Prognosis tools for short-term adverse events in older emergency department users: result of a Québec observational prospective cohort

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Abstract

Background: The “Program of Research on the Integration of Services for the Maintenance of Autonomy” (PRISMA-7) and “Emergency room evaluation and recommendations” (ER2) are both clinical tools used in Québec Emergency Departments (EDs) for screening of older ED users at higher risk of poor outcomes, such as prolonged length of stay (LOS) in EDs and in hospital. The study aimed to: 1) examine whether the PRISMA-7 and ER2 risk levels were associated with length of stays in ED and hospital, as well as hospital admission; and 2) compare the criteria performance (i.e., sensitivity, specificity, positive predictive value, negative predictive value, likelihood ratios and area under receiver operating characteristic curve) of the PRISMA-7 and ER2 high-risk levels for these three ED adverse events in Québec older patients visiting ED on a stretcher.

Methods: A total of 1905 older patients who visited the ED of the Jewish General Hospital (Montreal, Québec, Canada) on stretchers were recruited in this prospective observational cohort. Upon their ED arrival, PRISMA-7 and ER2 were performed. The outcomes were LOS in ED and in hospital, and hospital admission.

Results: The PRISMA-7 and ER2 risk levels were associated with length of stay in ED and hospital as well as with hospital admission. Prolonged stays and higher hospitalization rates were associated with high-risk levels, whereas those in low-risk level groups had significantly shorter LOS and a lower rate of hospital admission (P < 0.006). While performance measures were poor for both assessment tools, ER2 had a greater prognostic testing accuracy compared with PRISMA-7.

Conclusion: PRISMA-7 and ER2 were both associated with incidental short-term ED adverse events but their overall prognostic testing accuracy was low, suggesting that they cannot be used as prognostic tools for this purpose.

Keywords: Emergency department, Adverse event, Prognosis, Older adults

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Background
Overcrowding and delays in providing care have become increasingly problematic in emergency departments (EDs) in high-income countries [1, 2]. The limited access to primary care in Canada, in part related to the lack of primary care physicians, forces people in the community to seek care in EDs [1, 2]. A continuously enlarging proportion of ED users are aged over 75 and account for up to a quarter of all ED users. These ED users, compared to their younger counterparts, are at higher risk for short-term ED adverse events including prolonged length of stay in ED and in hospital, as well as hospital admission during an index ED visit [2, http://www.csbe.gouv.qc.ca/fileadmin/www/2016/Urgences/CSBE_Rapport_Urgences_2016.pdf]. A major concern of ED healthcare workers is the identification of older patients who are the most vulnerable and are at highest risk of adverse outcomes (e.g. long ED and hospital stay) and treating them in a timely manner [1–4]. The occurrence of adverse events may be reduced if the older ED users’ risk is determined early in the process of ED care (i.e., upon their ED arrival) [1, 3, 4]. One way for preventing short-term ED adverse events is to screen older users with the highest risk and provide timely and individualized geriatric interventions [5–9]. Therefore, the prognostic tools designed for this purpose may provide relevant information regarding potential adverse events early in the ED care plan, with the goal of improving the decision-making process and to reduce adverse events [10–15]. The identification of older patients at risk for short-term ED adverse events is, therefore, an important objective with regards to optimizing the care provided in the EDs. The “Program of Research on the Integration of Services for the Maintenance of Autonomy” (PRISMA-7) is a tool used to assess older ED users in the province of Québec (Canada) [16, 17]. PRISMA-7 was designed to screen disabilities in older community-dwellers and it stratifies them into two risk levels: low versus high-risk for disabilities. Disabilities are the result of chronic diseases burden, among which musculoskeletal and cardiovascular diseases are the main contributors [18]. Regardless the reasons of the ED visit, disabilities and related burden of chronic diseases characterize older ED users and this poorer overall health status may explain in part the high prevalence of the adverse events, suggesting that PRISMA-7 may be used as a prognosis tool to predict short-term ED adverse events [1–4, 16, 17]. To date, the association of the PRISMA-7 high-risk level with the occurrence of short-term ED adverse events has never been examined and therefore validated as a prognosis tool for this purpose. “Emergency room evaluation and recommendations” (ER2) is another clinical tool currently used in Québec EDs [4, 19]. Compared to PRISMA-7, ER2 has been particularly designed for assessing the risk for short-term ED adverse events in older patients [12–14]. The objective of the present study is to determine which clinical tool, between PRISMA-7 and ER2, is the most appropriate tool to use in EDs for detecting the older ED users most at risk for short-term ED adverse events. No previous study has compared PRISMA-7 and ER2 high-risk levels with regards to predicting the occurrence of short-term ED adverse events. Firstly, we hypothesised that both PRISMA-7 and ER2 high-risk levels would be associated with the occurrence of short-term ED adverse events. Disabilities of older ED users and related chronic diseases burden may explain in part the occurrence of the short-term ED adverse events [1–4]. PRISMA-7 screens disabilities, suggesting that it may be used as a prognostic tool of these ED adverse events [16, 17]. Secondly, we hypothesized that ER2 would be a significantly better prognostic tool compared to PRISMA-7 for predicting prolonged length of stay in ED and in hospital, as well as predicting hospital admission, because it was designed and validated for this specific goal unlike PRISMA-7 [4, 12–14, 19]. The study aimed to: 1) examine whether the PRISMA-7 and ER2 high-risk levels were associated with length of stays in ED and in hospital, as well as hospital admission, an 2) compare the performance criteria (i.e., sensitivity, specificity, positive predictive value [PPV], negative predictive value [NPV], likelihood ratios [LR] and area under receiver operating characteristic curve) of the PRISMA-7 and ER2 high-risk levels for the three previously stated ED adverse events in Québec older patients visiting ED on a stretcher.

Methods
Population and design
Older adults who consulted the ED of the Jewish General Hospital (Montreal, Quebec, Canada) between September 2017 and January 2018 were potential participants in this prospective observational cohort study. The inclusion criteria of the present study were: age ≥75, unplanned ED visit, being on a stretcher (non-ambulatory), assessed with PRISMA-7 and ER2 and agree to participate in the study. The exclusion criteria were participation in an experimental study, being in palliative care and transfer to another hospital during the ED visit. A total of 7017 older users aged ≥75 visited the ED of the Jewish General Hospital during the period of recruitment and 1905 (27.2%) among this group satisfied all selection criteria.

Baseline assessment
Firstly, PRISMA-7 was completed by the nurse in charge of the triage for medico-surgical emergencies. The nurse asked patients and/or their family the PRISMA-7 questions, details of PRISMA-7 items have been previously described [17]. The scoring method of PRISMA-7 is to
assign 1 point to the answer yes and no points to the an-
swer no, and to stratify patients into two risk levels: low
(i.e., score 0 to 2) and high (i.e., score 3 to 7) risk [17].

Secondly, once the patient is on a stretcher and the
triage is completed, their nurse completed the ER\textsuperscript{2}
assessment. ER\textsuperscript{2} is composed of 6 close-ended format
questions (i.e., yes versus no) that have been described
previously in detail [4, 19]. Compared to PRISMA-7,
which has a uniform scoring system, the ER\textsuperscript{2} is divided
into major and minor criteria with different assigned
scores. A score of five points is assigned to each major
criteria, which include “use of walking aid” and “temp-
oral disorientation”, whereas for the minor criteria,
which comprise the other four criteria of ER\textsuperscript{2} a score of
1 point is assigned. The range of score is 0 (lowest risk)
to 14 (highest risk). ER\textsuperscript{2} stratifies risk for short-term ED
adverse events into three levels based on obtained score:
low (0-3), moderate [3, 4] and high (≥ 6).

In addition, the following items were recorded using
the patients’ electronic medical record: age, sex, and
place of living (home versus residence versus other). The
reasons for ED visits were categorized into 5 sub-types:
organ failure defined as an acute organ decompensation;
mobility disorders defined as gait and/or balance impair-
ment and/or fall with or without fall-related injuries;
neuropsychiatric disorders defined as delirium, demen-
tia, and/or behavioral disorders; cancer defined as a
group of diseases involving abnormal cell growth with
the potential to invade or spread to other parts of the
body; and social issues defined as the absence of acute
disease symptoms combined with an acute increase in
the use of formal and/or informal home and social ser-
ices leading to an inability to cope at home.

This study has been performed following the usual
procedure of the ED of the Jewish General Hospital.
PRISMA-7 in this hospital is usually performed at initial
triage, which is the first step of assessment of ED users.
The triage is a process with the purpose of prioritizing
patients depending on the nature and urgency of their
health condition and to identify those require urgent
evaluation and treatment. This first step of ED visit as-
esses the type and severity of patient health conditions,
determines access to appropriate treatments and assigns
appropriate human health resources. The Canadian ED
Triage and Acuity Scale was used in this study [20]. This
scale is composed of 5 levels of urgencies which are:
level 1, defined as resuscitation; level 2, defined as emergent;
level 3, defined as urgent; level 4, defined as less urgent; and
level 5, defined as non-urgent. The triage process thus determines whether the ED
users need to be seen as ambulatory patients (level 4
and 5) or on a stretcher (level 2 and 3). There is no
need to perform ER\textsuperscript{2} in ambulatory patient, because
this category of ED users is habitually discharged
home after their ED visit. Since ER\textsuperscript{2} is a tool to
screen older ED users at risk for short-term ED ad-
verse events, it has only been performed in older ED
users with level 2 and 3 who were on stretchers
(non-ambulatory). PRISMA-7 and ER\textsuperscript{2} assessments
were therefore performed at different times, but the
delay between the two assessments was short (less
than 1 h) for older ED users on stretcher.

Follow-up and outcomes
The dates and hours of ED visit, hospital admission
and discharge were extracted from the patients’ digital
file when the patient was discharged from ED or hos-
pital. The length of stay in ED, expressed in hours,
was defined as the delay between the hour and day of
the ED visit and the hour and day of the ED dis-
charge (either to hospital or place of living). This out-
come was used as the primary outcome. The length
of stay in hospital, expressed in days, was defined as
the delay between day of the ED visit and the day of
the hospital discharge. The highest tertile of length of
stay in ED (i.e., > 26.9 hr) and hospital (i.e., > 13 days)
defined a prolonged length of stay. Hospital admission
after an index ED visit was considered when older
ED users were admitted to medical or surgical wards
at the Jewish General Hospital.

Power analysis
The number of participants required for this study has
been calculated to have enough statistical power in order
to compare the high-risk group to the low-risk group for
the primary outcome, ED length of stay, as well as for
the secondary outcomes (length of stay in hospital and
hospital admission). Based on a pilot study, comparisons
to be made between high-risk and low-risk groups are:
the length of stay in ED (23.4 ± 13.5 h versus 18.9 ± 13.1
h), the length of stay in hospital (21.8 ± 17.3 days versus
15.8 ± 14.5 days), and hospital admission (58.6% versus
40.2%). In order to detect differences for these 3 out-
comes with power 90% at the 0.05 significance level, this
study needed 1316 participants for ED length of stay,
1896 for hospital length of stay and 1901 for hospital
admission.

Ethics
This study was performed in accordance with the ethical
standards set forth in the Helsinki Declaration (1983).
All participants recruited in this study provided a verbal
informed consent because the study did not change the
usual clinical practice. The verbal informed consent was
obtained from the patients themselves in the presence of
their trusted person, usually a family member, who
helped them to make decision. Participants or their legal
guardian when appropriate, were informed that their medical information may be used for research purposes. If they disagreed, they informed the physician taking care of them and a note was recorded in their chart. The Research Review Office of the Jewish General Hospital and the Research Ethics Committee of the Integrated Health and Social Services University Network for West-Central Montreal approved this process.

Statistics
Means and standard deviations (SD) or frequencies were used to characterize the participants’ baseline characteristics. Between group-comparisons were performed using analysis of variance (ANOVA) with Bonferroni correction for multiple comparisons, unpaired t-test and Chi-square, as appropriate. Multiple linear and logistic regressions examined the association of the PRISMA-7 and ER² risk levels (used as independent variable with separated model for each tool and risk levels) with length of stay in ED and in hospital, and hospital admission (used as dependent variable with separated model for each variable).

Table 1: Comparison of baseline characteristics of participants on stretcher visiting emergency department separated in different subgroups based on their PRISMA-7 and ER² risk scores (n = 1905)

**Table 1** Comparison of baseline characteristics of participants on stretcher visiting emergency department separated in different subgroups based on their PRISMA-7 and ER² risk scores (n = 1905)

| Variable                        | PRISMA-7 risk | ER² risk | \( P \)-value² |
|---------------------------------|---------------|----------|----------------|
|                                 | Low (n = 885) | High (n = 1020) |               |
|                                 |               |          | Low (n = 722) | Moderate (n = 70) | High (n = 1113) |               |
|                                 |               |          |               |               |               |               |
| Age (years)                     |               |          |               |               |               |               |
| Mean ± SD                       | 84.5 ± 6.0    | 86.1 ± 6.0 | \( < 0.001 \) | 83.2 ± 5.4 | 86.5 ± 5.2 | 86.7 ± 6.1 | \( < 0.001 \) | \( < 0.001 \) | 1.000 |
| ≥ 85 years                      | 406 (45.9)    | 584 (57.3) | \( < 0.001 \) | 250 (34.6) | 55 (78.6) | 685 (61.5) | \( < 0.001 \) | \( < 0.001 \) | 0.004 |
| Male                            | 311 (35.1)    | 497 (48.7) | \( < 0.001 \) | 304 (42.1) | 57 (78.6) | 449 (40.3) | \( < 0.001 \) | 0.453 | \( < 0.001 \) |
| Place of living home            | 553 (62.5)    | 714 (70.0) | 0.001         | 547 (75.8) | 52 (74.3) | 668 (60.0) | 0.784 | \( < 0.001 \) | 0.018 |
| Home support                    | 565 (63.8)    | 709 (69.5) | 0.009         | 320 (44.3) | 55 (78.6) | 899 (80.8) | \( < 0.001 \) | \( < 0.001 \) | 0.651 |
| Polypharmacy³                   | 600 (67.8)    | 806 (79.0) | \( < 0.001 \) | 435 (60.2) | 55 (78.6) | 916 (82.3) | 0.003 | \( < 0.001 \) | 0.430 |
| Temporal disorientation⁴        | 196 (22.1)    | 294 (28.8) | 0.001         | 0            | 3 (4.3)  | 487 (43.8) | \( < 0.001 \) | \( < 0.001 \) | \( < 0.001 \) |
| Use of walking aid              | 353 (39.9)    | 592 (58.0) | \( < 0.001 \) | 0            | 12 (17.1) | 933 (83.8) | \( < 0.001 \) | \( < 0.001 \) | \( < 0.001 \) |
| Reason for ED visit, n (%)      |               |          |               |               |               |               |               |               |               |
| Organ failure⁵                  | 508 (57.4)    | 544 (53.3) | 0.075         | 411 (56.9) | 47 (67.1) | 594 (53.4) | 0.098 | 0.135 | 0.025 |
| Mobility disorders⁶             | 136 (15.4)    | 166 (16.3) | 0.589         | 104 (14.4) | 13 (18.6) | 185 (16.6) | 0.348 | 0.203 | 0.672 |
| Neuropsychiatric disorders⁷     | 98 (11.1)     | 114 (11.2) | 0.943         | 69 (9.6)  | 1 (1.4)  | 142 (12.8) | 0.022 | 0.036 | 0.005 |
| Cancer                          | 24 (2.7)      | 36 (3.5)  | 0.308         | 24 (3.3)  | 3 (4.3)  | 33 (3.0)   | 0.672 | 0.665 | 0.533 |
| Social issue⁸                   | 119 (13.4)    | 160 (15.7) | 0.168         | 114 (15.8) | 6 (8.6)  | 159 (14.3) | 0.108 | 0.377 | 0.181 |
| Length of stay, mean ± SD       |               |          |               |               |               |               |               |               |               |
| ED (hour)                       | 21.5 ± 13.8   | 23.5 ± 13.7 | \( < 0.001 \) | 19.9 ± 13.3 | 23.7 ± 15.7 | 24.3 ± 13.7 | 0.078 | \( < 0.001 \) | 1.000 |
| Hospital (days)                 | 12.4 ± 15.4   | 15.8 ± 20.0 | 0.004         | 10.8 ± 13.5 | 11.5 ± 18.7 | 16.3 ± 19.9 | 1.000 | \( < 0.001 \) | 0.334 |
| Hospital admission, n (%)       | 415 (46.9)    | 548 (53.7) | 0.003         | 310 (42.9) | 39 (55.7) | 614 (55.2) | 0.040 | \( < 0.001 \) | 0.929 |

PRISMA-7: “Program of Research on the Integration of Services for the Maintenance of Autonomy”; ER²: Emergency room evaluation and recommendation; ED: Emergency department; SD: Standard deviation; ³Score < 3; ⁴Score ≥ 3; ⁵Comparison based on unpaired t-test; ⁶Comparison based on an analysis of variance (ANOVA) with Bonferroni correction for multiple comparisons; ⁷Number of medications daily taken ≥ 5; ⁸Inability to give the current year and/or months; ⁹Defined as an acute organ decompensation; ¹⁰Defined as gait and/or balance impairment and/or fall with or without fall-related injuries; ¹¹Defined as delirium, dementia, behavioral disorders; ¹²Defined as the absence of symptoms of acute disease combined with an acute increase of the use of formal and/or informal home and social services leading to an inability to cope at home; P-values significant (i.e., < 0.05) in bold.
walking aid ($P \leq 0.001$). They had also a longer length of stay in ED ($P = 0.002$) and in hospital ($P = 0.004$), and were more frequently admitted to hospital ($P = 0.003$) compared to those with a PRISMA-7 low-risk level. Similar results for age, sex, home support, polypharmacy, temporal disorientation and use of walking aid were observed for comparisons between the three ER2 risk levels. More serious presenting conditions and higher incidence of adverse events were found in high-risk levels compared to moderate-risk and low-risk levels ($P \leq 0.003$). There was no significant different for place of living ($P = 0.784$) between mild and moderate-risk levels, as well as no significant difference for sex ($P = 0.453$) between mild and high-risk levels, and no significant difference for home support ($P = 0.651$) and polypharmacy ($P = 0.430$) between moderate and high-risk levels. Furthermore, neuropsychiatric disorders as a reason for ED visit was more prevalent in patients with low-risk levels compared to those in moderate-risk group ($P = 0.022$), but less prevalent compared to high-risk group ($P = 0.036$). Prevalence of this reason of ED visit was also higher in high-risk group of patients compared to those in moderate-risk group ($P = 0.005$). Organ failure was also a more prevalent reason of ED visit in patients with a moderate compared to a high-risk level ($P = 0.025$). In addition, longer stay in ED and hospital as well as hospital admission were observed in patients considered to have high-risk level compared to those with low-risk level ($P \leq 0.001$). Hospital admission was also more prevalent in patients with the moderate-risk level compared to those with the low-risk level ($P = 0.040$).

Regardless of the regression model used, the PRISMA-7 high-risk level was associated with a prolonged length of ED ($P \leq 0.001$) and hospital stay ($P \leq 0.007$) (Table 2). The PRISMA-7 low-risk level was associated with a short length of stay in ED and in hospital for linear regression model ($P \leq 0.007$). The ER2 high-risk level was associated with a prolonged length of ED and hospital stay, as well as elevated hospital admission rates ($P \leq 0.001$) for all regression models. The ER2 low-risk level was associated with short length of stay in ED and in hospital, as well as a lower hospital admission rate ($P \leq 0.001$).

Kaplan-Meier distributions showed significant differences in the time elapsed before hospital discharge between patients with a PRISMA-7 low and high-risk level; those with a high-risk level were discharged from ED ($P = 0.010$) and from hospital ($P = 0.002$) later compared to those with low-risk (Fig. 1). The three ER2 participants’s risk levels differed significantly for length of stay in ED and in hospital ($P \leq 0.001$), those with a high-risk level being discharged later than those in the low-risk ($P \leq 0.001$) (Fig. 2). Furthermore, patients with a high-

Table 2 Linear and logistic regression models showing the association of the length of stay in emergency department and hospital as well as the hospital admission (dependent variable with separated model for each variable) with PRISMA-7 and ER2 risk (independent variable) adjusted on place of living and reasons for emergency department visit ($n = 1905$)

| Dependent variable | Independent variable | Linear Regression\(a, b\) | Logistic Regression\(a, b, c\) |
|--------------------|----------------------|---------------------------|---------------------------------|
|                    |                      | $\beta$ [95%CI] P-Value   | OR [95%CI] P-Value              |
| Length of stay in emergency Department in hours | PRISMA-7 risk | High 2.28 [1.64;3.51] $\leq 0.001$ 1.57 [1.29;1.91] $\leq 0.001$ |  |
|                    | Low –2.28 [−3.51;−1.04] $\leq 0.001$ 0.64 [0.52;0.77] $\leq 0.001$  |
|                    | ER2 risk Low –4.11 [−5.38;−2.83] $\leq 0.001$ 0.58 [0.47;0.72] $\leq 0.001$  |
|                    | Moderate 1.09 [−2.19;4.38] 0.514 0.97 [0.58;1.62] 0.903  |
|                    | High 3.84 [2.58;5.09] $\leq 0.001$ 1.69 [1.38;2.07] $\leq 0.001$  |
| Length of stay in Hospital in days | PRISMA-7 score | Abnormal 3.21 [0.89;5.54] 0.007 1.48 [1.12;1.96] 0.006 |  |
|                    | Normal –3.21 [−5.54;−0.89] 0.007 0.68 [0.51;0.90] 0.006  |
|                    | ER2 risk level Mild –5.17 [−7.63;−2.72] $\leq 0.001$ 0.49 [0.36;0.68] $\leq 0.001$  |
|                    | Moderate –2.31 [−8.14;3.52] 0.438 0.62 [0.29;1.33] 0.218  |
|                    | High 5.31 [2.92;7.70] $\leq 0.001$ 2.10 [1.55;2.85] $\leq 0.001$  |
| Hospital admission | PRISMA-7 score | Abnormal – 1.32 [1.01;1.58] 0.003 |  |
|                    | Normal 0.76 [0.63;0.91] 0.003  |
|                    | ER2 risk level Mild – 0.59 [0.48;0.71] $\leq 0.001$  |
|                    | Moderate – 1.17 [0.72;1.91] 0.0516  |
|                    | High – 1.64 [1.36;1.98] $\leq 0.001$  |

PRISMA-7: “Program of Research on the Integration of Services for the Maintenance of Autonomy”; ER2: Emergency Room evaluation and recommendation; $\beta$: Coefficient of regression beta; OR: Odd ratio; HR: Hazard ratio; CI: Confident interval; “Model adjusted on place of living and reasons for emergency department visit; “Separate model for each risk level; “Hazard ratio; “Model adjusted on place of living.”
Fig. 1 Kaplan-Meier estimates of the probability of long emergency department (a) and hospital (b) stay with PRISMA-7 risk level in participants ($n = 1905$). a Emergency department, b Hospital.
Fig. 2 Kaplan-Meier estimates of the probability of long emergency department (a) and hospital (b) stay with ER$^2$ risk levels in participants ($n = 1905$). a Emergency department. b Hospital.
risk level were discharged later when hospitalized than those in the moderate-risk level ($P = 0.026$).

All performance criteria were poor (i.e., sensitivity, specificity, PPV and NPV $\leq 0.74$ LR $\leq 1.40$ and area under receiver operating characteristic curve $\leq 0.62$) with worse performances for PRISMA-7 compared to ER$^2$ (Table 3).

**Discussion**
The findings showed that PRISMA-7 and ER$^2$ risk stratifications were successfully associated with length of stay in ED and in hospital, as well as hospital admission rates. High-risk levels were associated with longer stays and higher rates of hospital admission, whereas the contrary association was found in low-risk levels for both tools. The strength of association was stronger for ER$^2$ compared to PRISMA-7. However, performance criteria for the high-risk level of these ED adverse events were low, with the lowest performance being reported with PRISMA-7.

PRISMA-7 and ER$^2$ high-risk levels were both associated with prolonged ED and hospital stays and higher rates of hospital admission. This association was expected for ER$^2$ because this clinical tool has especially been designed to determine prognosis for these short-term ED adverse events [4, 12–14, 19]. In particular, prolonged ED and hospital stays have previously been reported with the ER$^2$ high-risk levels, similar to the findings in the present study [4, 12, 14, 19]. In contrast, to the best of our knowledge, it is the first time that an association with prolonged ED and hospital stays is found with the PRISMA-7 high-risk level. PRISMA-7’s initial objective was to screen disability in community dwelling older adults [16, 17]. This tool has also been used to identify frailty in older ED users, but not related adverse events [21]. Frailty may be defined as a state of vulnerability related to homeostasis following a stressor and is independently associated with adverse events [22]. Frailty results from an accumulation of severe comorbid conditions and physiological aging [22, 23]. A tool which assesses frailty and its severity indirectly evaluates prognosis for frailty-related adverse events. The main characteristic of older ED users compared to younger users is an accumulation of severe and chronic morbidities and related-disabilities [3]. These particular health and functional conditions greatly influence the ED care plan [2, 3, 8]. Although older adults undergo more diagnostic tests and procedures than younger ED patients, their ED diagnoses tend to be less accurate [2, 7]. This has been attributed to atypical disease presentation due to multiple comorbidities, which can complicate the clinical presentation, diagnosis and care plan [1, 3, 5, 7]. Additionally, chronic morbidities may decompensate in a cascade and lead to disabilities and complications, regardless of the nature and the severity of the acute condition leading to the ED visit [3, 22]. This statement explains why the PRISMA-7 could be used as a prognostic tool to predict prolonged ED and hospital stays. Our findings also underscore that PRISMA-7 and ER$^2$ low-risk levels were associated with shorter ED and hospital stays. This finding is consistent with the former because it suggests that both tools also detect older ER users with a low level of frailty. The early identification of this category of older ED users may be useful for the ED care continuum, because these ED users with low levels of frailty have a higher propensity to be discharged to their place of living compared to those with higher stages of frailty [5–9, 24].

Our study revealed that, even if significant associations of PRISMA-7 and ER$^2$ risk levels with short-term ED adverse events were found, the predictive performance was poor for both tools. No study has assessed the ability of

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**Table 3** Performance criteria of PRISMA-7 and ER$^2$ high-risk for long length of stay in emergency department and hospital, and hospital admission (n = 1905)

| Criteria                        | Abnormal PRISMA-7 score$^a$ | ER$^2$ high-risk level$^b$ |
|--------------------------------|-----------------------------|-----------------------------|
|                                | Length of stay              | Hospital admission          | Length of stay          | Hospital admission |
|                                | Emergency department$^a$    | Hospital$^b$                | Emergency department$^a$| Hospital$^b$        |
| Sensitivity                    | 0.60                        | 0.63                        | 0.57                    | 0.67               |
| Specificity                    | 0.50                        | 0.56                        | 0.50                    | 0.46               |
| Positive predictive value      | 0.38                        | 0.36                        | 0.54                    | 0.38               |
| Negative predictive value      | 0.71                        | 0.72                        | 0.53                    | 0.73               |
| Likelihood ratio of positive test | 1.20                      | 1.18                        | 1.14                    | 1.25               |
| Likelihood ratio of negative test | 1.25                      | 1.26                        | 1.16                    | 1.40               |
| Area under receiver operating characteristic curve | 0.55                        | 0.56                        | 0.53                    | 0.59               |

PRISMA-7: “Program of Research on the Integration of Services for the Maintenance of Autonomy”; ER$^2$: Emergency room evaluation and recommendations; $^a$Score $\geq 3/7$; $^b$Score $\geq 6/14$; $^c$Length of stay used as a discontinuous variable defined as the highest tertile of lengths of stay (i.e., $> 26.9$ h); $^d$Length of stay used as a discontinuous variable defined as the highest tertile of lengths of stay (i.e., $> 13$ days)
PRISMA-7 or ER\(^2\) of predicting short-term ED adverse events, therefore, the comparison of this result with previous studies is difficult. However, our findings demonstrate similar results to other screening tools, such as The Identification of Seniors at Risk (ISAR), the Triage Risk Screening Tool (TRST), the eight-item questionnaire of Runciman, and the seven-item questionnaire of Rowland, if we take into consideration the ability of predicting return visits of older patients after they have been discharged from the ED. [25] Poor predictive performances were also reported in this study.

The imbalance between a significant association of the PRISMA-7 and ER\(^2\) high-risk levels with short-term ED adverse events and the poor performance criteria highlight that a significant statistical association between a high-risk level and an adverse event does not necessarily guarantee that the prognostic tool is accurate nor reliable. Similar results have been reported with prolonged hospital stays in older patients admitted to geriatric wards [4]. Despite the limited predictive performance, improvement in cost-effectiveness of targeted geriatric interventions using a screening tool has been previously demonstrated with low positive predictive value (i.e., between 20 and 30%) [24]. In addition, screening tools are known to be particularly useful in decision-making procedures to prevent events that depend on multiple factors [25, 26]. This result confirms that screening older patients with the highest risk level and providing them a timely and appropriate intervention is an efficient strategy.

The important number of older ED users recruited and the appropriate design to achieve the research objectives are the strengths of our study. However, some limitations need to be underscored. Firstly, the recruitment has been performed in a single centre, which limits the external validation of our results. Secondly, although adjustment for clinical characteristics influencing the studied short-term adverse events was performed, there are still potential residual confounders that may influence the short-term ED adverse events. Thirdly, both PRISMA-7 and ER\(^2\) do not take into consideration the reasons of ED visits which is usually an acute disease directly influencing the occurrence of short-term ED adverse events. For instance, an acute disease may compensate in a cascade related to chronic morbidities cumulated in older ED users, leading to multiple acute organ decompensations and disabilities and exposing this group to prolonged length of stays and hospitalisations [5–9]. However, in the regression analysis used to examine the association of between PRISMA-7 and ER\(^2\) risk levels with the three short-terms ED adverse events, all models were adjusted by the reasons of ED visits and a significant association was found.

**Conclusion**

In conclusion, even if PRISMA-7 and ER\(^2\) high-risk levels were associated with incidental short-term ED adverse events, their performance criteria for these risks were relatively low, suggesting that they cannot be used as prognostic tools for this purpose. However, they may be the starting point of an assessment that could provide pertinent information pertaining to ED tailor-made geriatric interventions.

**Abbreviations**

PRISMA-7: Program of Research on the Integration of Services for the Maintenance of Autonomy; ER\(^2\): Emergency room evaluation and recommendations; EDs: Emergency departments; SD: Standard deviations; ANOVA: Analysis of variance; ISAR: Identification of Seniors at Risk; TRST: Triage Risk Screening Tool

**Acknowledgments**

The authors are grateful for their cooperation: 1) the participants; 2) nurses of the Department of emergency of Jewish General hospital, and in particular Mrs. Valerie Shneidman and Mr. Jonathan Harroche; 3) Pharmacists of Jewish General hospital, and in particular Mrs. Julie Roy and Mrs. Nikki kampouris; 4) Mrs. Betty Elkaïm (Vice-President and Chief Development Officer of the Foundation of the Jewish General Hospital); 5) the Department of information and technology of Jewish General Hospital, and in particular Mrs. Isabelle Aumont, Mrs. Maria Veres, Mrs. Christine Bougie and Mrs. France Guimond; and 6) Mrs. Abbigail Shaw and Mr. Davis Schiller.

**Authors’ contributions**

CPL and OB: Principal investigators, study conception and design, obtaining funding and drafting first version of the manuscript and approval of the final manuscript; MA: Co-investigator, study conception and design, drafting first version of the manuscript and approval of the final manuscript; KG, EM and CV: recruitment of participants, study management, revising the manuscript critically for important intellectual content and approval of the final manuscript; JL: revising the manuscript critically for important intellectual content and approval of the final manuscript. All authors read and gave their approval of the final manuscript and for publication.

**Funding**

The study was financially supported by private donations (Mrs. Abbigail Shaw and Mr. Davis Schiller), the RUISSS McGill Centre of Excellence on Longevity and the Foundation of the Jewish General Hospital (Montreal, Quebec, Canada). The funding sources have no involvement in study design; in the collection, analysis and interpretation of data; in the writing of the manuscript; and in the decision to submit the article for publication.

**Availability of data and materials**

Data and materials are available on request. Request should be sent to the corresponding author: Cyrille Launay, MD, PhD; Department of Medicine, Division of Geriatric Medicine, Sir Mortimer B. Davis – Jewish General Hospital, McGill University, 3755 chemin de la Côte-Sainte-Catherine, Montréal, QC H3T 1E2, Canada; E-mail: . All requests needs a cover letter explaining the objective, justification and the referent ethic committee.

**Ethics approval and consent to participate**

This study was conducted in accordance with the ethical standards set forth in the Helsinki Declaration (1983). The study was classified as a clinical quality-improvement program for older ER users’ care plan by the Ethics Committee and the West-Central Montreal Health Review Office for quality programs of the Jewish General Hospital (Montreal, Quebec, Canada). All participants recruited in this study provided a verbal informed consent because the study did not change the usual clinical practice. The verbal informed consent was obtained from the patients themselves in the presence of their trusted person, usually a family member, who helped them to make decision. Verbal informed consent were obtained for all participants following a systematic and standardized process used in the ED ward where the study has been performed. Participants or their legal guardian when appropriate, were informed that their medical information may be used for research purpose.
they disagreed, they informed the physician taking care of them and a note was recorded in their chart. The Research Review Office of the Jewish General Hospital and the Research Ethic Committee of the Integrated Health and Social Services University Network for West-Central Montreal approved this process.

Consent for publication
All authors read and gave their approval of the final manuscript and for publication.

Competing interests
The authors CPL, JL, KG, CV, EM, MA and OB report no disclosures relevant to the manuscript.

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Received: 11 June 2020 Accepted: 29 December 2020

Published online: 22 January 2021

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