\(^9\) is intriguing. It raises the possibility that our focus on mechanical ventilation as the key risk factor in mediating late harm in patients surviving the ICU may be too narrow. For example, mechanical ventilation is often a key inclusion criteria for post-ICU rehabilitation studies,\(^3\) but may not target the most appropriate population. The contrast also suggests that the inflammation and multi-organ failure associated with ventilator-induced lung injury\(^3\) may resolve faster than the inflammation and immune-suppression that follows sepsis.\(^3\)

Future epidemiological and translational studies are needed to better understand the duration and recovery of these pathobiological derangements—and how the natural history differs between AHRF and sepsis.

Our study should be interpreted in the context of several limitations. First, this is an observational study, so cannot prove causation. We have attempted to control for confounding as best as possible through matching and regression adjustment, as well as by studying a cohort with prospectively collected data on a range of potential confounders. Second, we ascertained AHRF by International Classification of Diseases, Ninth Revision,
Clinical Modification (ICD-9-CM) coding for acute inciting events and respiratory failure. There may have been misclassification in both directions between at-risk and AHRF hospitalisations, but the negative and positive predictive values from our validation suggest that the degree of misclassification is likely small. Third, our study spanned a period of 15 years when treatment for AHRF changed. However, when limiting our analysis to only early or late hospitalisations, the associations were similar. Fourth, we were not able to examine the subset of patients with AHRF and ARDS because there is no reliable way to identify ARDS in administrative claims. Nonetheless, we believe AHRF to be a clinically meaningful study population, and postulate that our findings are reflective of ARDS patients as well. Fifth, we excluded patients aged <65 years because our study relied on Medicare claims, and only limited groups aged under 65 qualify for Medicare coverage.

Our study has several strengths. First, we believe that it is the only recent study to examine the excess late mortality associated with AHRF. Second, by examining a national cohort with detailed survey and claims data, we were able to achieve robust control for confounding. Third, by comparing patients with AHRF with two controls, we were able to disentangle the proportion of excess mortality related to hospitalisations for acute inciting events versus the development of AHRF. Finally, our findings were robust to several sensitivity analyses.

CONCLUSION
In a national cohort of older Americans, we have shown that AHRF is associated with a high rate of late mortality that is not explained by health status before AHRF. However, more than 70% of the excess risk for late mortality is explained by hospitalisation for an acute inciting event, not hypoxemic respiratory failure.

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