Prevalence of pulmonary hypertension in mitral regurgitation and its influence on outcomes

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ABSTRACT
Objective Pulmonary hypertension (PHT) commonly coexists with significant mitral regurgitation (MR), but its prevalence and prognostic importance have not been well characterised. In a large cohort of adults with moderate or greater MR, we aimed to describe the prevalence and severity of PHT and assess its influence on outcomes.

Methods In this retrospective study, we analysed the National Echocardiography Database of Australia (data from 2000 to 2019). Adults with an estimated right ventricular systolic pressure (eRVSP), left ventricular ejection fraction >50% and with moderate or greater MR were included (n=9638). These subjects were then categorised according to their eRVSP. The relationship between PHT severity and mortality outcomes was evaluated (median follow-up of 3.2 years, IQR 1.3–6.2 years).

Results Subjects were aged 76±12 years, and 62.6% (6038) were women. Overall, 959 (9.9%) had no PHT, and 2952 (30.5%), 3167 (32.7%), 1588 (16.4%) and 1017 (10.5%) patients had borderline, mild, moderate and severe PHT, respectively. A ‘typical left heart disease’ phenotype was identified with worsening PHT, showing rising E:e′, right and left atrial sizes increasing progressively, from no PHT to severe PHT (p<0.0001, for all). With increasing PHT severity, 1- and 5-year actuarial mortality increased from 8.5% and 33.0% to 39.7% and 79.8%, respectively (p<0.0001). Similarly, adjusted survival analysis showed the risk of long-term mortality progressively increased with higher eRVSP levels (adjusted HR 1.20–2.86, borderline to severe PHT, p<0.0001 for all). A mortality inflection was apparent at an eRVSP level >34.00 mm Hg (HR 1.27, CI 1.00–1.36).

Conclusions In this large study, we report on the importance of PHT in patients with MR. Mortality increases as PHT becomes more severe from an eRVSP of 34 mm Hg onwards.

INTRODUCTION
Mitral regurgitation (MR) is an increasingly prevalent valvular problem in developed countries and is the second-most common valve lesion requiring operative management in Europe and the USA, after aortic stenosis (AS). Symptoms and outcomes generally correlate with both the severity of the regurgitation and the myocardial response to volume overload. The identification of prognostic factors is important to risk stratify patients and to potentially guide treatment decisions. While it is recognised that pulmonary hypertension (PHT) is a potential complication of MR, this is actually not well documented. There are varying reports, in relatively small cohorts on the prevalence and echocardiographic phenotype of these patients and, thus, an incomplete understanding of the prognostic impact of PHT in significant MR.

Group 2 PHT, or PHT due to left heart disease (LHD), is the most common type of PHT. PHT is thought to be a predictive feature of deterioration in these patients and likely arises from elevated left atrial (LA) pressure causing back pressure into the pulmonary vasculature. In MR specifically, PHT is thought to be due to the direct effect of systolic backflow into the LA and...
may develop before patients experience symptoms or left ventricular (LV) systolic dysfunction.\textsuperscript{5,9,12}

Echocardiography remains the the most common screening tool for PHT and is widely used in clinical practice. Using the power of ‘big’ data from the National Echo Database of Australia (NEDA), a registry with contemporary community and hospital-based echo data on over 600,000 unique adult subjects from over 25 centres across Australia, we aimed to describe the prevalence of PHT in adults with moderate or greater MR and to assess the influence of PHT severity on outcomes.

METHODS

NEDA database and study design

The NEDA is a multicentre registry, and the purpose and methodology of which have been previously described.\textsuperscript{13-15} NEDA contains basic demographic and detailed echocardiographic data of adults from >25 centres across Australia. The database is linked with the National Death Index (NDI), provided by the Australian Institute for Health and Welfare; the NDI provides mortality data on each individual. The study period included >1 million echo reports from >600,000 individuals, studied between January 2000 and June 2019. Vital status was determined as of 21 May 2019 (median follow-up of 6.2 years, IQR 3.8–9.8 years); patients alive at this date were censored alive. NEDA is registered with the Australian New Zealand Clinical Trials Registry (ACTRN12617001387314), and human ethics approval was obtained from the Sydney Local Health District Human Research Ethics Committee, protocol X15-0587 and 2019/ETH069899. A retrospective waiver of consent was authorised as part of this ethics protocol.

Study cohort

Figure 1 shows our study flow diagram, and this is consistent with our previous work on PHT in left-sided valvular pathology.\textsuperscript{16,17} NEDA data at the time of study census were used to identify a cohort of patients with significant MR to characterise their relationship with PHT: (1) adults ≥18 years of age, (2) with at least one echocardiogram recorded on the system (where patients had multiple studies only the last study was included in the analysis), (3) with a recorded left ventricular ejection fraction (LVEF), estimated right ventricular systolic pressure (eRVSP) and (4) with moderate or greater MR. Text extraction was used to identify patients with moderate and severe MR as well as right ventricular (RV) size and function.\textsuperscript{14} While quantitative measures of MR were interrogated, sufficient data were not available in the majority of patients, a reflection of real-world practice within Australia. All participating laboratories use an integrative and semiquantitative approach for grading MR severity as recommended by the American Society of Echocardiography.\textsuperscript{18} Patients with mitral valve (MV) replacements were excluded from primary analysis as were patients with evidence of significant mitral stenosis (conservatively defined as mitral valve gradient >5 mm Hg), LVEF <60% and patients with less than moderate MR. eRVSP was conservatively derived using the Bernoulli equation (4×[(tricuspid regurgitation velocity) TRV]\textsuperscript{2}+assumed RA pressure of 5 mm Hg).\textsuperscript{19}

Study methods

Similar to our previous work,\textsuperscript{16,17} once the cohort of patients with moderate or greater MR was established, subjects were categorised according to their eRVSP, according to clinical guidelines\textsuperscript{10,20} to document the distribution of eRVSP and thence PHT severities. A ‘borderline PHT’ group which has previously been determined as potentially significant in both NEDA papers and other recent prospective publications\textsuperscript{11,21,22} was included. Defined categories were: (1) normal (eRVSP <30 mm Hg), (2) borderline (30.00–39.99 mm Hg), (3) mildly elevated (40.00–49.99 mm Hg), (4) moderately elevated (50.00–59.99 mm Hg) and (5) severely elevated (eRVSP ≥60 mm Hg).

We then analysed the eRVSP data according to decile distribution:\textsuperscript{14} first decile—5.00–30.00 mm Hg, second decile—30.01–34.00 mm Hg, third decile—34.01–37.04 mm Hg, fourth decile—37.05–39.16 mm Hg, fifth decile—39.17–43.00 mm Hg, sixth decile—43.01–46.00 mm Hg, seventh decile—46.01–48.44 mm Hg, eighth decile—48.45–53.00 mm Hg, ninth decile—53.01–61.00 mm Hg and 10th decile—>61.00 mm Hg.

All-cause mortality was determined during a median follow-up of 3.2 years (IQR 1.3–6.2 years). We explored the relationship between eRVSP level and survival, looking at both clinically defined groups (as above) and the eRVSP deciles.

Statistical analysis

All continuous variables are expressed as mean±SD, unless otherwise stated, and categorical data are expressed as frequency and percentages. For continuous variables, linear regression analysis using analysis of variance was used to test whether the trend of the mean across the categorical groups of eRVSP levels was linear. For binary variables, the \(\chi^2\) test was used to determine if there was a trend in the change in proportions across the groups.

Actuarial 1-year and 5-year survival rates for all-cause mortality were calculated from 9439 (97.9%) and 6780 (70.3%) subjects with complete follow-up for those time points. Multiple logistic regression models (entry at univariate p-value <0.05) were used to derive adjusted ORs for mortality models at fixed time points. Cox regression hazard models were used to derive adjusted HRs for mortality outcomes during follow-up (entry model at a univariate p-value <0.05). Adjusted analyses included age and sex. A sensitivity analysis was performed excluding patients with significant concurrent severe AS and moderate or greater aortic regurgitation (AR). Patients with moderate and severe MR were also assessed separately to determine if there were differences between these two groups. Severe MR was defined as ‘severe...
Pulmonary vascular disease

NEDA Registry 2.0
1060 231 investigations and 622 477 individuals

Last study only in 622 477 individuals
M = 327430 (52.6%) age 61 ± 17 yrs, F = 295 047 (47.4%) age 61 ± 18 yrs

184 813 no eRVSP documented
156 118 no LVEF documented
281 546 individuals
96 615 LVEF <60%
10 353 individuals
248 documented MVR

422 patients with ≥ moderate mitral stenosis

9683 patients with ≥ moderate MR, with preserved LVEF (≥ 60%)
Documented ≥ moderate MR on text extraction from echo report

Is PHT present?
eRVSP

No PHT (<30.00 mmHg)
959 individuals (9.9%)

Borderline (30.00-39.99)
2952 individuals (30.5%)

Moderate (50.00-59.99)
1588 individuals (16.4%)

Mild (40.00-49.99)
3167 individuals (32.7%)

Severe (>60.00)
1017 individuals (10.5%)

Figure 1  Study flow chart. This figure shows the analysis flow chart, performed in this study. eRVSP, estimated right ventricular systolic pressure; LVEF, left ventricular ejection fraction; MR, mitral regurgitation; MVR, mitral valve replacement; NEDA, National Echo Database Australia; PHT, pulmonary hypertension.

Figure 2 shows the frequency distribution of eRSP levels (median 43.00 mm Hg, IQR 35.46–50.96 mm Hg). The number of patients in each subgroup was as follows: no PHT (eRVSP <30 mm Hg)—959 individuals (9.9%), borderline PHT (eRVSP 30.00–39.99 mm Hg)—2952 individuals (30.5%), mild PHT (eRVSP 40.00–49.99 mm Hg)—3167 individuals (32.7%), moderate PHT (eRVSP 50.00–59.99 mm Hg)—1588 individuals (16.4%) and severe individuals (eRVSP >60 mm Hg)—1017 individuals (10.5%).

RESULTS
Prevalence of PHT and distribution of eRVSP
A total of 9638 patients with moderate or greater MR, normal left ventricular systolic function and eRVSP data were identified; the majority (62.6%) were women.

mitral regurgitation’ on text extraction. All analyses were performed with SPSS software version 25.0 (IBM, Armonk, New York, USA), and statistical significance was accepted at a two-tailed p-value of <0.05.
Cohort profile

Table 1 summarises the demographic and echocardiographic characteristics of the study cohort divided into subgroups based on measured eRVSP levels. Age was greater in those with higher eRVSP levels, from a mean of 69±17 years in patients with no PHT to 81±10 years in patients with moderate PHT, before a plateau was noted in those with severe PHT (80±11 years) (p<0.0001 for all). The proportion of patients with atrial fibrillation or an atrial arrhythmia was greater in those with higher eRVSP levels (24.0% vs 54.8%, no PHT vs severe PHT, respectively).

A typical pattern of worsening ‘LHD’ phenotypic response with worsening PHT was evident. E:e increased progressively with increased severity of PHT (12.90±5.63 vs 17.16±7.01, no PHT vs severe PHT, respectively, p<0.0001 for all). An increase in right atrial area and indexed left atrial volume was also noted (18.18±7.25 cm² vs 33.71±9.35 cm² and 43.84±24.21 vs 108.95±55.95 mL/m², no PHT vs severe PHT, respectively, p<0.0001 for all) though indexed left atrial volume plateaued in those with severe PHT. Qualitative observations showed an increased proportion of patients with increased RV dilation and functional impairments as eRVSP levels increased.

The differences between male and female patients are shown in online supplemental table 1. Female patients were older than their male counterparts (77±13 years vs 75±12 years) but did not show clinically meaningful differences in eRVSP level. Females had higher E:e but lower right atrial area and indexed LA volume compared with males.

Survival data

The survival profile of the cohort based on the severity of PHT as determined on echocardiography is summarised in table 2. All-cause mortality at 1 and 5 years (actuarial mortality) and long-term survival (all adjusted for age and gender) were reported between those with eRVSPs <30.00 mm Hg and the four categories of progressively elevated eRVSP. As predicted, the risk for mortality markedly increased with higher eRVSP levels. This was shown by the range in 1-year and 5-year actuarial mortality from a low of 8.5% and 33.0% to a high of 39.7% and 79.8%, in those with normal to severely elevated eRVSPs. This trend was mirrored in adjusted long-term mortality results which showed a 1.20-fold increase in risk in those with borderline PHT compared to a 2.86-fold increase in those with severe PHT (p<0.0001 for all) (figure 3A). Cardiovascular mortality trends showed an increased risk in those with moderate and severe PHT (table 2). Trends were less clear in those with smaller numbers and possible inaccurate coding for causes of death documented on death certificates, as possible contributing factors.

A sensitivity analysis was performed excluding patients with moderate or greater AR or severe AS. The trends were maintained in 1-year and 5-year actuarial analysis, though only achieved statistical significance from mild PHT onwards. Adjusted long-term mortality continued to
Pulmonary vascular disease progressively increase as eRVSP level increased (table 3, figure 3B). Indexed LA volume was also included as a variable in the survival models, with mortality trends mirroring the above results (online supplemental table 2). In all models, increasing age and male sex were also associated with increasing mortality (p<0.0001, for all).

Mortality was assessed when the cohort was divided into two cohorts based on severity (moderate MR—8214 patients, severe MR—1469 patients). In the moderate MR cohort, which had larger numbers, mortality outcomes mirrored that of the total cohort (online supplemental table 3). Mortality outcomes had a similar trend in those with severe MR, although statistical significance was not reached at milder elevations of eRVSP level, most likely a consequence of the smaller subject numbers (online supplemental table 4). Mortality outcomes among women (n=6038) matched with those in the total cohort (HR 1.32, 95% CI 1.10–1.58 for borderline PHT vs HR 2.92, 95% CI 2.42–3.53 for severe PHT) (online supplemental table 5). The male cohort (n=3645) had similar trends, though significance was only reached when pulmonary pressures were mildly elevated (online supplemental table 6).

Table 1  Baseline characteristics of study cohort (N=9683)

<table>
<thead>
<tr>
<th></th>
<th>eRVSP 0.00–29.99</th>
<th>eRVSP 30.00–39.99</th>
<th>eRVSP 40–49.99</th>
<th>eRVSP 50–59.99</th>
<th>eRVSP &gt;60.00</th>
<th>P value</th>
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<tr>
<td></td>
<td>n=959</td>
<td>n=2952</td>
<td>n=3167</td>
<td>n=1588</td>
<td>n=1017</td>
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<td>Demographics</td>
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<tr>
<td>Age, years</td>
<td>69±17</td>
<td>74±13</td>
<td>78±11</td>
<td>81±10</td>
<td>80±11</td>
<td>&lt; 0.0001</td>
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<td>Female (%)</td>
<td>611 (63.7)</td>
<td>1773 (60.1)</td>
<td>1992 (62.9)</td>
<td>1011 (63.7)</td>
<td>651 (64.0)</td>
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<td>BMI</td>
<td>25.92±5.15</td>
<td>26.02±5.40</td>
<td>26.52±5.76</td>
<td>26.56±6.03</td>
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<td>BSA</td>
<td>1.80±0.24</td>
<td>1.80±0.24</td>
<td>1.80±0.25</td>
<td>1.80±0.24</td>
<td>1.76±0.24</td>
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<td>Rhythm</td>
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<td>Atrial fibrillation/arrhythmia</td>
<td>230 (24.0)</td>
<td>947 (32.1)</td>
<td>1268 (40.0)</td>
<td>770 (48.5)</td>
<td>557 (54.8)</td>
<td>&lt;0.0001</td>
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<td>LV dimensions and function</td>
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<td>LVEF %</td>
<td>66.14±5.41</td>
<td>68.13±6.89</td>
<td>70.53±7.90</td>
<td>70.40±7.89</td>
<td>70.02±7.90</td>
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<tr>
<td>Stroke volume index (mL/m²)</td>
<td>44.06±11.97</td>
<td>44.98±14.37</td>
<td>44.96±14.79</td>
<td>43.49±14.09</td>
<td>42.15±14.99</td>
<td>0.06</td>
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<tr>
<td>E:e’ ratio</td>
<td>12.90±5.63</td>
<td>13.07±5.68</td>
<td>14.14±5.27</td>
<td>15.35±5.51</td>
<td>17.16±7.01</td>
<td>&lt;0.0001</td>
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<td>LVEDD</td>
<td>4.50±0.67</td>
<td>4.71±0.69</td>
<td>4.87±0.73</td>
<td>4.92±0.77</td>
<td>4.85±0.82</td>
<td>&lt;0.0001</td>
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<td>LVESD</td>
<td>2.87±0.53</td>
<td>2.87±0.57</td>
<td>2.84±0.64</td>
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<td>Atrial dimensions</td>
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<tr>
<td>LA volume index, mL/m²</td>
<td>43.84±24.23</td>
<td>67.90±41.04</td>
<td>95.53±47.82</td>
<td>108.92±54.28</td>
<td>108.95±55.95</td>
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<td>RA area, cm²</td>
<td>18.18±7.25</td>
<td>24.21±8.76</td>
<td>29.03±8.50</td>
<td>32.12±9.01</td>
<td>33.71±9.35</td>
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<td>Right heart dimensions and function</td>
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<td>eRVSP, mm Hg</td>
<td>25.66±3.63</td>
<td>35.43±2.94</td>
<td>44.69±2.76</td>
<td>53.97±2.93</td>
<td>71.10±10.84</td>
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<td>TR peak velocity, m/s</td>
<td>2.19±0.23</td>
<td>2.56±0.17</td>
<td>2.94±0.14</td>
<td>3.30±0.14</td>
<td>3.86±0.33</td>
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<td>RV basal diameter</td>
<td>3.53±0.75</td>
<td>3.30±0.47</td>
<td>3.36±0.37</td>
<td>3.44±0.40</td>
<td>3.56±0.47</td>
<td>&lt;0.0001</td>
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<td>Dilated RV</td>
<td>95 (9.9)</td>
<td>509 (17.2)</td>
<td>1051 (33.2)</td>
<td>673 (24.2)</td>
<td>521 (51.2)</td>
<td>&lt;0.0001</td>
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<td>Impaired RV function</td>
<td>14 (1.5)</td>
<td>28 (0.9)</td>
<td>59 (1.9)</td>
<td>54 (3.4)</td>
<td>82 (8.1)</td>
<td>&lt;0.0001</td>
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<td>Mitral valve dimensions and function</td>
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<td>MV mean gradient, mm Hg</td>
<td>2.32±1.18</td>
<td>2.63±1.20</td>
<td>3.01±1.16</td>
<td>3.31±1.18</td>
<td>3.09±1.22</td>
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<td>Concomitant valvular pathology</td>
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<td>Moderate or greater aortic regurgitation</td>
<td>85 (8.9)</td>
<td>246 (8.3)</td>
<td>369 (11.7)</td>
<td>229 (14.4)</td>
<td>156 (15.3)</td>
<td>&lt;0.0001</td>
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<td>Severe AS &lt;1 cm²</td>
<td>26 (2.7)</td>
<td>89 (3.0)</td>
<td>121 (3.8)</td>
<td>82 (5.2)</td>
<td>77 (7.8)</td>
<td>&lt;0.0001</td>
</tr>
</tbody>
</table>

Values are n (%) unless otherwise indicated.
AS, aortic stenosis; AV, aortic valve; BMI, body mass index; BSA, body surface area; eRVSP, estimated right ventricular systolic pressure (mm Hg); LA, left atrial; LV, left ventricle; LVEDD, left ventricular end diastolic diameter; LVEF, left ventricular ejection fraction; LVESD, left ventricular end systolic pressure; MV, mitral valve; RA, right atrial; RV, right ventricle; TR, tricuspid regurgitant; VTI, velocity-time integral.
Threshold for mortality

A Cox regression model was constructed using the decile distribution of eRVSP and adjusted for age and sex. This confirmed that the adverse effects of PHT were noted from the third decile (eRVSP 30.00–39.99 mm Hg) relative to the lowest decile <30.00 mm Hg (HR 1.16, 95% CI 1.00–1.36, p = 0.05). Risk increased progressively from those with borderline PHT in the third decile to the 10th decile (eRVSP 60.01–143.63; HR 2.89, 95% CI 2.51–3.32, p < 0.0001) (online supplemental table 7). Hence, the adjusted risk for mortality is markedly higher in those with borderline PHT in the third decile to the 10th decile (eRVSP 60.01–143.63; HR 2.89, 95% CI 2.51–3.32, p < 0.0001) (online supplemental table 7). Hence, the adjusted risk for mortality is markedly higher in those with borderline PHT and above regardless of age, gender or cause of death.

**DISCUSSION**

This large, ‘real-world’ cohort study, including over 9500 patients examines the relationship between MR and PHT, in patients with preserved ejection fraction (EF). The use of ‘big data’ through the NEDA, which includes over 1 million echoes in over 600 000 individuals, allowed for a more detailed, contemporary description of the prevalence and phenotype of these patients. We confirmed the negative short-term and long-term prognostic impact PHT has in MR and have documented that the threshold for excess mortality lies within the range of ‘borderline PHT’.

PHT is likely to occur in patients with significant MR via the following mechanism; in the initial phase, the direct effect of systolic backflow into the LA and subsequent volume overload leads to compensatory LA and LV dilation. Over time, there is decompensation leading to both LV systolic and diastolic dysfunction and reduced LA compliance. This causes elevated LA pressure and increases pulmonary capillary wedge pressure causing postcapillary PHT. This may develop before patients experience symptoms or LV dysfunction.

**Prevalence and phenotype of PHT with MR**

The prevalence of PHT complicating significant MR remains unclear with the rate thought to be tied to the grade of MR. Prevalence increases with the presence of symptoms and LV systolic dysfunction, with rates as high as 64% reported in those with New York Heart Association function classes III and IV and symptoms compared with <20% in asymptomatic patients with severe MR and preserved EF. Significant PHT (eRVSP >50 mm Hg) was reported in 23% in a contemporary cohort of patients with severe degenerative MR. In our large, contemporary cohort of hospital-based and community-based patients, who underwent an echocardiogram, we confirm the high prevalence of PHT in patients with significant MR and preserved LVEF; 59.6% of patients had some degree of PHT as defined by clinical guidelines (mild—32.7%, moderate—16.4%, severe—10.5%). Patients with ‘borderline PHT’ (eRVSP 30–39 mm Hg) represented 30.5% of the cohort.

Prior studies have shown that increased age, female sex, increased E/e’, and larger LA size are all independent predictors of raised pulmonary pressures in patients with significant MR. We confirmed this echocardiographic phenotype in our cohort and reported progressively higher proportions of RV dilation and dysfunction as PHT worsens. While there is a close relationship between PHT and both mitral stenosis and LV systolic dysfunction, the prospective exclusion of these patients from our cohort allows a greater understanding of the impact of PHT per se in patients with MR with preserved EF. A comprehensive understanding of this echocardiographic phenotype is important as it provides clinicians with an easily accessible, non-invasive method to monitor specific parameters associated with worsening PHT and thus prognosticate these patients more accurately.
Figure 3  Adjusted risk for all-cause mortality using Cox proportional hazards showing as estimated right ventricular systolic pressure (eRVSP) level increases based on clinical severity, risk of mortality increases in (A) the total cohort and (B) the cohort excluding patients with severe aortic stenosis and/or moderate or greater aortic regurgitation.
Outcomes of PHT in patients with MR
The presence of PHT negatively impacts outcomes with previous studies reporting increased postoperative heart failure, worse LV systolic dysfunction and poorer survival in patients with preoperative PHT compared with those without. Our much larger study confirms the serious impact of PHT on patients with significant MR, even in the absence of LV systolic dysfunction, with 59.6% of patients with eRVSP >40.00 mm Hg having a 1.20- to 2.86-fold adjusted increased risk of long-term all-cause mortality (dependent on PHT severity) compared with those with no PHT. Similar to our previous studies, we find that even ‘borderline’ PHT (eRVSP 30.00–39.99 mm Hg) confers an increased risk of all-cause mortality compared with those with normal eRVSP. This was evident in the short term, with 1-year actuarial mortality increasing 1.14-fold, and, in the long-term, with all-cause mortality risk increasing 1.20-fold. Furthermore, the large numbers provided by the NEDA allowed us to identify an inflection point for excess mortality risk, at eRVSP >34.00 mm Hg. These results were independent of age and gender and did not appear to be confounded by the presence of concomitant valvular pathology including moderate or greater AR or severe AS.

Clinical implications
Current clinical guidelines include the presence of significant PHT (eRVSP >50 mm Hg, confirmed on right heart catheterisation) as an indication for the operative management in asymptomatic patients with severe MR. Prior studies have shown that ‘early’ MV repair in asymptomatic patients with severe MR and normal LVEF leads to improvements in long-term mortality and HF hospitalisation. Furthermore, while there is a decrease in pulmonary pressures post-intervention, a significant degree of PHT remains despite amelioration of the valvular pathology. While we do not report on treatment effect, we demonstrate that even minor elevations in pulmonary pressures are associated with negative prognostic implications. With no specific medical therapy approved for PHT in the setting of MR, further studies will be needed to determine whether early mitral valve intervention might improve outcomes in those patients with MR with early stages of PHT or perhaps even before it develops.

Limitations
NEDA provides detailed echocardiographic data and linkage to mortality, and NEDA does not (yet) provide granular clinical data such as symptoms, comorbidities or pharmacological treatments. Most patients included in the database have undergone an echocardiogram for investigation of confirmed or suspected cardiac disease and should not be taken to reflect the population prevalence.

As noted in our previous studies, the data concerning PHT in NEDA are based on echocardiography-based measures, rather than haemodynamic assessment at right heart catheterisation. Prior studies have correlated eRVSP with invasive pulmonary artery systolic pressure, supporting the broad validity of our approach. Furthermore, echocardiography remains the most common screening tool to detect PHT and is the guideline-recommended diagnostic method of choice, to allow for monitoring and follow-up. We acknowledge that the definitive diagnosis of PHT and its underlying aetiology (precapillary vs postcapillary vs combined) should generally be confirmed on right heart catheterisation, after initial screening is suggestive of PHT. We also note that the absence of a tricuspid regurgitation jet does not preclude the presence of PHT and there may be a number of patients with MR and PHT who were not included in the study due to the lack of correct tricuspid regurgitant (TR) sampling or no quantifiable TR. Hence, both the prevalence and prognostic impacts reported in
this cohort should be interpreted as the minimum indicative values, from an epidemiological standpoint.

These studies were primarily derived from specialist centres or clinics across Australia, so some caution should be applied when applying these findings to other populations. However, Australia is a diverse and multi-ethnic population with universal health coverage aspects captured within the NEDA database. Finally, our data are lacking in quantitative measures of MR or RV dysfunction, a reflection of real-world echocardiographic practices in Australia. We were also unable to ascertain the underlying aetiology of the MR, so could not definitively differentiate between the primary versus secondary MR. However, patients with reduced LVEF were prospectively excluded, thus, our cohort likely represents a combination of patients with primary MR and atrial functional MR.

CONCLUSION
This is the largest series of adults to characterise the close, but incompletely understood, relationship between significant MR and complicating PHT. In patients with moderate or greater MR and preserved LV systolic function, mortality increases progressively as PHT becomes more severe from an eRSP 34 mm Hg onwards.

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Acknowledgements We would like to thank all the NEDA centres and their patients for contributing to these data.

Contributors GS and DP conceived and designed the NEDA Study. SR and DCC conceived this analysis and conducted study analyses, and all authors contributed to the interpretation of study data. SR and DCC wrote the manuscript, and all authors contributed to its revision. GS and DP are the guarantors of the overall veracity and accuracy of NEDA data presented in this manuscript.

Funding This research did not receive any specific grants from funding agencies in the public, commercial, or not for profit sectors. However, NEDA has received investigator initiated funding support from Janssen, Novartis Pharmaceuticals and Edwards Lifesciences in the past 3 years. SR is supported by the Heart Research Institute Australia, Emerging Cardiovascular Researcher Education Scholarship. Both NEDA (grant 1055214) and SS (grant 11358940) are supported by the National Health and Medical Research Council of Australia.

Competing interests SS, DP and GS have previously received consultancy/speaking fees from Edwards Lifesciences. DP and GS have received consultancy fees from Medtronic, Edwards Lifesciences, Abbott Laboratories and ECHO IQ Pty Ltd.

Ethics approval This study involves human participants and was approved by Sydney Local Health District Human Research Ethics Committee, protocol X15-0387 and 2019/ETH00899. A retrospective waiver of consent was obtained.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement Data are available on reasonable request.

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