Synthesis of evidence on the use of ecological momentary assessments to monitor health outcomes after traumatic injury: rapid systematic review

Rebecca J. Mitchell*, Rory Goggins and Reidar P. Lystad

Abstract

Background: With the increasing use of mobile technology, ecological momentary assessments (EMAs) may enable routine monitoring of patient health outcomes and patient experiences of care by health agencies. This rapid review aims to synthesise the evidence on the use of EMAs to monitor health outcomes after traumatic unintentional injury.

Method: A rapid systematic review of nine databases (MEDLINE, Web of Science, Embase, CINAHL, Academic Search Premier, PsychINFO, Psychology and Behavioural Sciences Collection, Scopus, SportDiscus) for English-language articles from January 2010–September 2021 was conducted. Abstracts and full-text were screened by two reviewers and each article critically appraised. Key information was extracted by population characteristics, age and sample size, follow-up time period(s), type of EMA tools, physical health or pain outcome(s), psychological health outcome(s), general health or social outcome(s), and facilitators or barriers of EMA methods. Narrative synthesis was undertaken to identify key EMA facilitator and barrier themes.

Results: There were 29 articles using data from 25 unique studies. Almost all (84.0%) were prospective cohort studies and 11 (44.0%) were EMA feasibility trials with an injured cohort. Traumatic and acquired brain injuries and concussion (64.0%) were the most common injuries examined. The most common EMA type was interval (40.0%). There were 10 key facilitator themes (e.g. feasibility, ecological validity, compliance) and 10 key barrier themes (e.g. complex technology, response consistency, ability to capture a participant’s full experience, compliance decline) identified in studies using EMA to examine health outcomes post-injury.

Conclusions: This review highlighted the usefulness of EMA to capture ecologically valid participant responses of their experiences post-injury. EMAs have the potential to assist in routine follow-up of the health outcomes of patients post-injury and their use should be further explored.

Keywords: Ecological momentary assessment, Experience sampling, Injury, Health outcome
populations in-hospital and post-discharge [2–4]. With the increasing use of mobile technology, one technique that may enable routine monitoring of patient health outcomes and patient experiences of care is ecological momentary assessments (EMAs).

EMAs, experiencing sampling methods or ambulatory assessments allow snap-shots into real-life moments by enabling self-collection of thoughts, behaviours, symptoms, activities, experiences, or biometric data (e.g. heart rate), in real-time from a defined population [5, 6]. Information is usually collected for short, specific periods either after a specific event or experience (i.e. event-contingent sampling), at fixed, regular intervals throughout a day (i.e. interval-contingent sampling) or at random time points during a day (i.e. signal-contingent sampling) [7]. These repeated measurements are being collected increasingly via mobile devices, including smart phones or sensor equipment.

The information collected using EMAs is usually recorded in a natural environmental situation and allows temporal sequences of symptoms or conditions to be monitored and relationships, and often interdependencies, between conditions to be explored [5, 8, 9]. The use of EMAs can improve data validity by reducing many data collection biases, such as retrospective recall bias, and data entry or transcription errors [5, 8]. EMAs also allows for the timely acquisition of information regarding an individual’s health outcomes and, when needed, swiftly acted upon [9].

EMAs have been frequently used to monitor risk factors and behaviours for intentional injuries, including suicidal thoughts, behaviours, and acts of self-harm [10], but their use has been fairly limited in monitoring health outcomes after other types of injuries. Around 973 million individuals sustain a traumatic injury (such as fractures, dislocations, open wounds, sprains or strains) worldwide each year that required some form of healthcare [11]. A serious injury can have an adverse impact on the individual, their family, and local community [12].

Many seriously injured individuals can experience ongoing mobility and functional limitations, depression, anxiety and post-traumatic stress disorder (PTSD) [13]. Therefore, the ability to monitor physical, psychological, and social health outcomes after injury, along with experiences of service use, and social participation would be advantageous to identify the need for interventions, ongoing service needs, and service planning, including use of primary care and allied health. The aim of this rapid review is to synthesise the evidence on the use of EMAs to monitor health outcomes after traumatic unintentional injury.

Method
This rapid review synthesises the evidence on the use of EMAs to monitor different types of health outcomes after sustaining an injury. The review examines information on the type of EMA tools used, follow-up periods, the different tools and methods used to monitor health outcomes, and the facilitators and barriers identified to using EMAs to monitor health outcomes post-injury. This review adhered to the Preferred Reporting Items for Systematic Review and Meta-analyses (PRISMA) statement [14].

Definitions
Research studies were included in the rapid review if they used EMAs to monitor the health outcomes of an individual after sustaining a traumatic injury. An EMA is a method by which information is captured at multiple time periods regarding an individuals’ current health (e.g. physical or psychological) state or experiences in real-time [5]. A traumatic injury was considered to be “a bodily lesion at the organic level caused by acute exposure to physical agents such as mechanical energy, heat, electricity, chemicals, and ionizing radiation interacting with the body in amounts or at rates that exceed the threshold of human tolerance. In some cases, injuries result from the sudden lack of essential agents such as oxygen or heat” [15]. There were no restrictions on the type of injury sustained. However, intentional injuries following self-harm or interpersonal violence were excluded from the review as EMAs have been frequently used to monitor risk factors and behaviours for intentional injuries, whereas use of EMAs has been limited to monitor outcomes after unintentional injuries.

Individual health outcomes could either capture health states in the short-, medium- or long-term. Health outcomes could include information on physical health or pain outcomes (e.g. physical functioning, mobility, activities of daily living (ADLs), pain, medication use), psychological health (e.g. depression, anxiety, stress, PTSD), or general health and social outcomes (such as quality of life (QoL), social activity participation, biometric data e.g. heart rate, hours spent sleeping, step count).

Data sources and eligibility criteria
A systematic search was conducted using nine databases: MEDLINE, Web of Science, Embase, CINAHL, Academic Search Premier, Psyclnfo, Psychology and Behavioural Sciences Collection, Scopus, SportDiscus. The search strategy was developed with a university librarian and included the following search terms: (ecological momentary assessment* OR Ecological Momentary Assessment OR momentary assessment* OR EMA OR experience sampling OR ambulatory assessment* OR
event sampling), AND (injur* OR accident OR trauma* OR accident* OR wound* OR lesion* OR bruise* OR abrasion* OR harm) (see Additional file 1 for full search strategy).

Studies were excluded if the article was a systematic or other type of review, a single case report, a study protocol, or if there was insufficient detail regarding the health outcome(s) examined. Results were limited to English-language articles that were published in peer-review journals from 1 January 2010 to 21 September 2021. Snowballing of reference lists from the articles was conducted to identify any potential articles not previously identified.

Abstract screening
The title, abstract and citation information relating to each study identified during the database searches was imported to Endnote X20 and duplicates removed. The abstracts were independently assessed for inclusion by three reviewers (RG, RL, RM), who met regularly to discuss any uncertainties. If the abstract did not report on how EMAs were used to monitor health outcomes after unintentional injury it was excluded. Any disagreements on abstract inclusion were discussed and consensus achieved. Abstract screening was independently verified for accuracy by dual screening each abstract in pairs, with 99.0% percent agreement achieved during the initial abstract screen (i.e. RG/RL and RG/RM). After discussion, consensus was obtained to include n = 34 abstracts to the full-text screening stage.

Full-text screening, data extraction and quality review
In the full-text screening each study was assessed by three reviewers (RG, RL, RM). Any study that did not meet the inclusion criteria was excluded. For studies that met the inclusion criteria, key characteristics of each study were extracted during the full-text review by one reviewer (RG), including: authors and publication year; review objective/aim; study type; country; and data collection timeframe, population characteristics, age and sample size, EMA type (i.e. signal, event or interval) and follow-up time period(s), EMA assessment tool(s), physical health or pain outcome(s), psychological health outcome(s), general health or social outcome(s), and facilitators or barriers of EMA methods identified during the study by the authors. Data extraction results were independently verified for accuracy by a second reviewer (RM) and any disagreements were discussed. The methodological quality of the articles was assessed by one reviewer (RG) using the CASP Cohort checklist [16] or the CASP RCT checklist [17]. Any clarifications regarding methodological quality were discussed between all reviewers.

Narrative synthesis
The information on the included studies in the data extraction table was compared and a narrative synthesis was undertaken of the facilitators and barriers by one reviewer (RM) and these were appraised by a second reviewer (RL). The narrative synthesis involved reading and reviewing each of the facilitators and barriers identified in the discussion section of each article. Then an inductive, iterative process was used to categorise each factor identified as either as a facilitator or a barrier based on the key factor theme (e.g. a facilitator of ‘data collection minimises recall bias’ was categorised as ‘reliability’ and a barrier of ‘EMA is potentially a time burden for study participants’ was categorised as ‘time-burden’). Each factor was allocated to one theme, but several facilitators or barriers could be identified in each article.

Results
There were 4418 records identified during the database searches. After removing duplicates, 2425 records remained. After abstract screening, 35 full-text records were assessed for eligibility, along with four records identified after snowballing. A final 29 articles using data from 25 unique studies were included in the rapid review (Fig. 1).

Almost all (84.0%) unique studies were prospective cohort studies, with four (16.0%) randomised control trial (RCTs) designs. Eleven (44.0%) studies were feasibility trials of the effectiveness of using EMA to record information with a traumatically injured cohort. The majority of studies were conducted in the United States (84.0%), with one (4.0%) study each in Canada, Germany, the Netherlands, and Spain. Traumatic brain injury (TBI) (i.e. n = 5 mild TBI, n = 2 severe TBI, and n = 2 TBI – severity not specified), concussion (n = 5), and acquired brain injury (n = 2) were the most common injuries examined (64.0%). Traumatic injury, such as fractures open wounds or anterior cruciate ligament injuries (20.0%; n = 5) and spinal cord injury (16.0%; n = 4) were also examined. The most common EMA type was interval (40.0%; n = 10), with nine (36.0%) random, and three (12.0%) event types. One (4.0%) study used both interval and event EMAs, and one study (4.0%) used all three EMA types. A variety of assessment measures were used to record information on physical or pain, psychological, general health and social outcomes post-injury using EMA. Only nine (36.0%) studies recorded information on social activity participation. Almost all (89.7%) authors identified at least one facilitator or barrier of using EMA (Table 1).

There were ten key facilitator themes and ten key barrier themes identified in studies using EMA to examine health outcomes post-injury (Table 2). Where facilitators
were identified, feasibility, ecological validity, and compliance were the most common themes identified for each EMA type (Fig. 2a). Complex technology, response consistency, and the ability to capture a participant’s full experience were the most common barrier themes identified for interval EMAs. Compliance decline and the potential for participants to be negatively affected by prompt frequency were the most common barrier themes identified for random EMAs (Fig. 2b). Quality assessment measures for articles varied and few studies (24.0%; n = 6) received all ‘Yes’ ratings (Tables 3 and 4).

**Discussion**

This rapid review identified 25 unique studies where EMAs were used to monitor symptoms and QoL after unintentional traumatic injury. Obtaining this sort of experiential information post-injury can assist in identifying temporal changes in clinically-relevant symptoms and health states, in making decisions regarding further treatment options, in allocating health service and resource requirements, and has the ability to identify any unmet health needs [48].

With mobile technology advances, the use of EMA is likely to become more commonplace to conduct follow-up studies for injured individuals in real-time and in real-world settings. This review has identified that each type of EMA (i.e. interval, random, and event) has demonstrated feasibility to monitor post-injury recovery, commonly for individuals who sustained brain or spinal cord injuries. Individuals who sustain either brain injuries or a spinal cord injury can have a long-term recovery and adjustment period, as individuals and their families adjust to living with the consequences of these injuries and their associated symptoms and related health issues [49–51]. It is possible that EMA could facilitate the long-term monitoring of the recovery of traumatically injured populations.

In general, compliance with data collection in prospective studies that have used EMAs has been reported as high [52], and that EMAs can reduce recall bias through using real-time data collection [23, 34, 47]. While participant responses using EMAs are considered to be more reliable than retrospective studies [22, 31, 48], several study authors did identify as a potential barrier the reliability of information from participants obtained using EMAs [20, 23, 27, 37, 42, 46, 47]. For some participant information, such as health service use (e.g. primary care, emergency department visit or hospital admission), there would
Table 1: Characteristics of studies using ecological momentary assessments to monitor health outcomes after injury

| Authors and publication year | Objective/aim | Study type | Country (data collection dates) | Injured population, age and sample size | EMA type & follow-up time period(s) | EMA assessment tool(s) | Physical/health or pain outcome(s) | Psychological health outcome(s) | General health or social outcome(s) | Facilitators or barriers of EMA methods identified by study authors |
|-----------------------------|---------------|------------|---------------------------------|----------------------------------------|-------------------------------------|------------------------|----------------------------------|-------------------------------|---------------------------------|---------------------------------------------------------------|
| Albania et al. 2021 [18]    | Test whether distress intolerance predicted traumatic brain injury (mTBI) or anxiety sensitivity. | Prospective cohort study | Florida, United States (unspecified date) | n = 70 aged ≥ 18 years with mild traumatic brain injury (mTBI) or anxiety sensitivity. | Interval. 1. 3h after film; 2. 10am, 3pm, & 8pm each day for 13 days after film. Event. 1. Study follow-up visits at time 3, 4 & 5. | 1. Text messages with survey link to Qualtrics online survey. | N/A | 1. Naturalistic trauma distress intolerance. 2. Depression Anxiety & Stress Scale (DASS-21). | N/A | Facilitators: 1. Feasibility of web-based ecological momentary assessment (EMA) demonstrated. 2. Study results support the use of EMA procedures to track an intervention. 3. A modification to EMA facilitated the study team to contact and encourage participants. Barriers N/A |
| Bethhaus et al. 2021 [19]   | Assessed the feasibility of design elements of a yoga-based interventional trial for post-concussive headaches among Veterans with mTBI, as well as the acceptability of the intervention. | Randomised controlled acceptability and feasibility trial | United States (2017–2019) | n = 27 to 54 Veterans aged ≥18 years with mTBI at various intervention stages. | Event. 1. Study follow-up visits at time 3, 4 & 5. | Other text messages sent to participants. 2. Brief Pain Inventory (HIT-6) impact of headaches on activities of daily living (ADL). 3. K Scale: Survey of Headache Impact (KSSI-H). 4. Short-Form McGill Pain Questionnaire (SF-MPQ). 5. Medication taken to treat headaches. | 1. Neuro-behavioural Symptom Inventory (NSI). 2. Perceived Stress Scale (PSS). 3. Barriers & facilitators to activity participation. 4. Home yoga practice. | 1. Patient-Reported Outcomes Measurement Information System (PROMIS). 2. Short Form Health Survey (SF-36). 3. Barriers & facilitators to activity participation. 4. Home yoga practice. | 1. Patients able to assess and monitor their own symptoms. 2. Patients able to assess and monitor their own symptoms. 3. Ability for results to be shared with the treating clinician. 4. Sustains ecologically valid results. Barriers N/A |
| Forster et al. 2020 [20]    | To establish the feasibility of using EMA (i.e., internet-based patient report of compliance) in patients with an acquired brain injury (ABI). To map fluctuations in patients’ responses. To determine whether patients’ compliance and their fluctuations in response behaviour were related to their level of functioning. | Prospective cohort pilot study | Germany (unspecified date) | n = 15 individuals with an ABI with cognitive and motor impairments. Aged 18–70 years | Random: 8 prompts per day for 7 days. Prompts were given between 8 am and 8 pm with at least an hour between prompts. | 1. The software moviXEOS, App version 1.3.0 via Android smartphones (Motorola Moto G, third generation) | 1. Rasch-Based Depression Screening using a Five-Point Likert Scale. 2. Attention Network Test to assess attention and reaction time compute-based reaction time. 3. The Verbal Learning and Memory Test to measure declarative verbal memory, learning performance, long-term encoding, and recognition performance. 4. Ability to conduct 10 basic daily functions using the Barthel Index. | N/A | 1. Compliance decreased with every passing testing day. 2. The higher frequency of EMA prompts correlated with lower compliance as burden increased. 3. Difficult to gauge how reliable and realistic the patients answers were. 4. Participant mood and emotion may have been influenced by high frequency of prompts. 5. The use of EMA might also be limited by other psychological factors such as social desirability and patients’ individual differences. Barriers N/A |
| Authors and publication year | Objective/aim | Study type | Country (data collection dates) | Injured population, age and sample size | EMA type & follow-up time period(s) | EMA assessment tool(s) | Physical health or pain outcome(s) | Psychological health outcome(s) | General health or social outcome(s) | Facilitators or barriers of EMA methods identified by study authors |
|-----------------------------|--------------|------------|---------------------------------|----------------------------------------|-------------------------------------|-----------------------|------------------------------------|-----------------------------|----------------------------------|--------------------------------------------------------------------------------|
| Hart et al. 2019 [21]       | To describe and provide the rationale for a randomized controlled trial for depression or anxiety after moderate to severe TBI, which will test two treatments based on behavioral activation (BA). | Randomised controlled trial | United States (unspecifed date) | n = 60 individuals with TBI that was sustained at least 6 months prior to enrollment. Consciousness must be altered and depression and anxiety present. Participants aged ≥ 18 years. | Random. 5 times per day within a 14-h window for 8 weeks. | 1. Collected via the LifeData System via RealLife Exp (a mobile app). | N/A | 1. Emotional status/behaviour - Global Severity Index of the Brief Symptom Inventory (BSI)-18, Environmental Reward Observation Scale (EROS), the Behavioral Activation for Depression Scale (BADS). | 1. Participant Assessment with Recomposed Tools-Objective (PART-O) to assess societal/community participation. 2. Satisfaction with life Scale (SWLS). 3. Quality of life after brain injury (QOLIBRI). 4. Patient Global Impression of Change. 5. Cognitive/functional status using Wechsler Abbreviated Scale of Intelligence. 6. Functional status - Extended Glasgow Outcome Scale. | Facilitators 1. The incorporation of commonly used technology in the form of smartphone apps has the potential to enhance the delivery of treatment in this population. Barriers N/A |
| Authors and publication year | Objective/aim | Study type | Country (data collection dates) | Injured population, age and sample size | EMA type & follow-up time period(s) | EMA assessment tool(s) | Physical health or pain outcome(s) | Psychological health outcome(s) | General health or social outcome(s) | Facilitators or barriers of EMA methods identified by study authors |
|------------------------------|---------------|------------|---------------------------------|----------------------------------------|--------------------------------------|---------------------|----------------------------------|----------------------------------|----------------------------------|------------------------------------------------------------------|
| Juengst et al. 2015a [22]    | A) Pilot feasibility and validity of a mobile health (mHealth) system for tracking mood-related symptoms after TBI. B) To investigate within-person variability in daily self-reported emotional and fatigue symptoms and factors associated with high within-person variability among individuals with chronic TBI. | A) Prospective cohort pilot study. B) Secondary analysis of prospective cohort pilot study | United States (unspecified date) | A) n = 20 individuals with TBI and aged ≥ 18 years. B) n = 18 adults with chronic TBI (2–27 years post-injury) who owned and could independently use an Apple or Android device. Participants aged 22–60 years. | Interval. Daily assessments for 8 weeks. | 1. Conducted via mHealth system delivered through iPerform application. | N/A | 1. Depressive symptoms using the Patient Health Questionnaire 2 (PHQ2). 2. Generalised Anxiety Disorders (GAD2). 3. Positive and Negative Affect Scale (PANAS). 4. Patient Health Questionnaire 9 (PHQ9). 5. Generalised Anxiety Disorders 7 (GAD7). | 1. Impact of fatigue on daily life using a 7-point Likert scale. | A) Facilitators 1. Participants showed good compliance and high satisfaction with the method of EMA. 2. Ecological valid results were demonstrated. 3. The simple interface on the app proved to be helpful with participants with TBI and could successfully complete assessments. 4. A schedule feature resulted in greater compliance and satisfaction. Barriers 1. Some participants frequently completed the wrong assessment. 2. Some participants completed too many assessments per day. B) Facilitators 1. Feasibility of mobile phone-based EMA demonstrated. 2. Use of EMA may reduce misidentification of individuals with clinically significant symptoms. 3. Higher frequency repeated symptom assessment in a natural environment over a short period could provide a more valid measure of emotional symptoms and a better indicator of clinically meaningful change at the individual level. Barriers 1. Single daily assessments may not be a valid representation of symptoms’ progression. 2. Method of self-reporting may contribute to variability. |
| Authors and publication year | Objective/aim | Study type | Country (data collection dates) | Injured population, age and sample size | EMA type & follow-up time period(s) | EMA assessment tool(s) | Physical health or pain outcome(s) | Psychological health outcome(s) | General health or social outcome(s) | Facilitators or barriers of EMA methods identified by study authors |
|------------------------------|--------------|------------|---------------------------------|---------------------------------------|-----------------------------------|-----------------------|-------------------------------|---------------------------------|-----------------------------------|------------------------------------------------------------------|
| Lenaert et al. 2019 [24]     | 1. To investigate the feasibility of using the experiencing sampling method (ESM) in individuals with ABI. 2. To explore the usability of ESM data on a clinical level by illustrating the interactions between person, environment, and affect. | Prospective cohort study | The Netherlands (July 2014–March 2015) | n = 17 individuals with ABI aged 18–65 years | Random. 10 semi-random beeps throughout the day between 7:30 am and 10:30 pm for 6 days. | 1. PsyMate (smart eHealth GmbH, Luxembourg) - a small electronic device with a touch-screen interface. | 1. Physical well-being using a 7-point Likert scale. | 1. PANAS. 2. Mood and self-esteem using a 7-point Likert scale. | | Facilitators 1. Data collection method had little interference on daily activities despite high volume of beeps. 2. User-friendly interface allowed for easy completion of assessments. 3. Familiarity with the questions over time reduced time and energy needed to answer them. 4. Data collection method had little influence on mood and feelings promoting ecological validity. 5. Users were able to monitor own progress and rehabilitation. Barriers N/A |
| Pavliscak et al. 2016 [25]   | 1. To examine engagement with a mobile application (i.e. mCare) for wounded Veterans rehabilitating in their communities. 2. To examine associations between Veterans’ background characteristics and their engagement with mCare. | Prospective Randomised Controlled Trial | United States (Unspecified specified) | n = 95 participants who received mCare. This included individuals with TBI (no distinction was made between those with TBI and those with other health issues). Participants were aged ≥ 18 years. | Interval. Daily assessment (specified time) for 36 weeks. | 1. EMA delivered via the mCare mobile application. 2. Text messages to notify of new information on the mCare application. | N/A | N/A | 1. General participation. | Facilitator 1. Findings suggest mCare can and will be adopted by Veterans in a community setting, even those with cognitive and emotional difficulties. Barriers 1. Participants’ exposure to mCare declined systematically as Veterans out processed from rehabilitation. |
| Rabinowitz et al. 2020 [26]  | To illustrate a novel framework for conceptualizing, collecting, and analyzing concussion symptom data. | Prospective cohort pilot study | United States (Unspecified date). | n = 10 recently concussed adolescents and young adults aged 15–33 years. | Interval. Five assessments per day for 20 days (morning, early afternoon, late afternoon, evening, and night). | 1. Collected via the LifeData System (via RealLife Exp (a mobile app)). | N/A | N/A | 1. 22-item Post-concussion Symptom Scale (PCSS). |
| Authors and publication year | Objective/aim | Study type | Country (data collection dates) | Injured population, age and sample size | EMA type & follow-up time period(s) | EMA assessment tool(s) | Physical health or pain outcome(s) | Psychological health outcome(s) | General health or social outcome(s) | Facilitators or barriers of EMA methods identified by study authors |
|-------------------------------|--------------|------------|-------------------------------|----------------------------------------|------------------------------------|----------------------|----------------------------------|-----------------------------|---------------------------------|----------------------------------------------------------------------------------------------------------------------------------|
| Rabinowitz et al. 2021 [27]  | To describe where, with whom, and how time was spent daily, and to characterise positive and negative affect, boredom, enjoyment, and perceived accomplishment as a function of time, activity, location, and social context, in people with chronic moderate-severe TBI and depression/anxiety. | Prospective cohort study | United States (5 April 2018-1 Feb 2020) | n = 23 individuals with TBI and at least mild depression or anxiety. Mean age 47.7 years. | Random. Notified 5 times in 14-h window for ~ 2 weeks. | 1. Collected via the LifeData System (via RealLife Exp (a mobile app)). | N/A | 1. Positive and Negative Affect using 20-item PANAS. | 1. General well-being and activity using multi-choice. Free-text responses were able to be provided for each question to qualify responses. Activities in last hour recorded, including who primarily doing activity with. | Facilitators: N/A. Barriers: 1. Closer monitoring with more proactive troubleshooting could have improved participants' response rates. |
| Smith et al. 2012 [28]  | To assess the utility of mHealth technologies, including personal digital assistant-based EMA and two-way interactive text (SMS) messaging, for providing treatment feedback to clinicians, encouraging and motivating Veterans throughout treatment, and monitoring participants for relapse after treatment discharge. | Prospective cohort pilot study | United States (Unspecified date) | n = 27 male veterans suffering from PTSD and/or mTBI. Distinctions between those with PTSD from those with mTBI were not made. Age unspecified. | Internal, Random and Event. Active phase: 1. Daily assessments. Was initially randomised then at routine intervals. Follow-up phase 3 months post-discharge 1. Daily at either 9am or 5pm. | 1. Electronic survey tool designed to support data collection via PDAs was used in active phase. 2. Assessments sent and completed via SMS (provided by LifeWIRE corporation) in follow-up phase | 1. Level of pain using a 1–10 Likert scale. 2. Symptoms Checklist-6. | 1. Depressive symptoms using BriefCOPE, and Beck Depression Inventory-II. | 1. Types of hassles and uplifts. 2. General activities. | Facilitators: 1. After data collection was changed from random to routine intervals, response rate improved. Barriers: 1. EMA at random times during waking hours was disruptive to participants' schedules and led to low response rate. 2. Participants were resistant to carrying two electronic devices and routinely left their assigned PDA in the housing unit during the day, which contributed to the low EMA response rate. 3. The PDAs were viewed as being clunky and out of date compared with smart phones. 4. Participants indicated that they did not believe the data they were providing were doing any good because they could not see any effect on their treatment. On a few occasions, participants temporarily ceased participating in EMA data collection after researchers and clinicians failed to respond to a stress or crisis event recorded in EMA data. |
| Authors and publication year | Objective/aim | Study type | Country (data collection dates) | Injured population, age and sample size | EMA type & follow-up time period(s) | Physical health or pain outcome(s) | Psychological health outcome(s) | General health or social outcome(s) | Facilitators or barriers of EMA methods identified by study authors |
|-----------------------------|--------------|------------|---------------------------------|----------------------------------------|----------------------------------|-----------------------------------|----------------------------------|----------------------------------|----------------------------------|
| Suffoletto et al. 2013 [29] | To examine whether patients with mTBI receiving text message-based education and behavioural support had fewer and less severe post-concussive symptoms than those not receiving text messaging support, and determine the feasibility of using text messaging to assess daily symptoms and provide support to patients with mTBI. | RCT pilot | United States, July–September 2012 | \(n = 43\) adults aged \(\geq 18\) years with mTBI \(n = 18\) in intervention group and \(n = 25\) in control group | Interval. Three daily questions at 9 am, 1 pm and 5 pm over 14 days | SMS messages. 1. Pain scale via have you had a headache in the last 24 h rating using 4-item Likert scale. | 1. Difficulty concentrating in last 24 h rated using 4-item Likert scale. 2. Irritability or anxiety in last 24 h rated using 4-item Likert scale. | N/A | Facilitators 1. Provision of self-care support messages focused on symptom-specific education, reassurance, and management guidance lowered irritability or anxiety during the acute recovery period. Barriers N/A |
| Sufrinko et al. 2019 [30] | To evaluate mobile EMA as an approach to measure sport-related concussion symptoms, explore the relationships between clinical outcomes and mobile EMA, and determine whether mobile EMA was advantageous for predicting recovery outcomes compared to traditional symptom report. | Prospective cohort study | United States (September 2016 – December 2016) | \(n = 20\) athletes aged 12–19 years with SRC | Random. Pseudo-randomised in 3 time blocks. i.e. Mon-Fri 7-8 am, 3-4 pm and 9-10 pm and Sat-Sun 9-10 am, 3-4 pm and 9-10 pm daily for 4 weeks and clinical assessment at visit 1 and visit 2. | 1. EMA surveys were administered via mobile EMA mobile application (Ilumivu, Inc) 2. The application utilized push notifications prompting the participant to open the application. | 1. Vestibular/Ocular Motor Screening rated using 0–10 EMA scale assessing intensity for 4 symptoms (i.e. headache, dizziness, nausea, and fogging). 2. Neurocognitive using the Immediate Post-Concussion Assessment and Cognitive Testing (ImPACT). | 1. 22-item Post-concussion Symptom Scale (PCSS) rated using 7 point Likert scale. | Facilitators 1. Mobile EMA better predicted recovery time than Post-Concussion Symptom Scale (PCSS). 2. Mobile EMA data across recovery better predicted recovery duration compared with PCSS score at any clinic visit, but illustrated symptom patterns that may further inform clinical profiles and guide treatment recommendations. Barriers 1. Participants were less likely to respond as days since injury increased 2. Results reflected a diminishing response rate throughout the course of the study. May be due to a. Mobile EMA data collected for a longer duration than previous studies. b. Intervals were less frequent than other studies, but yielded higher compliance with more intervals. c. Participants symptoms resolving and disinterest in completing mobile EMA. |
Table 1 (continued)

| Authors and publication year | Objective/aim | Study type | Country (data collection dates) | Injured population, age and sample size | EMA type & follow-up time period(s) | EMA assessment tool(s) | Physical health or pain outcome(s) | Psychological health outcome(s) | General health or social outcome(s) | Facilitators or barriers of EMA methods identified by study authors |
|-----------------------------|--------------|------------|-------------------------------|----------------------------------------|------------------------------------|------------------------|-----------------------------------|---------------------------------|----------------------------------|---------------------------------------------------------------|
| Trbovic et al. 2021 [31]    | To use actigraphy and mobile EMA to examine the relationship between sleep parameters and next day symptoms. | Prospective cohort study, pilot study | United States (September 2016 – December 2016) | n = 17 athletes aged 12–19 years with recent concussion | Interval: three scheduled assessments per day within 1h of prompt (i.e. morning – 7 am school days and 9 am weekends, afternoon – 3 pm, evening – 9 pm). By fourth week around half the participants had recovered, and sample size was underpowered beyond 3 weeks post-injury. Random: several random prompts daily for approximately 2 weeks after their initial office visit. | 1. EMA surveys were administered via mobile EMA mobile application (Lumivu, Inc). 2. The application utilized push notifications prompting the participant to open the application. 3. Participants were an Actigraph GT3x+ on their nondominant wrist for 24h per day. | 1. Somatic symptoms using PCSS and 7-point Likert scale. 2. Sleep and activity measures with real-time reports of cognitive and physical activity, and compare objective measures with real-time reported symptoms among youth during recovery after concussion | N/A | 1. Ecological validity was shown. | Facilitators 1. Youth able to use the SuperBetter app in conjunction with traditional medical care for post-concussive symptoms and were satisfied with use of the app. 2. Participants who used the app to complement medical care had more relief from concussion symptoms than those who had traditional medical care alone. 3. The gameful and/or social interactive design of SuperBetter was effective to improve optimism. 4. Patients able to assess and monitor their own symptoms.  | Barriers N/A |
| Wiebe et al. 2016 [32]      | To determine the feasibility of EMA following youth concussion, gather real-time reports of cognitive and physical activity, and compare objective measures with real-time reported symptoms among youth during recovery after concussion. | Prospective cohort pilot study | United States. (Unspecified date) | n = 34 recently concussed adolescents aged 11–19 years | Random: several random prompts daily for approximately 2 weeks after their initial office visit. | 1. EMA surveys were administered via mobile EMA mobile application (Lumivu, Inc). 2. The application utilized push notifications prompting the participant to open the application. 3. Participants were an Actigraph GT3x+ on their nondominant wrist for 24h per day. | N/A | 1. Daily cognitive rest and exertion were measured as number of text messages sent in minutes of screen time and gaming and minutes of reading or school work. | N/A | 1. Ecological validity was shown. | Barriers N/A |
| Worthen-Chaudhari et al. 2017 [33] | To evaluate whether a mobile health application that employs elements of social game design could complement medical care for unresolved concussion symptoms. | Prospective cohort study | United States (Phase I – 13 Aug 2014 – 7 Jan 2015; Phase II – 7 Jan 2015 – 4 Nov 2015) | Phase I n = 20, Phase II n = 19 adolescents aged 13–18 years, with concussion symptoms ≥ 3 weeks post-injury | Event: Participants asked to log activity at the frequency of one logged activity per day for 5 days each week, for a target of 15 logged activities over the first 3 weeks between pre-and post-test. | 1. EMA conducted via a mobile application – SuperBetter. | N/A | 1. Optimum, measured by the Centers for Disease Control’s Breakthrough to Better Concussion Management Final Report (BTB-CM). | 1. Optimism, measured by the Life Orientation Test-Revised (LOT-R). | Facilitators 1. Youth were able to use the SuperBetter app in conjunction with traditional medical care for post-concussive symptoms and were satisfied with use of the app. 2. Participants who used the app to complement medical care had more relief from concussion symptoms than those who had traditional medical care alone. 3. The gameful and/or social interactive design of SuperBetter was effective to improve optimism. 4. Patients able to assess and monitor their own symptoms. | Barriers N/A |
| Authors and publication year | Objective/aim | Study type | Country (data collection dates) | Injured population, age and sample size | EMA type & follow-up time period(s) | EMA assessment tool(s) | Physical health or pain outcome(s) | Psychological health outcome(s) | General health or social outcome(s) | Facilitators or barriers of EMA methods identified by study authors |
|------------------------------|---------------|------------|--------------------------------|----------------------------------------|-----------------------------------|-----------------------|----------------------------------|-------------------------------|-----------------------------|---------------------------------------------------------------|
| Carlozzi et al. 2018 [34] | Investigated the most efficient means of measuring pain intensity and pain interference comparing EMA to end-of-day (EOD) data, with the highest level of measurement reliability as examined in individuals with spinal cord injury (SCI). | Prospective observational study | United States (unspecified date) | n = 131 individuals with SCI aged ≥ 18 years. Participants also required to endorse ≥ 4 out of 10 average pain. | Interval 1: Five times throughout the day (upon waking, 11 am, 3 pm, 7 pm, and bedtime) | 1. Wrist-worn accelerometer enhanced with a user interface for entry of self-report data (i.e. the PRO-Diary; CamNTech, Cambridge, UK). | 1. Pain 5-point Likert scale. 2. PROMIS pain intensity. 3. SCI-QOL pain interference. 4. 10-item short form pain interference 5. Pain interference 5-point Likert scale. | N/A | N/A | 1. EMA is easy to complete as responses are less reliant on memory. 2. The timing of EMA assessments does not impact reliability. 3. The presence of a floor effect for EMA pain interference presented an analytical challenge in our data. 4. Calibration data may over- or under-estimate the pain ratings of participants in this study. 5. Pain in SCI is multifaceted, and thus ratings of pain intensity do not capture the full breadth of the pain experience in individuals with SCI. |
| Authors and publication year | Objective/aim | Study type | Country (data collection dates) | Injured population, age and sample size | EMA type & follow-up time period(s) | EMA assessment tool(s) | Physical health or pain outcome(s) | Psychological health outcome(s) | General health or social outcome(s) | Facilitators or barriers of EMA methods identified by study authors |
|-----------------------------|----------------|------------|---------------------------------|----------------------------------------|-------------------------------------|----------------------|-----------------------------------|---------------------------------|----------------------------------|--------------------------------------------------------------------------------|
| Carlozzi et al. 2021 [35]   | Examined the effect of sleep quality on same day Health Related Quality of Life (HRQoL). | Prospective cohort study. | United States (unspecified date) | \( n = 170 \) individuals with SCI aged \( \geq 18 \) years | Interval 1: Three times a day (i.e. morning, afternoon, evening) for 7 days. | 1. Smart phone or paper diary. 2. E4 wristband (Empatica) recorded heart rate variability, electrodermal activity, body movement (accelerometer data), & skin temperature. | 1. Pain 10-point Likert scale. 2. PROMIS pain intensity. 3. PROMIS pain interference. | 1. Thinking 10-point Likert scale. 2. PROMIS cognitive function abilities. 3. PROMIS depression. 4. PROMIS anxiety. | 1. Fatigue 10-point Likert scale. 2. PROMIS sleep disturbances. 3. PROMIS sleep-related impairment. 4. PROMIS fatigue. 5. PROMIS ability to participate in social roles and activities. | Facilitators: 1. Does not take very long for participants to habituate to a monitoring device (i.e. E4). 2. Ability to examine temporal relationships among symptoms. 3. Data collection minimises recall bias. 4. Maximises ecological validity of responses. Barriers: 1. The E4 wristband does not have established algorithms for evaluating sleep in the general population or in individuals with SCI. 2. Self-reported sleep quality and objective sleep measured by E4 was not consistent – potentially due to lack of established sleep algorithms for E4 and/or challenges for a wrist-worn device in capturing information on sleep for a SCI population with limited mobility and neurophysiological challenges. 3. EMA is a relatively intense data collection procedure that can be burdensome on the participant. 4. Potential for individuals to change behaviour when they are being monitored. |
| Authors and publication year | Objective/aim | Study type | Country (data collection dates) | Injured population, age and sample size | EMA type & follow-up time period(s) | EMA assessment tool(s) | Physical health or pain outcome(s) | Psychological health outcome(s) | General health or social outcome(s) | Facilitators or barriers of EMA methods identified by study authors |
|-----------------------------|--------------|------------|---------------------------------|----------------------------------------|----------------------------------|----------------------|-----------------------------|-----------------------------|---------------------------------|--------------------------------------------------|
| Kratz et al. 2017a [36]     | A) To examine whether pain acceptance moderates the momentary associations of pain intensity with pain interference and physical activity in people with chronic pain and SCI. B) To examine study compliance, protocol acceptability, and reactivity of intensive data collection methods in adults with chronic pain and SCI. C) To examine the moderating effect of within- and between-person pain acceptance on associations between pain and physical and psychological functioning. | A) Prospective observational cohort study. B) Secondary analysis of prospective observational cohort study. C) Secondary analysis of prospective observational cohort study. | A) United States (June 2014–January 2016) | n = 128 individuals with chronic pain and SCI aged ≥ 18 years. | Interval and event: 1. Five times throughout the day for a week (upon waking, 11 am, 3 pm, 7 pm, and bedtime). | 1. A wrist-worn accelerometer called the (PRO-Diary). 2. EOD electronic diaries (online collection site). | 1. Average physical activity per minute. 2. Brief Pain Inventory on 10-point scale. 3. Chronic Pain Coping Inventory-42 (CPI). 4. Chronic Pain Acceptance 8 (CPAQ8). 5. Pain Catastrophizing using the Catastrophizing subscale from the Coping Strategies Questionnaire. 6. Spinal Cord Injury – Quality of Life (SCI-QOL). | 1. Depressive symptoms using the PHQ-9. | | A) Facilitators 1. Dynamically demonstrates ecological validity. 2. Combination of objective and subjective measures reduced problems related to overlapping method variance. B) Barriers 1. User error or internet connection difficulties interfered with completion of assessments. 2. It was more difficult for subjects to enter ratings during busy wake and bed-time routines, which, for people with SCI, often involve lengthy and assisted self-care routines (e.g. bowel and bladder care). |
| Authors and publication year | Objective/aim | Study type | Country (data collection dates) | Injured population, age and sample size | EMA type & follow-up time period(s) | EMA assessment tool(s) | Physical health or pain outcome(s) | Psychological health outcome(s) | General health or social outcome(s) | Facilitators or barriers of EMA methods identified by study authors |
|-----------------------------|---------------|------------|---------------------------------|----------------------------------------|----------------------------------|---------------------|----------------------------------|----------------------------------|-------------------------------|------------------------------------------------------------------|
| Todd et al. 2018a [39]      | A) To utilize EMA to measure intra-individual diurnal variations in neuropathic pain and effect on exercise and non-exercise days. B) To describe strategies necessary to adapt EMA to measure neuropathic pain in adults with SCI, and explore participant perceptions of using EMA to measure pain sensations. | Prospective cohort study | Canada (Unspecified date) | n = 6 physically active men with SCI greater than 1 year post-injury with low neuropathic pain. Participants aged 27–50 years | Random. A total of 6 prompts per day, over 6 days. | 1. EMA surveys were administered via mobile EMA mobile application (Ilumivu, Inc). 2. The application utilised push notifications prompting the participant to open the application. 3. Fitbit Surge wrist-worn heart rate (HR) monitors were worn by participants to collect HR data. | 1. Modified Neuropathic Pain Scale rated using 10-item Likert scale. | 1. 11-point, single item Feeling Scale (FS) measured affect. 2. Felt Arousal Scale (FAS) measured arousal. | 1. Heart rate captured using the Fitbit surge HR monitor. | A) Facilitators 1. Little missing data and no participant dropout. 2. The measurement schedule was not cumbersome and therefore minimized the probability of reduced data quality and quantity. B) Barriers 1. Pain may have been exacerbated due to repeatedly asking participants to think about their pain levels. |
| Todd et al. 2019b [40]     | A) Prospective cohort study. B) Secondary analysis of prospective cohort study. | Secondary analysis of prospective cohort study | (Unspecified date) | n = 6 physically active men with SCI greater than 1 year post-injury with low neuropathic pain. Participants aged 27–50 years | Random. A total of 6 prompts per day, over 6 days. | 1. EMA surveys were administered via mobile EMA mobile application (Ilumivu, Inc). 2. The application utilised push notifications prompting the participant to open the application. 3. Fitbit Surge wrist-worn heart rate (HR) monitors were worn by participants to collect HR data. | 1. Modified Neuropathic Pain Scale rated using 10-item Likert scale. | 1. 11-point, single item Feeling Scale (FS) measured affect. 2. Felt Arousal Scale (FAS) measured arousal. | 1. Heart rate captured using the Fitbit surge HR monitor. | A) Facilitators 1. Participants provided positive responses regarding the practicability and usefulness of EMA to accurately capture their neuropathic pain experience. 2. Ecological validity was demonstrated. 3. Participants appreciated minimal morning/night-time interference. 4. Random EMA allowed for the dynamic phenomenon of neuropathic pain to be captured, while minimizing daily interference to participants. 5. The 6-day protocol could partially explain participants’ support for six assessments per day as the protocol length may have reduced the overall burden experienced by participants. 6. Easy user interface allowed for participant’s positive observations related to the “quick” nature of assessments. 7. Use of notifications to prompt participants was viewed as useful. B) Barriers 1. Participants reported that receiving multiple EMA prompts negatively influenced their neuropathic pain perception. |
| Authors and publication year | Objective/aim | Study type | Country (data collection dates) | Injured population, age and sample size | EMA type & follow-up time period(s) | EMA assessment tool(s) | Physical health or pain outcome(s) | Psychological health outcome(s) | General health or social outcome(s) | Facilitators or barriers of EMA methods identified by study authors |
|------------------------------|--------------|------------|--------------------------------|----------------------------------------|-----------------------------------|----------------------|-------------------------------|---------------------------------|---------------------------------|---------------------------------------------------------------------------------|
| Traumatic injury, including head injury | | | | | | | | | | |
| Gonzalez-Borato et al. 2021 [41] | To evaluate Psixport’s ability to gather real-time information about injured athlete’s psychological responses during the rehabilitation; to test the users’ perceived usability of Psixport; and to compare the reliability and differences between real-time data gathered with Psixport and the data gathered through the one-time retrospective method. | Prospective cohort feasibility study | Spain (unspecified date) | n = 28 severely injured athletes that require surgery and have a rehabilitation prescription therapist after surgery. Participants aged ≥ 18 years. | Event. Daily after completing rehab session for 15 days. | 1. Two questions from the Universal Pain Assessment Tool were included in Wong-Baker’s Faces Pain Rating Scale format. | 1. Psychological Responses to Sports Injury Inventory (PRSII) assessed cognitive appraisals regarding injuries on 6 dimensions: devastation, disinhibited, re-organisation, feeling cheated, restlessness, and isolation. | 1. An adaptation of the picture-oriented Self-Assessment Manikin to assess emotional valence and arousal. 2. Behavioural Responses captured using Sports Injury Rehabilitation Adherence Scale (SIRAS). | 1. Users found interface app simple and easy. 2. Relatively short questionnaire length allowed for commitment by participants. 3. Ecological validity and validity of responses maximised. 4. Users were able to monitor own progress and rehabilitation. | 1. Compliance reduced with each passing day. |
| Pacella et al. 2018 [42] | To examine changes in post-concussive symptoms (PCS) over the acute post-injury recovery period, focusing on how daily PCSs differ between mTBI and other injury types. | Prospective cohort study | United States (April 2013 – March 2014) | n = 108 adults with traumatic injury aged ≥ 18 years. | Interval. 3 times per day (9 am, 1 pm, 5 pm) for 14 days. Different symptoms (i.e. somatic, cognitive, emotional) were assessed at each interval. | 1. Assessments sent and completed via text message. | 1. Somatic measures (e.g. headaches) using a 5-point scale of headache intensity. | N/A | N/A | 1. Simple technology and using text-message allowed for easy completion for participants. 2. There may be reporting and recall biases associated with the EMA pattern. |
| Pacella et al. 2018 [43] | To apply ESM via daily text messaging to monitor and detect relationships among psychosocial factors and post-injury pain across the first 14 days after emergency department (ED) discharge | Prospective observational cohort study | United States (January 2016 – May 2017) | n = 75 adults with a trauma related injury aged 18–60 years | Interval. Five assessments at 5 pm for 14 days. | 1. Five assessments sent and completed via text message. | 1. Pain rated on a 1–10 Likert scale. | 1. Social support rated on a 1–7 Likert scale. | 1. Hyperarousal rated on a 1–7 Likert scale. 2. Intrusions rated on a 1–7 Likert scale. 3. Avoidance rated on a 1–7 Likert scale. | 1. ED patients were receptive to the manner of which assessments were conducted. 2. High compliance rate despite relatively high volume of text messages. 3. Validity was limited as assessment was completed once per day. |
| Authors and publication year | Objective/aim | Study type | Country (data collection dates) | Injured population, age and sample size | EMA type & follow-up time period(s) | EMA assessment tool(s) | Physical health or pain outcome(s) | Psychological health outcome(s) | General health or social outcome(s) | Facilitators or barriers of EMA methods identified by study authors |
|-----------------------------|---------------|------------|---------------------------------|----------------------------------------|-------------------------------------|-----------------------|-----------------------------------|----------------------------------|----------------------------------|----------------------------------|
| Price et al. 2014 [44]      | To determine the proportion of trauma patients that would consent to receiving daily text messages assessing mental health, determine response rates to daily text messages among trauma patients, identify predictors of higher rates of responding, assess patient satisfaction, and determine provider burden. | Prospective cohort pilot study | United States (Unspecified date) | n = 29 individuals with a traumatic injury. Mean age of 37.1 years. | Interval. A daily assessment for 15 days. One and three month follow-up. | 1. Assessments sent and completed via text message. | 1. Pain rated using a 1–10 Likert scale | 1. Hypervigilance rating using a 1–7 Likert scale. 2. Avoidance rated using a 1–7 Likert scale. 3. Re-experiencing rated using a 1–7 Likert scale. | 1. Social support rated using a 1–7 Likert scale. | Facilitators 1. Text messages are an efficient method of implementing a "watchful waiting" program after a traumatic event. Barriers 1. Technical difficulties reported as the primary reason for non-response. |
### Table 1 (continued)

| Authors and publication year | Objective/aim | Study type | Country (data collection dates) | Injured population, age and sample size | EMA type & follow-up time period(s) | EMA assessment tool(s) | Physical health or pain outcome(s) | Psychological health outcome(s) | General health or social outcome(s) | Facilitators or barriers of EMA methods identified by study authors |
|------------------------------|---------------|------------|---------------------------------|----------------------------------------|------------------------------------|-----------------------|-----------------------------------|---------------------------------|---------------------------------|------------------------------------------------------------------|
| Price et al. 2017 [45]       | To evaluate the use of a mobile phone application to collect symptom data during the acute post-trauma period. | Prospective cohort study | United States (Unspecified date) | n = 23 individuals with traumatic injury. Mean age of 27.6 years. | Random. Daily assessments between 7am-8pm for 30 days. One and three month follow-up. | 1. Conducted via mobile application, Metricwire (Waterloo, ON). 2. Follow-up interviews were conducted via telephone. | 1. Pain rated using a 1–10 Likert scale 2. Arousal rated using a 1–7 Likert scale. 3. Avoidance rated using a 1–7 Likert scale. 4. Re-experiencing rated using a 1–7 Likert scale. 5. Free-text response to what concerned them most that day. | 1. Arousal rated using a 1–7 Likert scale. 2. Avoidance rated using a 1–7 Likert scale. 3. Re-experiencing rated using a 1–7 Likert scale. 4. Free-text response to what concerned them most that day. | 1. Sleep rated using a 1–7 Likert scale. | Facilitators 1. The use of mobile devices to monitor symptoms presents a low-burden and low-cost method with substantial reach to learn about recovery. 2. Most participants felt that 1 survey per day was appropriate. 3. Participants stated that the notifications to complete tasks were helpful. 4. Ease of use of the interface, familiarity with the mobile device, and brevity of the survey allowed for easy completion of survey. 5. Using an individual’s personal device significantly reduced the cost associated with conducting studies or delivering intervention. Barriers 1. Participants may have preferred an opportunity to respond more frequently but did not feel an overwhelming obligation to do so. 2. Participants found the inclusion of the same questions in each survey repetitive, which may have diminished their willingness to respond to subsequent assessments. 3. Participants requested personalized questions and personalized feedback for a better experience. 4. Responding to assessments via a mobile device may be burdensome to the patient and might result in noncompliance. |
### Table 1 (continued)

| Authors and publication year | Objective/aim | Study type | Country (data collection dates) | Injured population, age and sample size | EMA type & follow-up time period(s) | EMA assessment tool(s) | Physical health or pain outcome(s) | Psychological health outcome(s) | General health or social outcome(s) | Facilitators or barriers of EMA methods identified by study authors |
|------------------------------|---------------|------------|--------------------------------|----------------------------------------|-------------------------------------|----------------------|-----------------------------------|-----------------------------------|-----------------------------------|------------------------------------------------------------------|
| Price et al. 2018 [46]       | To evaluate the acceptability of administering PTSD symptom assessments via a mobile application throughout the acute post-trauma period | Prospective cohort study | United States (Unspecified date) | n = 90 individuals with traumatic injury (n = 1 participant was injured following a physical assault). Mean age of 35.1 years. | Random. Daily assessments between 7am-8pm for 30 days. One and three month follow-up. | 1. Conducted via mobile application, Metricwire (Waterloo, ON). | 1. Pain rated on a 1–10 Likert scale. | 1. Post-Traumatic Stress Disorder (PTSD) checklist-5 (PCL-5) to assess PTSD symptoms (8-items). 2. Free-text response to what concerned them most that day. | 1. Sleep assessment using PCL-5. | Facilitators: 1. Assessments were perceived as moderately helpful and minimally burdensome. 2. Demonstrates possibility to obtain free response and Likert scale response via EMA. Barriers: 1. A subset of participants may not complete daily assessments due to technical difficulties or a lack of interest. 2. Variability in response time. Many took advantage of the 14-h response window. |

N/A Not applicable
| Theme                      | Examples                                                                 | Authors and EMA type                                                                 |
|----------------------------|--------------------------------------------------------------------------|--------------------------------------------------------------------------------------|
| **Facilitators**           |                                                                          |                                                                                      |
| Accepted technology        | Ease of use of the interface, familiarity with the mobile device, and brevity of the survey allowed for easy completion of survey. Does not take very long for participants to habituate to a monitoring device (i.e. E4). | Price et al. [45] - random Carlozzi et al. [47] - interval                           |
| Clinician-monitor          | Ability for results to be shared with the treating clinician.            | Forster et al. [20] - random                                                         |
| Compliance                 | Relatively short questionnaire length allowed for commitment by participants. A schedule feature resulted in great compliance and satisfaction. | Gonzalez-Borato et al. [41] - event Juengst et al. [22] - interval                     |
| Ecological validity        | Higher frequency repeated symptom assessment in a natural environment over a short period could provide a more valid measure of emotional symptoms and a better indicator of clinically meaningful change at the individual level. Data collection method had little influence on mood and feelings promoting ecological validity. | Juengst et al. [23] - interval Lenaert et al. [24] - random                           |
| Feasibility                | Simple technology and using text-message allowed for easy completion for participants. Text messages are an efficient method of implementing a "watchful waiting" program after a traumatic event. | Parcella et al. [42] - interval Price et al. [44] - interval                        |
| Methodological quality     | Combination of objective and subjective measures reduced problems related to overlapping method variance. Mobile ecological momentary assessments (mEMA) better predicted recovery time than Post-Concussion Symptom Scale (PCSS). | Kratz et al. [36] – interval and event Sufrinko et al. [30] - random                   |
| Personal device use reduced costs | Using an individual’s personal device significantly reduced the cost associated with conducting studies or delivering intervention. | Price et al. [45] - random                                                           |
| Reliability                | Use of EMA may reduce misidentification of individuals with clinically significant symptoms. Data collection minimises recall bias. | Juengst et al. [23] - interval Carlozzi et al. [47] - interval                       |
| Self-monitor               | Patients able to assess and monitor their own symptoms. Users were able to monitor own progress and rehabilitation. | Forster et al. [20] - random Gonzalez-Borato et al. [41] - event                     |
| Temporal relationships     | Ability to examine temporal relationships among symptoms. mEMA data across recovery better predicted recovery duration compared with PCSS score at any clinic visit, but illustrated symptom patterns that may further inform clinical profiles and guide treatment recommendations. | Carlozzi et al. [47] - interval Sufrinko et al. [30] - random                         |
| **Barriers**               |                                                                          |                                                                                      |
| Capture full experience    | Pain in spinal cord injury (SCI) is multifaceted, and thus ratings of pain intensity do not capture the full breadth of the pain experience in individuals with SCI. Single daily assessments may not be a valid representation of symptoms/ progression. | Carlozzi et al. [34] - interval Juengst et al. [23] - interval                        |
| Complex technology         | EMA requires more sophisticated analytical hardware for monitoring and data capture. User error or internet connection difficulties interfered with completion of assessments. | Carlozzi et al. [34] - interval Kratz et al. [37] – interval and event                 |
| Compliance decline         | Compliance decreased with every passing testing day. Participants were less likely to respond as days since injury increased. | Forster et al. [20] - random Sufrinko et al. [30] - random                           |
| Response consistency       | Difficult to gauge how reliable and realistic the patients answers were. Method of self-reporting may contribute to variability. | Forster et al. [20] - random Juengst et al. [23] - interval                           |
| Floor/ceiling effects      | The presence of a floor effect for EMA pain interference presented an analytical challenge in our data. | Carlozzi et al. [34] - interval                                                     |
| Potential biases           | Potential for individuals to change behaviour when they are being monitored. The use of EMA might also be limited by other psychological factors such as social desirability and patients’ individual differences. | Carlozzi et al. [47] - interval Forster et al. [20] - random                         |
be the potential to cross-check information provided by the participant with information from other sources, such as through record linkage of self-reported data collected using EMA with administrative health records [53].

Five studies [30, 33, 38, 40, 47] identified that EMAs facilitated the examination of temporal relationships, such as health symptoms like pain or PTSD and time since injury event. Examining temporal relationships can assist in providing information on factors associated with participant well-being and clinical improvement or where further treatment options could be considered. In addition, capturing information on the use of primary care and allied health services over time for injured individuals could also provide insight into the frequency of use of these services, along with details of treatment or rehabilitative activities undertaken.

For the post-injury cohorts in this review, only three studies collected biometric data using EMA, with heart rate [39, 40] and sleep and movement activity [31] recorded. Biometric data collection measures can be felt by some participants to be intrusive [54], but may become more frequently used over time. Motion-sensor apps have been incorporated into smart phones and it is possible that the ability to unobtrusively capture some participant activities, like walking, balance, and physical activity participation will grow over time [55].

---

**Table 2** (continued)

| Theme                                      | Examples                                                                 | Authors and EMA type |
|--------------------------------------------|--------------------------------------------------------------------------|----------------------|
| Personalised feedback                      | Participants requested personalized questions and personalized feedback for a better experience. Participants indicated that they did not believe the data they were providing were doing any good because they could not see any effect on their treatment. On a few occasions participants temporarily ceased participating in EMA data collection after researchers and clinicians failed to respond to a stress or crisis event recorded in EMA data. | Price et al. [45] - random Smith et al. [28] – interval, event, and random |
| Participant negatively affected by prompt frequency | Pain may have been exacerbated due to repeatedly asking participants to think about their pain levels. Participant mood and emotion may have been influenced by high frequency of prompts. | Todd et al. [39] - random Forster et al. [20] - random |
| Poor technology                           | The PDAs were viewed as being clunky and out of date compared with smart phones. | Smith et al. [28] interval, event, and random |
| Time-burden                               | EMA is a relatively intense data collection procedure that can be burdensome on the participant. It was more difficult for subjects to enter ratings during busy wake and bed-time routines, which, for people with SCI, often involve lengthy and assisted self-care routines (e.g. bowel and bladder care). | Carlozzi et al. [47] - interval Kratz et al. [37] – interval and event |

---

**Fig. 2** Key facilitator (A) and barrier (B) themes identified for ecological momentary assessment (EMA) by EMA type.

1Multiple themes could be identified for each article.
| Authors                      | Q1 | Q2 | Q3 | Q4 | Q5a | Q5b | Q6a | Q6b | Q7                  | Q8 | Q9 | Q10 | Q11 | Q12 |
|-----------------------------|----|----|----|----|-----|-----|-----|-----|---------------------|----|----|-----|-----|-----|
| **Traumatic brain injury, concussion, and acquired brain injury** |    |    |    |    |     |     |     |     |                     |    |    |     |     |     |
| Albanese et al. [18]        | Y  | Y  | Y  | Y  | Y   | C   | Y   |     | Greater distress intolerance predicted a poorer ability to volitionally suppress intrusions during the monitoring period. | Y  | Y  | Y  | Y  |     |
| Forster et al. [20]         | Y  | Y  | Y  | Y  | N   | Y   | Y   |     | Patients’ mean compliance rate for EMA was 71.6%. Across all variables, a mean of 55.1% variability in responses. No correlation between patients’ compliance or mean fluctuation. | Y  | Y  | Y  | Y  |     |
| Juengst et al. [22]         | Y  | Y  | Y  | Y  | Y   | Y   |     |     | Participants correctly completed 73.4% of all scheduled assessments | Y  | Y  | Y  | Y  |     |
| Juengst et al. [23]         | Y  | Y  | Y  | Y  | N   | Y   | Y   |     | Significant temporal within-person variability occurred for all measures. | N  | Y  | Y  | Y  |     |
| Lenaert et al. [24]         | Y  | Y  | Y  | Y  | N   | Y   | Y   |     | Demonstrated feasibility with a 71% response rate and a 99% completion rate. There were no dropouts and method indicated as user-friendly. | N  | Y  | Y  | Y  |     |
| Rabinowitz et al. [26]      | Y  | Y  | Y  | Y  | Y   | Y   |     |     | Network modelling revealed marked heterogeneity across participants in terms of acute concussion symptoms. | Y  | Y  | C  | Y  |     |
| Rabinowitz et al. [27]      | Y  | Y  | Y  | Y  | Y   | N   | Y   |     | EMA response rate was positively correlated with integrity of episodic memory and education. Activities associated with positive or negative affect were able to be characterised. | N  | Y  | Y  | Y  |     |
| Smith et al. [28]           | Y  | Y  | Y  | Y  | N   | N   | Y   |     | mHealth technologies are feasible adjuncts to traditional medical treatment in the Veteran population. | N  | Y  | Y  | Y  |     |
| Sufrinko et al. [30]        | Y  | Y  | Y  | Y  | Y   | N   | Y   |     | Post-concussion symptoms able to be measured using mobile EMA and symptoms captured able to be used to determine associations with recovery. | Y  | Y  | Y  | Y  |     |
| Trbovich et al. [31]        | Y  | Y  | Y  | Y  | Y   | C   | Y   |     | Sleep efficiency and total sleep time were negatively associated with next day concussion symptoms. | Y  | Y  | Y  | Y  |     |
| Wiebe et al. [32]           | Y  | Y  | Y  | Y  | N   | N   | Y   |     | EMA feasible. Concussion symptoms decreased as the 2-week follow-up period progressed. | Y  | Y  | C  | Y  |     |
| Worthen-Chaudhari et al. [33]| Y  | Y  | Y  | Y  | N   | Y   | Y   |     | Mobile apps using social gaming may promote health management in teens with unresolved concussion symptoms. | Y  | Y  | Y  | C  |     |
| **Spinal cord injury**       |    |    |    |    |     |     |     |     | Identified minimum of number of End of Day (EOD) and Ecological Momentary Assessments (EMAs) needed to achieve different levels of reliability (“adequate”>0.70, “good”>0.80 and excellent >0.90). | N  | Y  | Y  | Y  |     |
| Carlozzi et al. [34]        | Y  | Y  | Y  | Y  | Y   | Y   |     |     | Fluctuations in sleep quality were significantly associated with ratings of Health-related Quality of Life (HRQOL). | Y  | Y  | Y  | Y  |     |
| Carlozzi et al. [35]        | Y  | Y  | Y  | Y  | Y   | Y   |     |     | Pain acceptance significantly moderated the momentary association between pain intensity and pain interference; those with higher pain acceptance experienced a blunted increase in interference when pain was high. | N  | Y  | Y  | Y  |     |
| Kratz et al. [36]           | Y  | Y  | Y  | Y  | Y   | Y   |     |     | Participant compliance was related to time of day/ presence of audible prompts, mobility aid use, race, and baseline levels of pain and pain interference, with more missing data at wake and bedtimes/ no prompts, and for those who used hand-held mobility devices, identified as African American, and/or reported higher baseline pain and pain interference. | Y  | Y  | Y  | Y  |     |
| Kratz et al. [37]           | Y  | Y  | Y  | Y  | Y   | Y   |     |     | Bivariate correlations indicated moderate to large between-person linear associations between pain acceptance, intensity, and catastrophizing. | N  | Y  | Y  | Y  |     |
| Authors                  | Q1 | Q2 | Q3 | Q4 | Q5a | Q5b | Q6a | Q6b | Q7 | Q8 | Q9 | Q10 | Q11 | Q12 |
|-------------------------|----|----|----|----|-----|-----|-----|-----|----|----|----|-----|-----|-----|
| Todd et al. [39]        | Y  | Y  | Y  | Y  | Y   | Y   | Y   | Y   |    | Y  | Y  | Y   | Y   |     |
|                         |    |    |    |    |     |     |     |     | Participants experienced a significant decrease in neuropathic pain following completion of at least one bout of exercise. |    |    |    |    |    |
| Todd et al. [40]        | Y  | Y  | Y  | Y  | N   | N   | Y   | Y   |    | N  | Y  | Y   | Y   |     |
|                         |    |    |    |    |     |     |     |     | Participants reported that EMA protocol was unobtrusive to their daily routines, and effectively captured their neuropathic pain sensations |    |    |    |    |    |

**Traumatic injury, including head injury**

| Authors                  | Q1 | Q2 | Q3 | Q4 | Q5a | Q5b | Q6a | Q6b | Q7 | Q8 | Q9 | Q10 | Q11 | Q12 |
|-------------------------|----|----|----|----|-----|-----|-----|-----|----|----|----|-----|-----|-----|
| Gonzalez-Borato et al. [41] | Y  | Y  | Y  | Y  | N   | N   | N   | Y   |    | Y  | Y  | Y   | Y   |     |
|                         |    |    |    |    |     |     |     |     | Psixport can gather information about injured athletes' cognitive appraisals, emotional responses, behaviours, and pain perceptions. EMA more accurate than retrospective reports. |    |    |    |    |    |
| Pacella et al. [42]     | Y  | Y  | Y  | Y  | Y   | Y   | Y   | Y   |    |    |    |    |    |    |
|                         |    |    |    |    |     |     |     |     | Greater odds of headache and concentration difficulties on day 1 post-injury among the head injured and mild Traumatic Brain Injury (mTBI) groups vs non-head injured trauma controls. |    |    |    |    |    |
| Pacella et al. [43]     | Y  | Y  | Y  | Y  | Y   | Y   | Y   | Y   |    |    |    |    |    |    |
|                         |    |    |    |    |     |     |     |     | Pain scores decreased over time, and daily fluctuations of hyperarousal were associated with daily fluctuations in reported pain level within each person. |    |    |    |    |    |
| Price et al. [44]       | Y  | Y  | Y  | Y  | Y   | N   | N   | Y   |    | N  | Y  | Y   | Y   |     |
|                         |    |    |    |    |     |     |     |     | Response rates were correlated with PTSD symptoms at baseline but not at other times. |    |    |    |    |    |
| Price et al. [45]       | Y  | Y  | Y  | Y  | Y   | N   | N   | Y   |    | N  | Y  | Y   | Y   |     |
|                         |    |    |    |    |     |     |     |     | Responses rates were uncorrelated with PTSD symptoms or depression symptoms at 1-and 3-month post-injury. |    |    |    |    |    |
| Price et al. [46]       | Y  | Y  | Y  | Y  | N   | N   | N   | Y   |    | N  | Y  | Y   | Y   |     |
|                         |    |    |    |    |     |     |     |     | Response rate was 61.1%. Participants reported that the daily assessments were not bothersome and were moderately helpful. |    |    |    |    |    |

Y Yes, N No C Can't tell, N/A Not applicable

*CASP Cohort Appraisal Checklist questions:
1. Did the study address a clearly focussed issue?
2. Was the cohort recruited in an acceptable way?
3. Was the exposure accurately measured to minimise bias?
4. Was the outcome accurately measured to minimise bias?
5a. Have the authors identified all important confounding factors?
5b. Have they taken into account of the confounding factors in the design and/or analysis?
6a. Was the follow up of the subjects complete enough?
6b. Was the follow up of subjects long enough?
7. What are the result of this study?
8. How precise are the results?
9. Do you believe the results?
10. Can the results be applied to the local population?
11. Do the results of this study fit with other available evidence?
12. Does the study have implications for practice?
Conducting follow-up studies of injured individuals post-discharge is not without challenges, such as participant retention [19] and compliance with data collection protocols [56]. EMA studies have generally shown good retention [35, 42] and participant compliance with data collection prompts [52]. However, this review has identified several potential barriers to EMA use, including the acknowledged behaviour-altering effects associated with social desirability bias and the Hawthorne Effect [20, 47]. Six studies queried whether it was possible that the full experience of an injured individual was being captured using EMAs [23, 30, 34, 37, 43, 46]. By limiting the number of follow-up questions that participants are asked to reduce time-burden, it is possible that key aspects of a participant's post-injury experiences are not being captured. Including a free-text option for participants to record any additional information they would like to provide would allow participants the ability to provide pertinent information of their own choosing and could allow a more complete picture of participant experience to be obtained [57].

This review identified that studies that use EMAs should have the functionality to provide feedback to participants regarding their progress in real-time. In general, health symptoms tend to be reported as more intense and longer lasting using EMAs [5]. In four studies, the authors acknowledged that participants appeared to be negatively affected by prompt frequency [20, 30, 39, 40], demonstrating the need to have in place appropriate risk management practices [59] to reduce any adverse impact on study participants. Risk management practices could include the use of clinical thresholds for participant responses (including for pain scores or psychological health measures) to trigger immediate follow-up of the participant from a health professional. There is the potential that some health outcome measures may need to be adapted for use with EMA to reduce the potential for participant re-experiencing [46] of the injurious event or feelings around the injurious event, such as might occur through the collection of information on PTSD symptoms.

**Strengths and limitations**

The strengths of this rapid review were that it used a comprehensive search strategy involving multiple databases, the review followed the PRISMA guideline, a specialist university librarian assisted with the development of the key search terms, and multiple reviewers were involved in the data extraction phase with high interrater reliability. Any clarifications or disagreements were discussed between reviewers and consensus was obtained. However, there were limitations. Only unintentional injuries were considered, therefore articles that used EMA to examine

### Table 4  Quality assessment criteria of each article using CASP RCT Checklist

| Authors | Q1 | Q2 | Q3 | Q4 | Q5 | Q6 | Q7 | Q8 | Q9 | Q10 | Q11 |
|---------|----|----|----|----|----|----|----|----|----|-----|-----|
| Betthauser et al. [19] | Y | Y | Y | N | Y | Y | Y | Y | Y | C |
| Hart et al. [21] | Y | Y | C | N | Y | Y | Y | N | Y | Y | Y |
| Pavliscak et al. [25] | Y | Y | Y | N | Y | Y | Y | C | Y | Y | Y |
| Suffoletto et al. [29] | Y | Y | Y | Y | N | Y | Y | C | Y | Y | Y |

**CASP RCT Appraisal Checklist questions:**

1. Did the trial address a clearly focussed issue?
2. Was the assignment of patients to treatments randomised?
3. Were all of the patients who entered the trial properly accounted for at its conclusion?
4. Were the patients, health workers and study personnel ‘blind’ to treatment?
5. Were the groups similar at the start of the trial?
6. Aside from the experimental intervention, were the groups treated equally?
7. How large was the treatment effect?
8. How precise was the estimate of the treatment effect?
9. Can the results be applied to the local population?
10. Were all clinically important outcomes considered?
11. Are the benefits worth the harms and costs?
Future directions
There are several further opportunities for research using EMAs. As meanings can differ for participant responses using EMAs in different countries, further exploration and establishment of response norms for different types of injuries would be advantageous [60]. There is also potential for other factors, such as type of setting or presence of peers, to influence participant data recording practices, therefore the generation of normative samples (e.g. using non-injured participants) could assist to tease out the influence of some of these factors [61]. Normative samples may also provide information on exposure and risk factors for injury [59], along with the potential for exposure-time estimates for different activities (e.g. worker hazard exposure, time spent playing or training for different sports) [9, 62, 63] using activity sensor technology. There is also the potential for EMAs to be used to record information regarding the perceptions and experience of family members [64] of an injured individual. More broadly, health systems could incorporate long-term symptom and QoL measures using EMAs as part of routine patient follow-up, with any symptom reports outside a normative range being flagged for clinical follow-up and assessment [23].

Conclusion
This review summaries the literature on the use of EMAs to capture symptoms, health states, behaviours, QoL, and activities post-injury. It highlighted the usefulness of EMA to capture ecologically valid participant responses of their experiences post-injury and has identified common facilitators and barriers regarding the use of EMAs. EMAs have the potential to assist in routine clinical follow-up of health outcomes of patients post-injury and their use should be further explored.

Abbreviations
ABI: Acquired brain injury; ADL: Activity of daily living; ED: Emergency department; EMA: Ecological momentary assessment; EOD: End of day; ESM: Experience sampling method; HRQoL: Health-related quality of life; mTBI: Mild traumatic brain injury; N: not applicable; PCS: Post-concussive symptoms; PTSD: Post-traumatic stress disorder; QoL: Quality of life; RCT: Randomised control trial; SCI: Spinal cord injury; TBI: Traumatic brain injury.

Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s12874-022-01586-w.

Acknowledgements
The authors thank Ms. Jan Van Balen, medical research librarian, for their expertise and guidance regarding the development of the search terminology.

Authors’ contributions
All authors contributed to the study conception and design. RG conducted the database search and all authors conducting abstract and full-text screening. The first draft of the manuscript was written by RM and all authors commented on previous versions of the manuscript. All authors read and approved the final manuscript.

Funding
This research did not receive any specific grant from funding agencies in the public, commercial, or not-for-profit sectors.

Availability of data and materials
All data generated or analysed during this study are included in this published article.

Declarations
Ethics approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

Competing interests
The authors declare they have no competing interests.

Received: 4 November 2021 Accepted: 23 March 2022 Published online: 22 April 2022

References
1. Card AJ. Moving beyond the WHO definition of health: a new perspective for an aging world and the emerging era of value-based care. World Med Health Policy. 2017;9(1):127–37.
2. Staiger R, Schwandt H, Puhan MA, Clavien P. Improving surgical outcomes through benchmarking. J Afri Surg. 2019;106(1):59–64.
3. Preyde M, Brassard K. Evidence-based risk factors for adverse health outcomes in older patients after discharge home and assessment tools: a systematic review. J Evid Based Soc Work. 2011;8(5):445–68.
4. Forster AJ, Muijf HJ, Peterson JF, Gandhi TK, Bates DW. The incidence and severity of adverse events affecting patients after discharge from the hospital. Ann Intern Med. 2003;138(3):161–7.
5. Shiffman S, Stone AA, Hufford MR. Ecological momentary assessment. Annu Rev Clin Psychol. 2008;4:1–32.
6. Arneric SP, Cedarbaum JM, Khoo S, Papapetropoulos S, Hill DL, Ropacki M, et al. Biometric monitoring devices for assessing end points in clinical trials: developing an ecosystem. Nat Rev Drug Discov. 2017;16(10):736.
7. Wheeler L, Reis HT. Self-recording of everyday life events: origins, types, and uses. J Pers. 1991;59(3):339–54.
8. Mofsen AM, Rodebaugh TL, Nicol GE, Depp CA, Miller JP, Lenze EJ. When all else fails, listen to the patient: a viewpoint on the use of ecological momentary assessment in clinical trials. JMIR Mental Health. 2019;6(5):e11845.
9. Runyan JD, Steinke EG. Virtues, ecological momentary assessment/intervention and smartphone technology. Front Psychol. 2015;6:481.
10. Gee BL, Han J, Benassi H, Batterham PJ. Suicidal thoughts, suicidal behaviour and self-harm in daily life: a systematic review of ecological momentary assessment studies. Digital Health. 2020;6(25):20762963958.

11. Haagmans J. And global burden of disease study collaborators, the global burden of injury: incidence, mortality, disability-adjusted life years and time trends from the global burden of disease study 2013. Inj Prev. 2016;22:18–31.

12. Leske JS, Jiricka MK. Impact of family demands and family strengths and capabilities on family well-being and adaptation after critical injury. Am J Crit Care. 1998;7(5):383.

13. Richmond TS, Kaufer D. Predictors of psychological distress following serious injury. J Trauma Stress. 2000;13(1):681–92.

14. Page MJ, McKenzie JE, Bossuyt PM, Boutron I, Hoffmann TC, Boutron I, et al. The PRISMA 2020 statement: an updated guideline for reporting systematic reviews. BMJ. 2021;372:n71.

15. Baker S, O’Neill B, Gimsburg M, Guo-Hua L. The injury fact book. New York: Oxford University Press; 1992.

16. Rabinowitz AR, Fisher AJ. Person-specific methods for characterizing the course and temporal dynamics of concussion symptomatology: a pilot study. J Head Impair Rehabil. 2019;3(5):567–73.

17. Betthauser LM, Forster JE, Bortz A, Penzenik M, Hernandez TD, Bahraini N, et al. Strength and awareness in action: feasibility of a yoga-based intervention for post-acute mild TBI headaches among veterans. Contemp Clin Trials Commun. 2021;22:100076.

18. Albanese BJ, Preston TJ, Bedford C, Macatee RJ, Schmidt NB. Distress in persons with chronic pain and spinal cord injury. Qual Life Res. 2017;26(3):587–600.

19. Simon RM, Martin G, VA. An examination of diurnal variations in neuroendocrine pain and affect, on exercise and non-exercise days, in adults with spinal cord injury. Spinal Cord Ser Cases. 2018;4:94.

20. Todd KR, Shaw RB, Kramer JL, Ginz KAM. Using ecological momentary assessment to evaluate neuroendocrine pain experienced by adults with SCI. Rehabilitation 2018;4(3):113–9.

21. Pacella M, Prabhu A, Morley J, Huang S, Suffoletto B. Postconcussive symptoms over the first 14 days after mild traumatic brain injury: an experience sampling study. J Head Trauma Rehabil. 2018;33(3):E31–9.

22. Pacella ML, Girard JM, Wright AGC, Suffoletto B, Callaway CW. The association between daily posttraumatic stress symptoms and pain over the first 14 days after injury: an experience sampling study. Acad Emerg Med. 2018;25(8):844–55.

23. Price M, Ruggiero KI, Ferguson PL, Patel SK, Treiber F, Couillard D, et al. A feasibility pilot study on the use of text messages to track PTSD symptoms after a traumatic injury. Gen Hosp Psychiatry. 2014;36(3):249–54.

24. Price M, van Stolk-Cooke K, Ward HL, O’Keeffe M, Gratton J, Skalka C, et al. Tracking post-trauma psychopathology using mobile applications: a usability study. J Technol Behav Sci. 2017;21(1):41–8.

25. Price M, van Stolk-Cooke K, Legrand AC, Brier ZMF, Ward HL, Connor JP, et al. Implementing assessments via mobile during the acute posttrauma period: feasibility, acceptability and strategies to improve response rates. Eur J Psychotraumatol. 2018;9:1500822.

26. Carozi NE, Freedman J, Troost JP, Carson T, Molton IR, Ehde DM, Najarian K, Miner JA, Boileau NR, Kraul AT. Daily variation in sleep quality is associated with health-related quality of life in people with spinal cord injury. Arch Phys Med Rehabil. 2021;103(2):263–73.e4.

27. Wiebe DJ, Nance ML, Houskanech E, Grady MF, Otto T, Sandmark DK, et al. Ecological momentary assessment to accomplish real-time capture of symptom progression and the physical and cognitive activities of patients daily following concussion. JAMA Pediatr. 2016;170(11):1108–10.

28. Worthen-Chaudhari L, McGonigal J, Loggan K, Bockbrader MA, Yeates KO, Mysiw W. Reducing concussion symptoms among youth: evaluation of a mobile health app. Brain Inf. 2017;31(10):1279–86.

29. Sajid A, Rabinowitz AR, Jiricka MK, Guo-Hua L. The injury fact book. New York: Oxford University Press; 1992.

30. Spence P, O’Keeffe MC, Hurley B, Mannion C, Glynne J, Slane K, et al. The relationship between accelerometer-measured sleep and next-day psychological momentary assessment symptom report during sport-related concussion recovery. Sleep Health. 2021;28:28.
52. Axén I, Jensen I, Butler Forslund E, Grahn B, Jørgensen V, Opava C, et al. Frequently repeated measurements—our experience of collecting data with SMS. BMC Med Res Methodol. 2020;20:1–11.

53. Mitchell R, Braithwaite J. Evidence-informed healthcare policy and practice: using record linkage to uncover new knowledge. J Health Serv Res Policy. 2021;26(1):62–7.

54. Liu J, Zhao Y, Lai B, Wang H, Tsui KL. Wearable device heart rate and activity data in an unsupervised approach to personalized sleep monitoring: algorithm validation. JMIR Mhealth Uhealth. 2020;8(8):e18370.

55. Dunton GF, Dzubur E, Intille S. Feasibility and performance test of a real-time sensor-informed context-sensitive ecological momentum assessment to capture physical activity. J Med Internet Res. 2016;18(6):e106.

56. Rosenberg GM, Shearer EJ, Zion SR, Mackey SC, Morris AM, Spain DA, et al. Implementation challenges using a novel method for collecting patient-reported outcomes after injury. J Surg Res. 2019;241:277–84.

57. Rich JL, Chojenta C, Loxton D. Quality, rigour and usefulness of free-text comments collected by a large population based longitudinal study-ALSWH. PLoS One. 2013;8(7):e68832.

58. Juengst SB, Terhorst L, Nabasny A, Wallace T, Weaver JA, Osborne CL, et al. Use of mHealth Technology for Patient-Reported Outcomes in community-dwelling adults with acquired brain injuries: a scoping review. Int J Environ Res Public Health [Electronic Resource]. 2021;18(4):23.

59. Bai S, Babeva KN, Kim MI, Asarnow JR. Future directions for optimizing clinical science & safety: ecological momentum assessments in suicide/self-harm research. J Clin Child Adolesc Psychol. 2021;50(1):141–53.

60. Stone AA. Thoughts on the present state of real-time data capture. The science of real-time data capture: Self-reports in health research; 2007. p. 361–70.

61. Balasundaram AP, Athens J, Schneiders AG, McCrory P, Sullivan SJ. Psychological and lifestyle factors that influence the serial reporting of postconcussion-like symptoms in a non-concussed population. PM&R. 2017;9(9):866–73.

62. Mitchell R, Driscoll T. Calculating the risk of injury: do we underestimate the true risk of injury? Aust Epidemiol. 2012;19:4–6.

63. Allegrante JP, Mitchell RJ, Taylor JA, Mack KA. Injury surveillance: the next generation. Inj Prev. 2016;22(Suppl 1):i63–5.

64. Everhart RS, Borschuk AP, Mladich SA, Barell J, Heron KE. Caregiver daily experiences associated with child asthma symptoms. Am J Health Behav. 2018;42(2):50–60.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.