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The use and predictive performance of the Peninsula Health Falls Risk Assessment Tool (PH-FRAT) in 25 residential aged care facilities: a retrospective cohort study using routinely collected data

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Abstract

Background: The Peninsula Health Falls Risk Assessment Tool (PH-FRAT) is a validated and widely applied tool in residential aged care facilities (RACFs) in Australia. However, research regarding its use and predictive performance is limited. This study aimed to determine the use and performance of PH-FRAT in predicting falls in RACF residents.

Methods: A retrospective cohort study using routinely-collected data from 25 RACFs in metropolitan Sydney, Australia from Jul 2014-Dec 2019. A total of 5888 residents aged \geq 65 years who were assessed at least once using the PH-FRAT were included in the study. The PH-FRAT risk score ranges from 5 to 20 with a score > 14 indicating fallers and \leq 14 non-fallers. The predictive performance of PH-FRAT was determined using metrics including area under receiver operating characteristics curve (AUROC), sensitivity, specificity, sensitivity_{Event Rate(ER)} and specificity_{ER}.

Results: A total of 27,696 falls were reported over 3,689,561 resident days (a crude incident rate of 7.5 falls /1000 resident days). A total of 38,931 PH-FRAT assessments were conducted with a median of 4 assessments per resident, a median of 43.8 days between assessments, and an overall median fall risk score of 14. Residents with multiple assessments had increased risk scores over time. The baseline PH-FRAT demonstrated a low AUROC of 0.57, sensitivity of 26.0% (sensitivity_{ER} 33.6%) and specificity of 88.8% (specificity_{ER} 82.0%). The follow-up PH-FRAT assessments increased sensitivity_{ER} values although the specificity_{ER} decreased. The performance of PH-FRAT improved using a lower risk score cut-off of 10 with AUROC of 0.61, sensitivity of 67.5% (sensitivity_{ER} 74.4%) and specificity of 55.2% (specificity_{ER} 45.6%).

Conclusions: Although PH-FRAT is frequently used in RACFs, it demonstrated poor predictive performance raising concerns about its value. Introducing a lower PH-FRAT cut-off score of 10 marginally enhanced its predictive performance. Future research should focus on understanding the feasibility and accuracy of dynamic fall risk predictive tools, which may serve to better identify residents at risk of falls.

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Introduction

Falls in older people are a common and challenging health problem causing significant morbidity, mortality and economic burden [1-3]. Approximately one in three people aged 65 years and over fall every year [4]. Globally, the age-standardised prevalence of falls is estimated to be over 5000 per 100,000 people [5]. According to the 2017 global burden of disease study from 195 countries, falls resulted in over half a million deaths, 16.7 million years of life lost, 19.3 million years lived with disability and 35.9 million disability-adjusted life years [5]. The occurrence and consequences of falls are even more concerning in residential aged care facilities (RACFs) [6] (also called care homes, nursing homes or long-term care). In Australia, people aged 65 years and over in RACFs are approximately five times more likely to experience a fall and six times more likely to experience fall-related injury compared to people of the same age in the community [4, 7]. In 2018/19, nearly 11% of permanent residents aged 65 years or over in RACFs were admitted to hospitals or visited emergency departments due to falls [8].

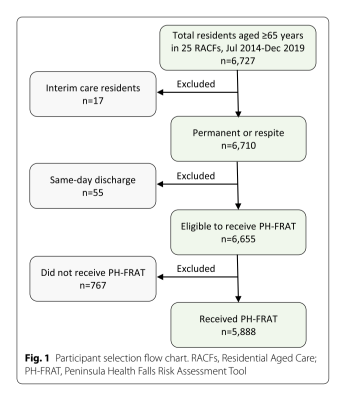
Falls risk assessment tools (FRATs) have been utilised in acute, subacute and aged care settings to help identify those at highest risk of falling [9]. FRATs provide fall risk profiles to predict the likelihood of future occurrences of falls and therefore can play a critical role in targeted fall prevention programs [10]. Several FRATs have been developed over the past two decades [9]. Examples of commonly used FRATs for older people in hospital or community settings include STRAT-IFY (St Thomas's risk assessment tool in falling elderly inpatients) [11], Morse Fall Scale [12], Berg Balance Scale [13] and FROP-Com (falls risk for older people in the community) [14]. A recent systematic review identified fifteen FRATs suitable for use among older adults in RACFs, [15] including the Peninsula Health Falls Risk Assessment Tool (PH-FRAT) [16].

PH-FRAT, developed in 1999 by Peninsula Health in Victoria, Australia, is a validated and easy to use tool that can be used for both screening (early identification of individuals at risk of falls) and assessment (including identification of risk factors) and management strategies for reducing fall risk [17]. The tool was developed using a sample of 291 patients from a single site receiving subacute and residential aged care services. The original validation study reported a moderate predictive performance with a sensitivity of 58.4% and specificity of 90.1% [16]. Although the tool has been widely used in Australia [16, 18], there has been little further evidence of its effectiveness in predicting falls in settings. To the best of our knowledge, only one small study has evaluated its performance in RACFs [19] and there have been no published studies reporting its performance in routine clinical practice. Our aim was to conduct a large pragmatic study to understand the extent of routine use and the performance of PH-FRAT in predicting the occurrence of falls in 25 RACFs managed by one large aged care provider..

Methods

Study design and settings

We conducted a retrospective, longitudinal, cohort study using routinely collected de-identified aged care data from 1 July 2014 to 31 December 2019 extracted from 25 RACFs managed by a large not-forprofit aged care provider in New South Wales (NSW), Australia. We followed the REporting of studies Conducted using Observational Routinely-collected health Data (RECORD) statement [20] when writing this paper.



Participants

The study population flow chart is shown in Fig. 1. The eligibility criteria included residents aged \geq 65 years, who stayed at the facility for more than 24h and had at least one PH-FRAT assessment. We excluded interim care (temporary stay) and same-day discharge (i.e. residents with a length of stay <24h at the facility) residents as these residents stayed in the facility for a short period of time and therefore had lower rates of falls and PH-FRAT assessment data relevant to this study.

Data sources

The data used for this study were sourced from electronic health records used to collect clinical and care management data, data from PH-FRAT assessments and fall incident data. Twenty-five RACFs used the PH-FRAT tool [16] to assess residents' falls risk. Assessments are conducted upon entry to facilities and over time to monitor a residents' falls risk, identify risk factors and create a personalised management plans for high-risk residents. The PH-FRAT comprises three parts: fall risk status, risk factor checklist and action plan. For this study, we focused on the fall risk status data. Fall risk status is calculated by identifying whether four major fall risk factors (recent falls, medications, psychological status and cognitive status) were evident. The total risk score ranges from 5 to 20 with a score of >14 indicating likely fallers and \leq 14 nonfallers. A three-level fall risk classification of low (scores 5-11), medium (12-15) and high (16-20) has been used to identify and provide targeted fall preventive services to high-risk residents [16].

All residents' falls were reported using a standardised incident form containing information on incident date and time, location of the incident, body region injured if there was an injury, and whether transfer to hospital was required. In addition to the fall risk and incident data, we extracted relevant data on socio-demographics (e.g. age, gender, country of birth), length of stay and clinical characterisitics (e.g. fall history at admission, health status).

Statistical methods

Descriptive statistics including medians with inter-quartile ranges (IQR) were reported. We compared the characteristics of included and excluded participants using χ^2 statistics for categorical variables and the Wilcoxon rank-sum test for continuous variables. To determine the predictive performance of the tool, the PH-FRAT scores were compared against the actual (observed) falls. The actual (observed) falls data were obtained from the facility incident form. We limited the follow-up period to the first six months after the PH-FRAT assessment as the tool was not designed for a long-term prediction. Observed falls were defined as the occurrence of any fall regardless of whether an injury was involved or hospitalisation required. Initially, we evaluated the performance of PH-FRAT using resident's baseline risk status. Given PH-FRAT can be reapplied multiple times and its performance may change over time, we also reported the performance of the tool using the risk status at follow-up assessments. We present the results of the second to fifth applications of the tool. As we aimed to assess the performance of the tool in predicting falls, falls that occurred before the completion of the PH-FRAT assessment were excluded from the analysis. For instance, when evaluating the performance of the second PH-FRAT application against the occurrence of falls, only falls that occurred after the completion of the second PH-FRAT were included in the analysis.

The predictive performance of PH-FRAT was determined using commonly used performance metrics including sensitivity, specificity, positive predictive value (PPV), negative predictive value (NPV) and area under receiver operating characteristics (ROC) curve (AUROC) along with their 95% confidence intervals (CI). AUROC is an indicator of the discriminatory power of a given tool [21] (that is the ability of PH-FRAT to efficiently discriminate between fallers and non-fallers in this case). The AUROC values range from 0.5 to 1 with values from 0.5-0.6, 0.6-0.7, 0.7-0.8, 0.8-0.9 and 0.9-1.0 suggesting respectively poor, sufficient, good, very good and excellent discrimination [21]. We also report Youden's index (sensitivity + specificity -1), a point on ROC curves that is farthest away from the diagonal/reference line indicating optimal sensitivity and specificity of a given tool [21, 22]. Youden's index ranges from 0 (no discrimination) to 1 (perfect discrimination) [21, 22].

We used two methods to calculate the performance measures including the standard and *event rate* (ER) methods. The formulas used to calculate these measures using both methods are presented in Supplementary Table 1. The *event rate* method was based on an approach proposed previously [23] and is the preferred method as it accounts for the recurrent nature of falls. The 95% CIs for sensitivity_{ER} and specificity_{ER} were determined using a bootstrapping technique as described by Haines et al. [23].

In addition to the original cut-off value of 14 to define fall risk status, we also determined a cut-off that best fitted our data. We used the highest Youden's index_{ER} (a value on ROC curves associated with the highest sensitivity and specificity) to determine the optimal cut-off [22]. We report all performance measures of the new cutpoint in a similar way as the original cut-point. A subgroup analysis by resident type (permanent vs respite) was also reported. All *p*-values were 2-tailed and alpha

was set at P < 0.05. Analysis was conducted using Stata version 16 (StataCorp LP, College Station, TX).

Results

Participants

Of the total 6727 residents, 839 were excluded from the study (767 did not receive the tool, 55 had a same-day discharge and 17 were interim care residents). Of the 6655 eligible residents, 5888 (88.5%) received at least one PH-FRAT assessment (Fig. 1). Table 1 compares the baseline and follow-up characteristics of the included (n = 5888) and excluded (n = 839) participants. Except

Table 1 Comparison of included and excluded participar
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	Included (<i>n</i> = 5888)	Excluded (<i>n</i> = 839)	P-value
Gender, n(%)			
Male	2006 (34.1)	316 (37.7)	0.038
Female	3882 (65.9)	522 (62.3)	
Age at admission, median (IQR)	86 (81–90)	86 (80–90)	0.352
Age at admission, mean (SD)	84.9 (7.7)	84.6 (8.2)	
Age category in year, n (%)			
65–74	617 (10.5)	107 (12.7)	0.215
75–84	1822 (30.9)	259 (30.9)	
85–94	2993 (50.8)	415 (49.5)	
≥95	456 (7.8)	58 (6.9)	
Country of birth, n (%)			
Australia	3737 (63.5)	437 (52.1)	< 0.001
Other country	2151 (36.5)	402 (47.9)	
Resident status at the end of the stud	dy ^a , n (%)		
Active	2390 (40.6)	591 (70.4)	< 0.001
Deceased	3498 (59.4)	248 (29.6)	
Resident type, n (%)			
Permanent admission ^b	4794 (81.4)	331 (39.5)	< 0.001
Respite admission only	1094 (18.6)	508 (60.5)	
Selected health status, n (%)			
Dementia	3044 (51.7)	234 (27.9)	< 0.001
Depression, mood and affective disorders	2479 (42.1)	177 (21.1)	< 0.001
Cognitive impairment	1980 (33.6)	178 (21.2)	< 0.001
Anxiety and stress-related disor- ders	1893 (32.2)	100 (11.9)	< 0.001
Cerebrovascular accident	1494 (25.4)	129 (15.4)	< 0.001
Diabetes mellitus	1331 (22.6)	159 (19.0)	0.017
Visual impairment	984 (16.7)	74 (8.8)	< 0.001
Delirium	593 (10.1)	33 (3.9)	< 0.001
Parkinson's disease	442 (7.5)	40 (4.8)	0.004

 χ 2 was used to compare the categorical variables and the Wilcoxon rank-sum tests was used to compare the continuous variables respectively. ^aThe status of the 77 residents (same-day discharge or interim care) were recorded at the time of discharge or end of the follow-up period. ^bIncluded some residents who had respite admission before permanent admission

for age, all other characteristics were significantly different between the included and excluded participants. Included participants had a higher prevalence of comorbidities, were born in Australia, and died before the end of the study. The median age in the included participants was 86 (IQR 81–90) and 65.9% were female (Table 1).

PH-FRAT utilisation

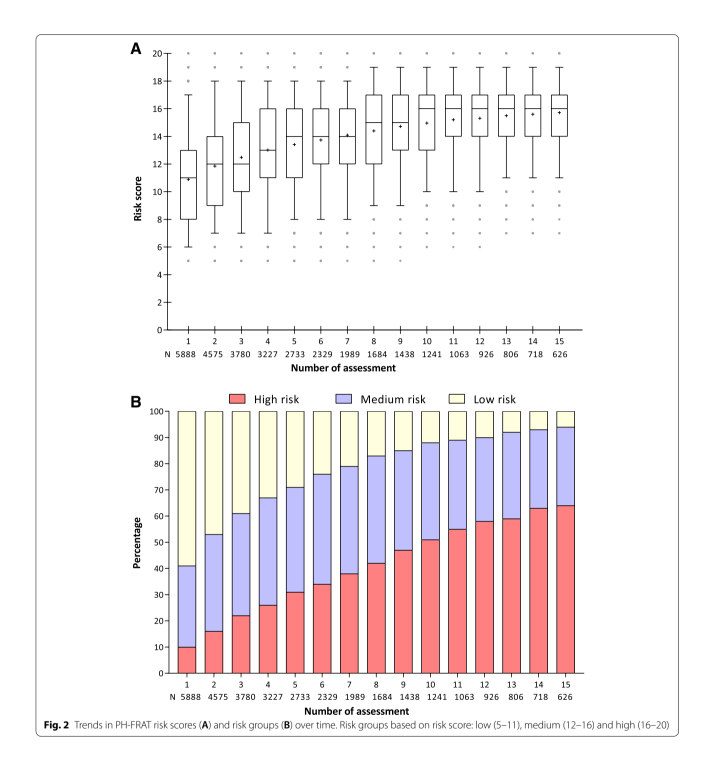
For the 5888 residents who received at least one PH-FRAT assessment, 38,931 assessments were performed with a median of 4 assessments per resident (IQR 2–8; range 1–107). For residents with multiple PH-FRAT assessments, the median time between assessments was 43.8 days (IQR 10.7–144.0). The overall median PH-FRAT risk score was 14 (IQR 11–16).

Figure 2 shows the risk scores and risk groups for residents' first and subsequent PH-FRAT assessments. We reported the first fifteen assessments for residents receiving PH-FRAT on multiple occasions. There was an upward trend in risk scores from first to the 10th assessment (increasing from median scores of 11 to 16) and then little change after the 10th assessment. The proportion of residents classified as high risk (risk score of 16-20) increased consistently over time from 10.0% at the first assessment to 63.7% at assessment fifteen (Fig. 2). It is important to note that having a fall in RACFs increases the subsequent PH-FRAT score as a history of 'recent falls' is one of the components of the PH-FRAT assessment [16]. For example, having a fall in the past 3 months while in a RACF automatically increases the subsequent PH-FRAT score by 8 points [16].

PH-FRAT performance

A total of 27,696 falls were reported over 3,689,561 resident days (a crude incident rate of 7.5 falls/1000 resident days; 95% CI 7.4–7.6) for the 5888 residents during the entire follow-up period. Of the 27,696 fall incidents, 26,448 (95.5%) were reported after a baseline PH-FRAT assessment was completed. Sixty-two percent of residents (n = 3627) experienced at least one fall after the baseline PH-FRAT assessments. In the first 6 months, there were 7487 falls over 820,752 resident days (a crude incident rate of 9.1 falls/1000 resident days; 95% CI 8.9–9.3).

Table 2 presents the performance of baseline PH-FRAT assessments to accurately categorise residents as 'fallers' or 'non fallers' by comparing against the actual (observed) occurrence of falls after limiting the follow-up period to the first 6 months of PH-FRAT assessment. The ROC curve is shown in Supplementary Figure 1. Using a risk score cut-off of 14 (recommended by the original developer) [16], the AUROC was 0.57 indicating poor predictive performance. The tool had



high specificity (88.8%; 95% CI 87.8–89.9) indicating the tool accurately predicted 88.8% of residents as unlikely to fall but had a low sensitivity (26.0%; 95% CI 24.3–27.8) in that it was only able to predict 26.0% of residents who were likely to fall. When the *event rate* method was utilised, the sensitivity_{FR} increased to

33.6% (95% CI 30.3–36.8) while maintaining a moderately high specificity_{ER} of 82.0% (95% CI 80.9–83.1).

By examining the ROC curve (Figure S1) we were able to identify the cut-off point at which PH-FRAT presents the optimal sensitivity and specificity. The highest Youden's index was obtained at a risk score cut-off

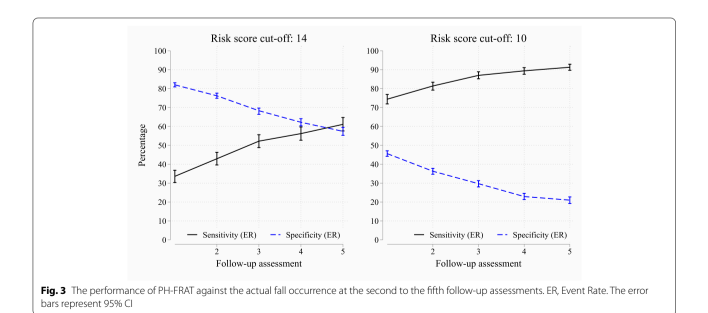
PH-FRAT Predicted Falls	Risk score cut-off 14				Risk score cut-off 10			
	Observed Falls		No. of	Resident	Observed Falls		No. of	Resident
	Fallers	Non-fallers	falls	days	Fallers	Non-fallers	falls	days
Fallers	627	388	2512	147,454	1626	1559	5570	446,080
Non-fallers	1782	3091	4975	673,298	783	1920	1917	374,672
Total	2409	3479	7487	820,752	2409	3479	7487	820,752
Standard method								
AUROC (95% CI)	0.57 (0.56–0.59)			0.61 (0.60–0.63)				
Sensitivity (95% CI)	26.0 (24.3–27.8)			67.5 (65.6–69.4)				
Specificity (95% CI)	88.8 (87.8–89.9)			55.2 (53.5–56.9)				
PPV (95% CI)	61.8 (58.7–64.8)			51.1 (49.3–52.8)				
NPV (95% CI)	63.4 (62.1–64.8)			71.0 (69.3–72.7)				
Youden's index	0.148			0.227				
Event rate method								
Sensitivity _{ER} (95% CI)	33.6 (30.3-	-36.8)			74.4 (71.9-	-76.9)		
Specificity _{ER} (95% CI)	82.0 (80.9-	-83.1)			45.6 (44.2-	-47.1)		
Youden's index _{ER}	0.156				0.201			

Table 2 The performance of baseline PH-FRAT against the actual fall occurrence within the first six months of PH-FRAT assessment

AUROC Area Under Receiver Operating Characteristics curve, PPV Positive Predictive Value, NPV Negative Predictive Value. AUROC interpretation: poor (0.5–0.6), sufficient (0.6–0.7), good (0.7–0.8), very good (0.8–0.9) and excellent discrimination (0.9–1.0). Youden's index ranges from 0 (no discrimination) to 1 (perfect discrimination)

of 10 (a score of >10 indicating fallers and \leq 10 non-fallers). Using this cut-off, the tool had an AUROC of 0.61, the sensitivity improved significantly to 67.5% (95% CI 65.6–69.4) with a specificity of 55.2% (95% CI 53.5–56.9). When the *event rate* method was utilised, sensitivity further increased to 74.4% (95% CI 71.9–76.9) and specificity_{ER} dropped to 45.6% (95% CI 44.2–47.1) (Table 2).

Figure 3 presents the sensitivity_{ER} and specificity_{ER} of PH-FRAT using the follow-up assessments (i.e. after residents have been reassessed on multiple occasions). The results from the second to the fifth assessments are presented. For both cut-off values, the sensitivity_{ER} improved with subsequent assessments, while the specificity_{ER} decreased. For instance, using the cut-off value of 14, the sensitivity_{ER} increased from 42.9% at the second



assessment to 61.1% at the fifth assessment, while the specificity_{FR} decreased from 76.2 to 57.4% (Fig. 3).

Sub-group analysis

A subgroup analysis by resident type showed no major difference in the performance of PH-FRAT by resident type. The AUROC for respite vs permanent residents was 0.55 vs 0.57 (using a cut-off of 14) and 0.61 for both groups (using a cut-off of 10). The performance of PH-FRAT was slightly better in respite compared to permanent residents when the *event rate* method was used (Youden's index_{ER} of 0.224 vs 0.150 using a cut-off of 14 and 0.312 vs 0.191 using a cut-off of 10) (Table 3).

Discussion

Key findings

This study is one of very few studies to describe the utilisation pattern of PH-FRAT and determine its predictive performance against actual falls observed in RACFs in a large sample. We retrospectively evaluated the frequency of use and ability of the PH-FRAT risk assessment tool to predict falls among nearly 6000 residents in 25 RACFs. We found the PH-FRAT was frequently used with 89% of eligible residents receiving at least one assessment. However, the predictive performance of the tool was poor, accurately predicting a fall (within 6 months of PH-FRAT assessment) in only 33.6% of residents. With subsequent assessments, sensitivity improved but the specificity decreased. By changing the cut-off score by which the PH-FRAT categorises 'fallers' and 'non-fallers' from 14 to 10, the sensitivity and specificity of the PH-FRAT changed to 74 and 46% respectively.

Interpretation and comparison with existing literature

The predictive performance PH-FRAT in our study was lower than that reported in the original validation study [16]. Stapleton et al. [16] reported a moderate predictive performance of the tool with a sensitivity of 58.4% (sensitivity_{ER} 68.8%), specificity of 90.1% (specificity_{ER} 70.2%) and Youden's index of 0.49 (Youden's index_{ER} 0.39). The main reason for the difference may be related to the dissimilar characteristics of our study population and that of the origin study [16]. In the original study (n = 291), only 20% of the patients were from nursing homes while the remainder were admitted for rehabilitation services (60%) or to hostels (20%) unlike in our study where all patients were from RACFs. Our study population was also older (mean age of 85.2 vs 79.9 years) with higher levels of comorbidities compared to the population used to develop the tool. It is likely that the original study overstated their predictive accuracy due to the retrospective validation design [24], a limitation recognised by the original developers [16].

External validation of FRATs beyond the original data is fundamental to establishing their generalizability. However, external validation is rarely performed, and in the domain of falls, only a small number of prediction models for community-dwelling older adults have been externally validated [25, 26], showing modest predictive accuracy [26–28]. Barker et al. [19] conducted a prospective external validation study to evaluate the psychometric properties (predictive, evaluative and discriminative validity) of four FRATs widely utilised in Australian RACFs including the PH-FRAT, Queensland Fall Risk Assessment Tool (QFRAT), Melbourne Fall Risk

	Risk score cut-off 14		Risk score cut-off 10			
	Permanent (<i>n</i> = 4794)	Respite (<i>n</i> = 1094)	Permanent (<i>n</i> = 4794)	Respite (<i>n</i> = 1094)		
Standard method						
AUROC (95% CI)	0.57 (0.56–0.58)	0.55 (0.52–0.58)	0.61 (0.59–0.62)	0.61 (0.57–0.65)		
Sensitivity (95% CI)	26.9 (25.1–28.8)	16.0 (11.1–21.9)	68.1 (66.1–70.1)	60.3 (53.1–67.2)		
Specificity (95% CI)	87.0 (85.7–88.3)	94.0 (92.2–95.5)	53.0 (51.1–54.9)	61.4 (58.2–64.6)		
PPV (95% CI)	66.1 (60.9–67.2)	36.5 (26.3–47.6)	55.5 (53.6–57.3)	25.2 (21.3–29.4)		
NPV (95% CI)	58.1 (56.5–59.7)	83.8 (81.4–86.1)	65.9 (63.9–68.0)	87.8 (85–90.2)		
Youden's index	0.139	0.100	0.211	0.217		
Event rate method						
Sensitivity _{ER} (95% CI)	33.7 (30.4–37.0)	30.8 (15.6–45.9)	74.6 (72.0–77.2)	71.3 (61.9–80.7)		
Specificity _{ER} (95% CI)	81.3 (80.1–82.4)	91.6 (89.1–94.1)	44.5 (43.0–46.0)	59.9 (55.5–64.2)		
Youden's index _{ER}	0.150	0.224	0.191	0.312		

Table 3 The performance of baseline PH-FRAT assessments compared to predict actual fall occurrence for respite and permanent residents within the first six months of PH-FRAT assessment

AUROC Area Under Receiver Operating Characteristics curve, PPV Positive Predictive Value, NPV Negative Predictive Value. AUROC interpretation: poor (0.5–0.6), sufficient (0.6–0.7), good (0.7–0.8), very good (0.8–0.9) and excellent discrimination (0.9–1.0). Youden's index ranges from 0 (no discrimination) to 1 (perfect discrimination)

Assessment Tool (MFRAT), and the Falls Assessment Risk and Management Tool (FARAM). Although the study had a small sample consisting of 87 aged care residents, multiple sites including nursing homes and hostels were used. All tools exhibited poor psychometric properties. PH-FRAT had moderate sensitivity (52%), specificity (66%) and a Youden's index (0.18) which was comparable with our findings when a risk score cut-off 10 was used. However, when the researchers compared the predictive accuracy of the tools against a single screening question *"has the resident fallen in the past 12 months?"*, all four tools performed no better than the screening question.

Our finding that the sensitivity_{ER} values increased while specificity_{ER} decreased with reapplications of the tool is consistent with the observation by Haines et al. [23] when evaluating the risk of falls among hospitalised patients (n = 316). Haines reported a dramatic increase in sensitivity_{ER} when a fall risk screening tool was reapplied [23]. This is likely to occur due to an increased risk of a resident falling and therefore being classified as true over time, which equates to increased sensitivity_{SF}. As sensitivity (true positive rate) and specificity (true negative rate) are reciprocal to each other [29], the inverse relationship between sensitivity_{FR} and specificity_{FR} values over multiple assessments is not surprising. Additionally, given specificity values tend to decrease with an increasing prevalence of disease [30], a possible increase in the prevalence of falls over time might have contributed to the decrease in the specificity_{ER} values at follow-up times.

Implications for clincal practice and recommendations for further research

PH-FRAT is one of the most frequently used tools in RACFs in Australia [15] and is currently adopted for inclusion in subacute fall prevention guidelines by the Australian Commission on Safety and Quality in Health Care [4]. Previous studies have shown its use to be highly feasible taking only 2-3 min to complete and it has demonstrated good inter-rater reliability [16, 19]. The high uptake of PH-FRAT in RACFs confirms its feasibility in routine clinical practice. However, PH-FRAT might need to be updated to reflect the current aged care population profile. Indeed, the poor predictive performance of PH-FRAT raises practical concerns about its utility and on whether it may be contributing to poor or untimely care decisions [31, 32]. This has safety implications as potentially high-risk residents eligible for specific fall prevention programs are likely to miss out on receiving the intervention due to incorrect risk profiling. We found that reducing the risk score cut-off point from 14 to 10 substantially improved the tool's performance with a dramatic increase in the sensitivity_{ER} from 33.6 to 74%. We recommend using a revised cut-off score of 10 to define falls risk status – for example – a risk score of >10 to indicate high risk and \leq 10 low risk. For RACFs already using the tool, this change can be implemented easily given the current procedures already in place.

However, it is important to emphasise that most predictive tools, in general, do not perform well outside the original study population [33]. A study of 31 clinical prediction models that were externally validated, only 6 showed a comparable predictive performance in the validation studies [34]. This could be due to possible differences in health care practices and population charactistics [31]. Indeed, predictive factors relevant in the original population may no longer be applicable in another setting. For instance, of the items included in PH-FRAT that were predictive of falls in the original study, only fall history and psychological status predicted actual falls in a study by Barker et al. [19] whereas increasingly recognised risk factors for falls such as walking aids use and certain medical conditions (e.g. Parkinson's disease) were not included in PH-FRAT [35].

Updating PH-FRAT through model recalibration by incorporating any new, relevant predictors may enhance the predictive performance of the tool. Although opinion varies regarding modifying the existing tool by incorporating new predictors among researchers [32], recalibration can help to optimise the predictive performance of a tool in a new setting [31].

Future research should also focus on understanding the feasibility and effectiveness of dynamic fall risk prediction models using routinely collected data which can reflect contemporary changes in residents' risk factors. The existing FRATs used in RACFs [15] have been based on a static prediction using input variables collected at a single time point without incorporating the potential changes in the status of input variables over time. A model that incorporates all potentially useful information about input variables on an ongoing basis (dynamic prediction) could a play critical role in improving the prediction of falls. Dynamic predictions involve the use of real-time or near real-time data to enable up-to-date risk predictions. In long-term care settings such as RACFs risk factors for falls (e.g., medication utilisation) changes over time. However, electronic data containing relevant fall risk factors including both time-invariant (e.g. demographic) and time-varying (e.g. medication) factors are now collected as part of routine care in RACFsproviding unique opportunities to develop and test dynamic falls risk prediction tools [36, 37]. Several studies have identified that certain medications that are used for the treatment of conditions affecting cardiovascular (e.g., beta-blockers, diuretics) or central nervous systems (e.g., antipsychotics, sedatives) are known to increase the risk of falling [38–45]. As older people in RACFs are the primary users

of these medications, it is important to utilise medication data as one of the main time-varying factors to obtain a robust and accurate dynamic prediction and monitoring of falls risk over time. Advanced statistical methods such as *joint models* [46], *landmark models* [47] and deep learning-based machine learning approaches [48] have previously been utilised to develop dynamic prediction models in other settings.

Strengths and limitations

This is the first study to evaluate the use and performance PH-FRAT using routinely collected aged care data. The strength of our study lies in the methodology. Firstly, our study is a multi-centre study that involved a large sample of nearly 6000 residents from 25 RACFs. Secondly, unlike previous studies [16, 19], we utilised a longitudinal cohort design with long-term follow-up which allowed us to track residents over a combined period of greater than 2.5 million resident days. Routinely collected data has the added advantage of not being influenced by the study aims, minimal selection bias as a loss to follow-up or non-response is not an issue, and not subject to recall bias and differential misclassification [49]. Finally, in addition to reporting the performance metrics using the standard method, we utilised a modified approach based on event rate which serves a better indicator for recurrent events like falls [23].

The main limitation of this study was our focus on part 1 of the tool (risk score), which meant that the potential effect of subsequent fall prevention interventions was not accounted for in the current analysis. In addition to the risk score, PH-FRAT provides possible actions that can be implemented to prevent falls. For instance, residents that were predicted to be in a high-risk group at baseline might have received targeted fall prevention interventions and thus potentially decreased their risk of falling. This can confound the relationship between PH-FRAT and the likelihood of falls occurring. Our study was further restricted to RACFs in metropolitan areas and one aged care provider only, thus our findings may not be representative of all RACFs.

Conclusion

In conclusion, although PH-FRAT is frequently used in RACFs, it demonstrated poor predictive performance against the occurrence of actual falls. This raises concerns about its utility and value and may be preventing some residents from gaining access to necessary fall prevention interventions. Reducing the PH-FRAT score to a lower cut-off value of 10 may optimise its predictive performance. Future research should also focus on understanding and improving the feasibility and effectiveness of dynamic fall risk predictive tools using routinely collected electronic aged care data to address the underlying limitation of static falls risk assessment tools.

Abbreviations

AUROC: Area under receiver operating characteristics curve; CI: Confidence interval; ER: Event rate; FARAM: Falls Assessment Risk and Management Tool; FRATs: Falls risk assessment tools; FROP-Com: Falls risk for older people in the community; IQR: Inter-quartile ranges; MFRAT: Melbourne Fall Risk Assessment Tool; NPV: Negative predictive value; NSW: New South Wales; PH-FRAT : Peninsula Health Falls Risk Assessment Tool; PV: Positive predictive value; QFRAT: Queensland Fall Risk Assessment Tool; RACFs: Residential aged care facilities; ROC: Receiver operating characteristics curve; STRATIFY: St Thomas's risk assessment too in falling elderly inpatients.

Supplementary Information

The online version contains supplementary material available at https://doi. org/10.1186/s12877-022-02973-0.

Additional file 1: Table S1. Measures of performance of PH-FRAT against actual fall occurrence.

Additional file 2: Figure S1. ROC curve for the baseline PH-FRAT in predicting falls in six month.

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Informed consent

This is a retrospective cohort study utilising, de-identified, existing routinely collected aged care data. A waiver of informed consent was granted by Macquarie University Human Research Ethics Committee following the Australian National Statement on Ethical Conduct in Human Research section 3.3.14 (a) as the study used previously collected de-identified data.

Authors' contributions

Conception– NW; Design and data analysis – NW with input from other team members; Data interpretation – all authors; Initial manuscript drafting – NW; Critical revision – all authors; Final approval for publication – all authors.

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Availability of data and materials

The data that support the findings of this study are available from the aged care provider (Anglicare), but restrictions apply to the availability of these data, which were used under license for the current study, and so are not publicly available. Data are however available from the authors upon reasonable request and with permission of Anglicare.

Declarations

Ethics approval and consent to participate

The study was reviewed and approved by Macquarie University Human Research Ethics Committee (reference number, 6144). The study was performed in accordance with the principles of the Declaration of Helsinki.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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References

- 1. Berková M, Berka Z. Falls: a significant cause of morbidity and mortality in elderly people. Vnitrni lekarstvi. 2018;64(11):1076–83.
- 2. Peel NM. Epidemiology of falls in older age. Can J Aging. 2011;30(1):7–19.
- Watson W, Clapperton A, Mitchell R. The burden of fall-related injury among older persons in New South Wales. Aust N Z J Public Health. 2011;35(2):170–5.
- Australian Commission on Safety and Quality in Health Care. Preventing falls and harm from falls in older people: best practice guidelines for Australian residential aged care facilities. Sydney: Commonwealth of Australia; 2009.
- James SL, Lucchesi LR, Bisignano C, Castle CD, Dingels ZV, Fox JT, et al. The global burden of falls: global, regional and national estimates of morbidity and mortality from the global burden of disease study 2017. Injury Prevention. 2020;26(Suppl 2):i3.
- Rapp K, Becker C, Cameron ID, König HH, Büchele G. Epidemiology of falls in residential aged care: analysis of more than 70,000 falls from residents of bavarian nursing homes. J Am Med Dir Assoc. 2012;13(2):187.e181–6.
- AlHW. Pointer S 2019. Trends in hospitalised injury due to falls in older people, 2007–08 to 2016–17. Injury research and statistics series no. 126. Cat. No. INJCAT 206. AlHW: Canberra; 2019.
- Australian Commission on Safety and Quality in Health Care. Hospitalisations in Australian aged care: 2014/15–2018/19. Sydney: Commonwealth of Australia; 2021.
- 9. Park SH. Tools for assessing fall risk in the elderly: a systematic review and meta-analysis. Aging Clin Exp Res. 2018;30(1):1–16.
- Walker GM, Armstrong S, Gordon AL, Gladman J, Robertson K, Ward M, et al. The falls in care home study: a feasibility randomized controlled trial of the use of a risk assessment and decision support tool to prevent falls in care homes. Clin Rehabil. 2016;30(10):972–83.
- Oliver D, Britton M, Seed P, Martin FC, Hopper AH. Development and evaluation of evidence based risk assessment tool (STRATIFY) to predict which elderly inpatients will fall: case-control and cohort studies. BMJ (Clin Res). 1997;315(7115):1049–53.
- 12. Morse JM, Morse RM, Tylko SJ. Development of a scale to identify the fallprone patient. Canadian Journal on Aging. 1989;8(4):366–77.
- Berg K, Wood-Dauphine S, Williams J, Gayton D. Measuring balance in the elderly: preliminary development of an instrument. Physiother Can. 1989;41(6):304–11.
- Russell MA, Hill KD, Day LM, Blackberry I, Gurrin LC, Dharmage SC. Development of the falls risk for older people in the community (FROP-com) screening tool. Age Ageing. 2009;38(1):40–6.
- Nunan S, Brown Wilson C, Henwood T, Parker D. Fall risk assessment tools for use among older adults in long-term care settings: a systematic review of the literature. Aust J Ageing. 2018;37(1):23–33.
- Stapleton C, Hough P, Oldmeadow L, Bull K, Hill K, Greenwood K. Fouritem fall risk screening tool for subacute and residential aged care: the first step in fall prevention. Aust J Ageing. 2009;28(3):139–43.
- 17. Department of Health & Human Services. Falls Risk Assessment Tool (FRAT) instructions. Melbourne, Victoria. Available at: https://training.aacs. com.au/wp-content/uploads/2016/08/Falls-Risk-Assessment-Tool-FRAT. pdf.
- Sax Institute. Translational research framework: testing innovation in policy, programs and service delivery. Available at: https://www.saxin

stitute.org.au/wp-content/uploads/Translational-Research-Framework. pdf. [Accessed 12 Jan 2022].

- 19. Barker AL, Nitz JC, Low Choy NL, Haines T. Measuring fall risk and predicting who will fall: Clinimetric properties of four fall risk assessment tools for residential aged care. The J Gerontol Series A. 2009;64A(8):916–24.
- Benchimol El, Smeeth L, Guttmann A, Harron K, Moher D, Petersen I, et al. The REporting of studies conducted using observational routinely-collected health data (RECORD) statement. PLoS Med. 2015;12(10):e1001885.
- 21. Šimundić A-M. Measures of diagnostic accuracy: basic definitions. EJIFCC. 2009;19(4):203–11.
- Perkins NJ, Schisterman EF. The inconsistency of "optimal" cutpoints obtained using two criteria based on the receiver operating characteristic curve. Am J Epidemiol. 2006;163(7):670–5.
- Haines TP, Hill KD, Bennell KL, Osborne RH. Recurrent events counted in evaluations of predictive accuracy. J Clin Epidemiol. 2006;59(11):1155–61.
- 24. Haines TP, Hill K, Walsh W, Osborne R. Design-related bias in hospital fall risk screening tool predictive accuracy evaluations: systematic review and meta-analysis. J Gerontol A Biol Sci Med Sci. 2007;62(6):664–72.
- Hendrich AL, Bufalino A, Groves C. Validation of the Hendrich II fall risk model: the imperative to reduce modifiable risk factors. Appl Nurs Res. 2020;53:151243.
- Vlaeyen E, Poels J, Colemonts U, Peeters L, Leysens G, Delbaere K, et al. Predicting falls in nursing homes: a prospective multicenter cohort study comparing fall history, staff clinical judgment, the care home falls screen, and the fall risk classification algorithm. J Am Med Dir Assoc. 2021;22(2):380–7.
- Gates S, Smith LA, Fisher JD, Lamb SE. Systematic review of accuracy of screening instruments for predicting fall risk among independently living older adults. J Rehabil Res Dev. 2008;45(8):1105–16.
- Barry E, Galvin R, Keogh C, Horgan F, Fahey T. Is the timed up and go test a useful predictor of risk of falls in community dwelling older adults: a systematic review and meta-analysis. BMC Geriatr. 2014;14:14.
- Parikh R, Mathai A, Parikh S, Chandra Sekhar G, Thomas R. Understanding and using sensitivity, specificity and predictive values. Indian J Ophthalmol. 2008;56(1):45–50.
- Leeflang MMG, Rutjes AWS, Reitsma JB, Hooft L, Bossuyt PMM. Variation of a test's sensitivity and specificity with disease prevalence. CMAJ. 2013;185(11):E537–44.
- Kappen TH, van Klei WA, van Wolfswinkel L, Kalkman CJ, Vergouwe Y, Moons KGM. Evaluating the impact of prediction models: lessons learned, challenges, and recommendations. Diagn Prog Res. 2018;2(1):11.
- 32. Ramspek CL, Jager KJ, Dekker FW, Zoccali C, van Diepen M. External validation of prognostic models: what, why, how, when and where? Clin Kidney J. 2021;14(1):49–58.
- McKechnie D, Pryor J, Fisher MJ. Predicting falls: considerations for screening tool selection vs. screening tool development. J Adv Nurs. 2016;72(9):2238–50.
- Siontis GCM, Tzoulaki I, Castaldi PJ, Ioannidis JPA. External validation of new risk prediction models is infrequent and reveals worse prognostic discrimination. J Clin Epidemiol. 2015;68(1):25–34.
- Deandrea S, Bravi F, Turati F, Lucenteforte E, La Vecchia C, Negri E. Risk factors for falls in older people in nursing homes and hospitals. A systematic review and meta-analysis. Arch Gerontol Geriatr. 2013;56(3):407–15.
- Seaman KL, Jorgensen ML, Raban MZ, Lind KE, Bell JS, Westbrook JI. Transforming routinely collected residential aged care provider data into timely information: Current and future directions. Aust J Ageing. 2021;40(3):e262–8.
- Ludlow K, Westbrook J, Jorgensen M, Lind KE, Baysari MT, Gray LC, et al. Co-designing a dashboard of predictive analytics and decision support to drive care quality and client outcomes in aged care: a mixed-method study protocol. BMJ Open. 2021;11(8):e048657.
- Leipzig RM, Cumming RG, Tinetti ME. Drugs and falls in older people: a systematic review and Meta-analysis: I. Psychotropic Drugs. J Am Geriatr Soc. 1999;47(1):30–9.
- Leipzig RM, Cumming RG, Tinetti ME. Drugs and falls in older people: a systematic review and Meta-analysis: II. Cardiac and analgesic drugs. J Am Geriatr Soc. 1999;47(1):40–50.
- Woolcott JC, Richardson KJ, Wiens MO, Patel B, Marin J, Khan KM, et al. Meta-analysis of the impact of 9 medication classes on falls in elderly persons. Arch Intern Med. 2009;169(21):1952–60.

- Hartikainen S, Lönnroos E, Louhivuori K. Medication as a risk factor for falls: critical systematic review. J Gerontol Series A. 2007;62(10):1172–81.
- Hartikainen S, Lönnroos E. Use of sedatives and hypnotics, antidepressants and benzodiazepines in older people significantly increases their risk of falls. Evid Based Med. 2010;15(2):59.
- de Vries M, Seppala LJ, Daams JG, van de Glind EMM, Masud T, van der Velde N, et al. Finish group on fall-risk-increasing D: fall-risk-increasing drugs: a systematic review and Meta-analysis: I. cardiovascular drugs. J Am Med Dir Assoc. 2018;19(4):371.e371–9.
- Seppala LJ, Wermelink A, de Vries M, Ploegmakers KJ, van de Glind EMM, Daams JG, et al. Fall-risk-increasing drugs: a systematic review and Metaanalysis: II. Psychotropics. J Am Med Dir Assoc. 2018;19(4):371.e311–7.
- 45. Seppala LJ, van de Glind EMM, Daams JG, Ploegmakers KJ, de Vries M, Wermelink A, et al. EUGMS Task and finish group on fall-risk-increasing D: Fall-risk-increasing drugs: a systematic review and Meta-analysis: III. Others. J Am Med Dir Assoc. 2018;19(4):372.e371–8.
- Yuen HP, Mackinnon A, Hartmann J, Amminger GP, Markulev C, Lavoie S, et al. Dynamic prediction of transition to psychosis using joint modelling. Schizophr Res. 2018;202:333–40.
- Musoro JZ, Struijk GH, Geskus RB, Ten Berge I, Zwinderman AH. Dynamic prediction of recurrent events data by landmarking with application to a follow-up study of patients after kidney transplant. Stat Methods Med Res. 2018;27(3):832–45.
- Gupta G, Sunder V, Prasad R, Shroff G. CRESA: a deep learning approach to competing risks, recurrent event survival analysis. In: Advances in knowledge discovery and data mining: 2019// 2019. Cham: Springer International Publishing; 2019. p. 108–22.
- Thygesen LC, Ersboll AK. When the entire population is the sample: strengths and limitations in register-based epidemiology. Eur J Epidemiol. 2014;29(8):551–8.

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