Lessons in Linkage: combining administrative data using deterministic linkage for surveillance of sports and recreation injuries in Florida, United States

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Abstract

Objectives
Previous and ongoing epidemiological surveillance of sports and recreation injuries (SRI) has been cross-sectional in nature, utilised a subset of injuries based on athletic trainer availability, or focused on elite and professional athletes. In the United States, surveillance is often prohibitively expensive and not well funded by national organisations or agencies, except for the case of some professional and elite sports. This paper details the methodology, barriers, and successes of using deterministic linkage to combine emergency department and hospitalisation data with a single identifier for use in surveilling sports injuries for persons aged 5 to 18 years.

Design
Data linkage of a population cohort.

Methods
We performed deterministic linkage of administrative emergency department and hospitalisation data from the state of Florida in the US. Data was acquired from the Florida Agency for Health Care Administration. With limited identifiers available due to privacy, we combined data across multiple years using a near universal identifier. We identified sport and recreation injuries using a modified External Cause of Injury Morbidity Matrix and ICD codes across all possible diagnoses. Finally, we obtained descriptive statistics of records that were successfully linked and those that were not to assess similarities between the groups.

Results
We found 384,731 visits for SRI over a seven-year period. We were able to link approximately 70% of the records using a single identifier. There were statistically significant differences by age, sex, payer, and race/ethnicity for the records that were linked compared to the records that were not linked.

Conclusions
This study is significant because while similar methods have been used to examine other conditions (e.g. asthma), few have linked multiple types of administrative data especially with nearly no identifiers to examine sports and recreation injuries. This method was found useful to identify injuries over time for the same individuals seeking care in emergency departments, or in hospital inpatient settings, though future work will need to address the limitations of this method. If we expect to move health surveillance forward as budgets for it become even more limited, we must develop and improve methods to do it with fewer resources, including using data that has great limitations.

Keywords
deterministic linkage; epidemiological methods; epidemiology; surveillance; public health; data cleaning; data linkage; reformatting; SAS; Python; injury surveillance; disease surveillance; health surveillance; biostatistics

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Introduction

De-identified hospital and emergency department data are often used when assessing the health of a population. This is advantageous when examining the prevalence of health problems because each visit is treated as a discrete occurrence, but disadvantageous when examining the issues such as the effect of repeat injuries on individuals (person-level). In certain situations, we have access to identifiable information and this permits a different type of epidemiological assessment that is often limited to primary data collection. It becomes possible to identify how many encounters an individual has with the medical system over time and a better understanding of the variance in health care in the population based on patient characteristics such as prior health history. This could be useful in determining, for example, how primary care physicians can better intercede with patients to reduce the number of preventable sports injuries.

Involvement in sport and recreation promotes health and overall well-being [1, 2]. However, these activities expose individuals to a variety of elements that heighten risk for injury. Studies indicate millions of sport and recreation related injuries (SRI) are treated in emergency departments (ED) each year [3–8] and the incidence and severity of injuries within sport continue to garner increased attention from prevention professionals, the media, and participants [9–11]. Data gathered from EDs often represent the best available resources for exploring SRIs across a broad geographical region. Administrative records from EDs have been utilized to establish the prevalence of SRIs and common mechanisms of injury among different populations [8, 12–16]. In the United States, all fifty states and Washington, DC capture administrative data related to clinical visits, though the information included from each state varies in depth and public availability. Despite continuing work to explore both the risk factors for SRIs as well as the nature and severity of these injuries, limited information exist regarding the burden of these injuries on the healthcare system, particularly the burden associated with recurring injuries to the same individual.

Public health surveillance is the ongoing systematic collection of data for quantifying the burden of a problem (and thus developing priorities for addressing it); for monitoring trends of an issue; for identifying new problems; and for the evaluation of prevention efforts [17]. Surveillance can take place at various levels, e.g. national, state, and local community, and can consider both geographic, demographic, and policy differences between groups. Surveillance provides the information necessary to change or support policy at an appropriate intervention level. It also identifies where more resources are necessary to make a difference. Some issues, such as sports and recreation injuries, require various interventions at multiple policy levels for prevention and evaluation. Previous and ongoing epidemiological surveillance of SRI tends to be cross-sectional in nature [18, 19], be collected for a limited geographic area [20], utilise a subset of injuries based on athletic trainer availability [21], or focus on elite and professional athletes [22–26]. This surveillance is necessary to better identify what injuries and other health concerns are happening to sports and recreation participants, identify factors for intervention, and test prevention programs [27–29]. But surveillance is also often prohibitively expensive [30–33] and, in the United States, not well funded by national organizations or agencies except for the case of some professional and elite sport.

This problem is not without potential solutions. Multiple countries have identified that we should work more often to use existing, or secondary, data for injury surveillance and prevention [34]. It has also been recommended that we improve or establish injury surveillance systems given the high burden of morbidity and mortality due to unintentional injuries [34]. A sustainable injury surveillance system that captures the same or most of the same information as traditional active, passive, or syndromic surveillance could be a feasible way to reduce costs without reducing quality. Optimally, this alternative would also allow data to be combined with other data sources such as community level, census, education, or geographic data to provide richer context for health findings. Health administrative data, such as records about emergency department discharges, is an existing data source that is regularly updated. This type of data is used in the United States for required reporting, for research, and for policy development. Discharge data has been used to surveil various health issues [35] including diabetes [36], asthma [37], and arthritis [38]. The purpose of this paper is to describe the data linkage methodology we used to establish a surveillance data set for SRI using discharge data in a single state of the United States. We expect that this strategy can be applied to a variety of data in and outside of the US to study issues in SRI.

Methods

This paper follows the guidelines of the RECORD statement (REporting of studies Conducted using Observational Routinely-collected health Data), an extension of the STROBE statement (Strengthening the Reporting of Observational Studies in Epidemiology) [39] (https://www.equator-network.org/reporting-guidelines/record/).

Institutional review board approval

This study was approved by the Florida Agricultural and Mechanical University Institutional Review Board (#014-47). A HIPAA Waiver of Authorization was granted to permit the research without needing individual patient participation approval.

Data sources

In Florida, data from hospitals, ambulatory care centers, and EDs is captured on a quarterly basis by the Agency for Health Care Administration (AHCA) [40]. Inpatient data has been available since 1988, and ED data since 2005. The inpatient data includes short- and long-term acute care hospitals and psychiatric hospitals. The ED data includes hospitals that have EDs but does not include freestanding EDs. Freestanding EDs are licensed EDs that are physically separated from a hospital or other clinical facility and can be independently operated or as an offsite or satellite facility of a hospital [41]. Only facilities affiliated with hospitals are recognized by the Centers for Medicare and Medicaid Services (CMS), meaning
independently operated (non-hospital affiliated facilities) can be more selective about who they serve and cannot bill Medicare (a primarily income-based insurance program) or Medicaid (a primarily age-based insurance program). A listing of all independently owned and operated facilities is available from AHCA. By default, the AHCA data includes all discharges from non-independent inpatient, and ED facilities no matter the state or national residency of the patient. Diagnosis information in all AHCA data is codified using the International Classification of Diseases (ICD), specifically the United States Clinical Modification (CM). Procedure details are coded using Current Procedural Terminology (CPT) codes. A listing of variables captured can be found for each data set on the Florida Health Finder website for researchers using AHCA data [40] and in the Data Catalog [42]. Details on more secure variables such as identifiers are available after contacting the Agency. All variables included in this project can be found in Table 1.

Identifying a target population

Every data linkage project should clearly define the population under study. For surveillance purposes, the population source needs to be consistent over time and inclusive of all events in that time and space. This is in line with three of the criteria for a successful surveillance system – quality data, stable, and representative [43]. Optimally, a census (a complete enumeration of a population) rather than a sample (finite part of a statistical population) should be used to identify the prevalence (burden) of the health problem. Not only is it necessary to capture the universe of potential visits, it is imperative to ensure that the population under surveillance is consistent across databases used for linkage. The goal of this project was to create a data set that was representative of Florida residents aged 5 to 18 years of age (childhood/adolescence) seen in an ED or hospital with an unintentional SRI (Figure 1). Age in this study is captured as the patient age on the date of ED attendance or hospital admission. The age range was selected in this project to best align with childhood and adolescence in the US as aligned with typical primary and secondary school attendance.

A major step to assess data consistency is to determine the availability and format of data across time. As expected, the variables collected and the format of how those variables were collected/encoded changed in the duration of each AHCA database’s (i.e. inpatient, ED) existence. The type of errors that this can cause are explained in detail here [44].

There are more codes for SRI in ICD-10-CM than exist in ICD-9-CM. The increased level of coding specificity allows for better capture of injuries, yet affects the comparability of rates across time periods. Future changes to the ICD (such as the use of ICD-11-CM) can create a similar problem and needs to be accounted for. As we started this work, the United States was using ICD-9-CM and migrating toward the use of ICD-10-CM. Now, that switchover is complete and if used States was using ICD-9-CM and migrating toward the use of needs to be accounted for. As we started this work, the United

Strategy to establish the study population

When linking data, an initial action should be distinguishing what the final data set should look like – from what variables and observations should be included to how they should be included and formatted. The RECORD Statement recognizes “three levels of population hierarchy that are relevant in studies using routinely collected data” – the source population [the population that the data comes from], the database population [people with data in the data source], and the study population [those identified within the database population that fit the study criteria] [39] (Figure 1). As previously stated, our goal was to create a data set that was representative of Florida residents aged 5 to 18 years of age injured during sport or recreational activities (source population). There currently is no surveillance system that completely tracks this population and measures every health issue or outcome. However, we had access to administrative data that could be used to address the topic. This data – hospitalisation and ED data from AHCA in Florida – is known as the database population. We further narrowed the database population down to the study population using a variety of ICD codes and tools discussed below. We were most concerned with keeping the information for each visit a person had to an ED or hospital during the period January 1, 2006 to December 31, 2012.

Data linkage often includes two steps – merging and concatenation. Merging refers to putting data together side by side that might be from the same or different sources; as an example, one data set contains demographic information and the other data set contains visit information for those same individuals. There are several types of merges. If one data set contains multiple rows for the same individual and other data has a single row, this is referred to as a ‘many to one’ or ‘one to many’ merge depending on which set is the ‘main’ data set. If both sets only have one row per individual, this is a ‘one to one’ merge. When merging data, if columns have the same name, same named columns will often be deleted.

When information is available for different groups of people over time, such as health visits over multiple years, data should be stacked instead of being put side by side. This is called concatenation. If you are unsure whether observations in a given data set belong to an individual, they can be treated as discrete pieces of information and thus you will likely use concatenation more than merging. A major consideration when concatenating data is that the variables in each data set being stacked have the same names and formatting. Failure to abide by this can result in loss of data and/or a data set resulting in duplicate columns. For brevity, many existing papers refer to either of these processes simply as merging or data linkage. We will distinguish between merging and concatenation throughout this paper for clarity.

We obtained access to an identifier variable (the masked social security number is discussed below) and wanted our resulting data for an individual to be on a single row, meaning even if an individual had multiple visits their data would still be a single row. When using data from any clinical system,
Table 1: Variables included in the longitudinal sports and recreation injury study

<table>
<thead>
<tr>
<th>Variable name</th>
<th>ED</th>
<th>Hospital</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>E-Code</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Length of Stay</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Masked Identifier</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Principal Diagnosis</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Other Diagnosis</td>
<td>9</td>
<td>30</td>
</tr>
<tr>
<td>Principal Procedure</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Procedure</td>
<td>4</td>
<td>30</td>
</tr>
<tr>
<td>Payer</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Patient Residence</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Patient Status at Discharge</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Quarter of the Year</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Admitting Diagnosis</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Day of the Week</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Sex</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Race and Ethnicity</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>ED Hour of Arrival</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Year of Service</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Admission Source</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Type of Admission/Priority of Admission</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Time of Admission</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Days to Principal or Other Procedure</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Present on Admission Indicator for E-Codes</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Present on Admission Indicator for Other Diagnoses</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Present on Admission Indicator for Principal Diagnosis</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>ER Charges</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Trauma Charges</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Physical or Occupational Therapy Charges</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Total Gross Charges</td>
<td>1</td>
<td>1</td>
</tr>
</tbody>
</table>

Numbers indicate the number of variables for a particular item (e.g. there is 1 sex variable).

Identifying the identifiers

When linking administrative data, particularly discharge data for surveillance of a health issue, there are several important considerations including the necessity to have a consistently used universal patient level identifier, or combination of identifiers, across years of data collection such as a date of birth or a national identification number. In the United States, there is no national identification number so the Social Security Number (SSN), a unique number allocated to each individual person, is often used as a proxy. The SSN is issued by the US Social Security Administration, an agency established to oversee an anti-poverty program – Social Security – in 1935 via the Social Security Act [49]. Social Security includes benefits for retiring workers after age 65, disability insurance, health insurance (Medicare), survivors’ insurance, and supplemental income for those with disabilities including children. SSNs are issued by the US Social Security Administration when applications are filed, often at birth, but registration is voluntary. An SSN is required for people that file US income taxes, open bank accounts, buy savings bonds, apply for government services, or get a job in the United States. It allows the agency to “identify and accurately record your covered wages or self-employment earnings”, and “to monitor your record once you start getting benefits” [49]. Non-citizens that are authorized to work in the United States can get an SSN but, to be clear, as it is not required, there
are people seeking health care in the US that do not have an SSN. Something like a driver’s license number or residential (or mailing) address is not sufficient as a universal identifier as 1) these values can change with varying frequency and 2) everyone does not have them. When possible, these identifiers can be used in combination with other identifiers such as the SSN to successfully identify a single individual. As societies change and values or constructs of those societies change, the utilisation of socio-demographic variables such as race, ethnicity, sex, and gender may be inconsistently captured. Race and ethnicity are defined and utilized in many ways across time, vary in terms of categorization, and vary by source; some data may be patient identified and other data may be nurse identified, among other ways of capture. Many scientists have previously misused the word gender to identify biological differences in people so many sets designed and collected in the past may interchangeably use the term sex and gender to mean the same thing – biology. However, as we know better, newer data sets are better defining and utilising the term gender and doing the same for sex. The identifiers and demographic variables are not just data points but markers of society. The longer the time period being linked, the more difficult it may be to identify a consistent set of identifiers no matter which ones are used.

In the present study, we had limited identifiers available in the data that is available to researchers. We chose to use a masked version of the SSN as the lone identifier. This was a value that was randomly and securely created using alphanumeric characters by AHCA in lieu of the actual SSN. This is done to anonymize the actual value and safeguard the patient’s SSN. Every time data is drawn from the original source for research purposes, the mask changes. For example, an actual SSN is a nine-digit number (e.g. 000000000) while the masked version might be Az024M12NGq81a. Note that the example value of 0 is not always recreated as the same alphanumeric digit but every visit for the individual with the SSN of 000000000 would be Az024M12NGq81a. Also note that the masked number is not the same length of the original SSN. The rationale behind using the masked SSN in this study was – 1) changes in the data across time to the race and ethnicity variables to make it difficult to know that using the variables would consistently match patients; 2) the benefit of being able to identify most individual patients using the masked SSN; 3) no other address information was available; and 4) no other consistent identifier was available in data researchers could access to prevent re-identification of patients.

Cleaning the study population data

Data obtained from AHCA included a variable to indicate residence of the patient which we used to exclude any visits that were not for Florida residents. We then identified the unintentional injury discharges using the ICD-9-CM external cause-of-injury codes (E-codes). E-codes are used to classify injury by mechanism (e.g. struck by/against) and intent (e.g. unintentional). These codes are in addition to the “main” ICD codes. In ICD-10-CM and beyond, external cause of injury codes are included as activity codes (V00-Y99) and provide a greater level of detail about the event. Because they are better integrated into ICD in versions 10 and beyond, we anticipate greater adoption and utilization which will improve linkage potential. Tools and recommendations to examine injury data can be found on the website of the Centers for Disease Control
Table 2: Altered external cause of injury morbidity matrix SAS syntax

Original:*

IF (CAUSEDET GE 800 AND CAUSEDET LE 869) OR (CAUSEDET GE 880 AND CAUSEDET LE 929) THEN INJ = 1;

Updated:

IF (CAUSEDET GE 800 AND CAUSEDET LE 869) OR (CAUSEDET GE 880 AND CAUSEDET LE 929) THEN INJ = 1;
IF (CAUSEDET2 GE 800 AND CAUSEDET2 LE 869) OR (CAUSEDET2 GE 880 AND CAUSEDET2 LE 929) THEN INJ2 = 1;
IF (CAUSEDET2 GE 800 AND CAUSEDET2 LE 869) OR (CAUSEDET2 GE 880 AND CAUSEDET2 LE 929) THEN INJ3 = 1;
UNINTENTIONAL = 0;
IF INJ = 1 OR INJ2 = 1 OR INJ3 = 1 THEN UNINTENTIONAL = 1;

* GE means Greater than or equal to.
† LE means Less than or equal to.

a Original Syntax provided by the Centers for Disease Control and Prevention National Center for Injury Prevention and Control [50, 51]. One of these tools, the External Cause of Injury Morbidity Matrix, was designed to use the first, or primary, of available E-codes in recorded data. Our data contained three E-code variables (Figure 3). In Figure 3, you can see an example data frame with an identifying variable (ID), five diagnosis variables (DX1-DX5), and three E-codes (E1-E3). As we found, the user may have an inconsistent number of diagnosis variables or E-codes across time. The method as described is adaptable to any number that the user has available [52]. It is also adaptable if the user has more than one identifying variable.

Due to the potential arbitrary placement of E-codes in the available fields and a want to include all possible instances of unintentional sport and recreational injury, we altered the SAS [53] syntax provided by the CDC [54] to include all E-code variables. The original syntax [54] created and used three variables – CAUSEDET (first 3 digits of the ICD code after the E), CAUSET4 (first 4 digits of the ICD code after the E for morbidity data), and PERSINJ (4th digit of the ICD code after the E) – to reorganize data. Our alteration was to make a CAUSEDET variable for all three E-codes we had available (Table 2). The original first E-code variable (E1) in Figure 3 was used to create the new variable CAUSEDET, the second (E2) was used to create CAUSEDET2, and the third (E3) was used to create CAUSEDET3. For example, if the data for the first visit for patient ‘Az024M12NGq81a’ had a valid response for the first E-code (E1). This response is the value ‘E849.4’ and so the new variable CAUSEDET would contain the value 849 because they were the first three digits after the E. We could then use CAUSEDET, CAUSEDET2, and CAUSEDET3 to create variables (INJ, INJ2, and INJ3) indicating whether any of the three E-code variables contained data on an unintentional injury as opposed to some other external cause of injury [Table 2]. Since our intention was to only retain observations where an unintentional injury had occurred, we then created an indicator variable (UNINTENTIONAL) to identify and restrict the data set. Complete example syntax with instructions can be found online [52].

Finding and including sports and recreational injuries

The next step of the cleaning process was to identify which records contained SRI. A number of ICD-9-CM E-codes were available for this purpose (Table 3) [55]. While not the most inclusive, these variables narrowed down the number of observations to injuries that occurred in places for sport and recreation (E849.4), using recreational equipment (e.g. E885.2 [fall from a skateboard]), or while participating in sport or recreational activities without equipment (e.g. E910.2 [accidental drowning and submersion while engaged in other sport or recreational activity without diving equipment]). If the value of any E-code variable matched these codes singularly
Final cleaning steps

Normal considerations for combining data include standardizing inconsistent coding or formatting data sets and regular updates to data categories such as sociodemographic factors or causes of injury that improve the usefulness of the data. To combat the latter, we used a two-step process involving available codebooks and examination of the data sets. We identified every possible version of a variable and determined how we could make them consistent. For example, in Figure 4 you can see that from 2006 to 2009, race and ethnicity were in all of the data sets as a single combined variable (RACE). In 2010, however, they were separated into two separate variables (RACE and ETHNICITY). These two variables were more inclusive of possible categorizations. To reformat the 2006 to 2009 data into separate race and ethnicity variables likely would have resulted in a larger degree of misclassification of observations than combining the data from the race and the ethnicity variables from 2010–2012. Thus, we elected to use the older categories from the 2006–2009 RACE variable for the linkage. We updated all data prior to concatenating years. While this method was often used to prevent a massive loss of data, it also meant we were unable to take advantage of newer and improved variable formats and categories.

Merging and concatenating data

Figure 5 shows each data set as it is restricted by patient residence, unintentional injury status, and then SRI status. Next the data is concatenated by source and finally the sources are merged into our final study population data set. The next section and Figure 6 provide more detail for preparing the data from each source for merging. In order to complete this process, we needed at least one identifying variable for individual people. We used the masked SSN.

Merging all the data together for surveillance

The data sets for each type (ED and Hospitalisation) were obtained in annual extractions (e.g. 2006, 2007, 2008) and needed to be combined (concatenated). After concatenating the data by type (Figure 6, Step 2) we then sorted each of the two new data sets (ED and Hospitalisation) by masked SSN (unique identifier). This allowed us to see how many total visits existed in each set and how many visits existed for each unique individual. After creating a visit specific variable (Figure 6,
Step 3), we then used this new variable to transform (or rearrange) the data frame. The type of transformation used is also known as transposing. To transpose data generally means one of two things. The first is often called changing the data from wide and long (one row per visit) to long and narrow (one row per variable). This can be seen in Figure 6, Step 4. We change the data frame from a point where the columns are the individual variables (wide) and the rows represent individual visits (long) to a point where each row represents individual variables related to specific visits (long) and the columns are for each individual person (not visit) in the data set (narrow).

We used this transformation to add a prefix (‘V1_‘ in Figure 6, Step 4) to each visit specific variable. For example, in Figure 6, Step 4 you can see that the first diagnosis variable (DX1) for the first visit for subject ‘ABC’ is now row V1_DX1 in the data frame.

The second type of transpose is known as changing the data from long and narrow to short and wide. Think of it as almost reversing the first type of transpose. We can see how this works as we move from Step 4 to Step 5 in Figure 6. Because we are using this transformation on our new data frame (the one with the prefixes), when we transform the table...
Figure 6: Steps to transform the single data sets into concatenated data sources in preparation for merging.

Each individual person’s data will be on a single row instead of across multiple rows. This makes the data set (top to bottom) shorter and (side to side) wider than the original data frame, hence the phrase ‘short and wide’. The original data frame (the result of Figure 6, Step 2) is noticeably different than the resulting data frame (the result of Figure 6, Step 5). This transformation put each individual person’s discharges on a single row and created a requisite number of variables to retain all discharge information.

At this point either dataset (ED or hospital inpatient) could be analysed separately or merged to create one single and complete data set with ED and inpatient data.
Identifying the injuries

For injury surveillance purposes, it was important to align the now combined data to existing tools such as the Barell Matrix [56] to classify the injuries by body region (e.g. shoulder) and nature of the injury (e.g. fracture). One observed barrier to using the Barell Matrix (and the subsequent Injury Mortality Diagnosis (IMD) matrix for ICD-10) is that it was developed to only use the primary diagnosis on any record no matter how many diagnoses were available. However, our experience has indicated that the order of diagnoses, is arbitrary or based on the cost of the illness or injury to facilities and insurers rather than in the order of seriousness as intended. Because all aspects of injury regardless of seriousness impact the population, our goal was to capture and count all injuries, not just primary diagnoses or the total number of individuals injured. As such, we modified the Centers for Disease Control and Prevention’s SAS syntax [57] for the Barell Matrix to include all available diagnoses in the data sets in a fashion similar to our alteration of the E-code variables. Because of the importance of how this was done and to not detract from the current work, specific methods for this process and population level statistics using this data will be published separately.

Results

Evaluating linkage quality

We calculated the frequency of the masked SSN in order to identify any outliers. We found that 2 identifiers had an improbable number of discharges (e.g. greater than 100 observations in a single year). We verified with the data owner, the Agency for Health Care Administration, that these particular identifiers were catch-all identifiers for people that did not have an SSN recorded on their discharge. In lieu of enough available data to perform probabilistic linkage, we decided to exempt these records (30.7% of ED data; 36.8% of hospitalization data) from the data set for individual level (person-level) analyses (Table 4). This data was still useful when looking at population wide measures such as counting the total number of SRI especially by sex or race/ethnicity.

As eloquently explained and demonstrated by both Gilbert [58] (GUILD guidance) and Harron [59], there is a significant need to not only report the link or the methods to achieve the link, but also certain factors such as bias that can indicate whether a link is low, medium, or high quality linkage. Lacking a gold standard for our project, we opted for sensitivity analyses that compared linked (those with a valid masked SSN) and unlinked data (those without a valid masked SSN) then identified potential biases that could occur and how they might affect the results. These analyses indicated that there were narrow but statistically significant differences by sex, age, and race/ethnicity (Tables 5, 6).
Table 4: Deterministic linkage statistics in the longitudinal sports and recreation injury study – individual persons represented

<table>
<thead>
<tr>
<th>Source</th>
<th>Total visits</th>
<th>Excluded visits</th>
<th>Visits after exclusion</th>
<th>Individual people represented</th>
</tr>
</thead>
<tbody>
<tr>
<td>Emergency Department</td>
<td>376,179</td>
<td>115,467</td>
<td>260,712</td>
<td>215,718</td>
</tr>
<tr>
<td>Hospital (Inpatient)</td>
<td>8552</td>
<td>3145</td>
<td>5407</td>
<td>5329</td>
</tr>
</tbody>
</table>

Table 5: Deterministic linkage statistics in the longitudinal sports and recreation injury study – differences in emergency department population demographics

<table>
<thead>
<tr>
<th>Linkage status</th>
<th>Excluded</th>
<th>Included</th>
<th>Comparison test</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age* in years</td>
<td>Mean 12.2</td>
<td>Mean 13.0</td>
<td>T-Test −75.4</td>
</tr>
<tr>
<td>*Normality tested using the KS test</td>
<td></td>
<td></td>
<td>p &lt; 0.0001</td>
</tr>
<tr>
<td>Sex (Male)</td>
<td>71.3%</td>
<td>74.1%</td>
<td>X²315.7</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>p &lt; 0.0001</td>
</tr>
<tr>
<td>Race/Ethnicity</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>American Indian/Alaska Native</td>
<td>0.2%</td>
<td>0.1%</td>
<td>X²6062.8</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>p &lt; 0.0001</td>
</tr>
<tr>
<td>Asian/Pacific Islander</td>
<td>0.8%</td>
<td>0.5%</td>
<td></td>
</tr>
<tr>
<td>Black</td>
<td>14.9%</td>
<td>26.5%</td>
<td></td>
</tr>
<tr>
<td>White</td>
<td>60.7%</td>
<td>52.5%</td>
<td></td>
</tr>
<tr>
<td>White Hispanic</td>
<td>16.2%</td>
<td>14.0%</td>
<td></td>
</tr>
<tr>
<td>Black Hispanic</td>
<td>0.4%</td>
<td>0.4%</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>6.9%</td>
<td>6.2%</td>
<td></td>
</tr>
<tr>
<td>Payer (Commercial health insurance)</td>
<td>32.9%</td>
<td>18.8%</td>
<td>X²23535.91</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>p &lt; 0.0001</td>
</tr>
</tbody>
</table>

Table 6: Deterministic linkage statistics in the longitudinal sports and recreation injury study – differences in hospital population demographics

<table>
<thead>
<tr>
<th>Linkage status</th>
<th>Excluded</th>
<th>Included</th>
<th>Comparison test</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age* in years</td>
<td>Mean 12.15</td>
<td>Mean 13.04</td>
<td>T-Test −11.2</td>
</tr>
<tr>
<td>*Normality tested using the KS test</td>
<td></td>
<td></td>
<td>p &lt; 0.0001</td>
</tr>
<tr>
<td>Sex (Male)</td>
<td>77.3%</td>
<td>81.0%</td>
<td>X²16.6</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>p &lt; 0.0001</td>
</tr>
<tr>
<td>Race/Ethnicity</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>American Indian/Alaska Native</td>
<td>0.2%</td>
<td>0.2%</td>
<td>X²84.8</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>p &lt; 0.0001</td>
</tr>
<tr>
<td>Asian/Pacific Islander</td>
<td>0.9%</td>
<td>0.6%</td>
<td></td>
</tr>
<tr>
<td>Black</td>
<td>16.3%</td>
<td>24.1%</td>
<td></td>
</tr>
<tr>
<td>White</td>
<td>58.2%</td>
<td>53.6%</td>
<td></td>
</tr>
<tr>
<td>White Hispanic</td>
<td>14.8%</td>
<td>14.4%</td>
<td></td>
</tr>
<tr>
<td>Black Hispanic</td>
<td>0.5%</td>
<td>0.4%</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>9.2%</td>
<td>6.6%</td>
<td></td>
</tr>
<tr>
<td>Payer (Commercial health insurance)</td>
<td>35.0%</td>
<td>22.8%</td>
<td>X²294.3</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>p &lt; 0.0001</td>
</tr>
</tbody>
</table>

These differences persisted across years. This means that we must acknowledge that any results gained about the impact of sports and recreational injuries from this linked data is biased and may be less applicable to certain subgroups of the population. Further examination and potential linkages must be explored. With such a degree of missingness, imputation alone may not be a reasonable option, however may be one of the tools necessary to minimize loss of records and improve both accuracy and representativeness of the study population.

Discussion

In this manuscript, we discuss in detail the methodology, barriers, and successes of using deterministic linkage to
combine emergency department and hospitalisation data with a single identifier for use in surveilling sports injuries across a seven-year period. This is significant because while similar methods have been used to examine other conditions (e.g. asthma), few have linked multiple types of administrative data especially with nearly no identifiers, nor did they specifically use this methodology to examine SRI. The steps described in this manuscript will not only allow the authors to complete future work looking at how well we can track SRI as the ICD continues to change, but others can use these methods with country specific data and expand the method to add other sources that use similar identifiers. If we expect to move data surveillance forward as budgets for data surveillance become even more limited, we must develop and improve methods to do it with fewer resources including data that has great limitations.

The use of deterministic linkage is not novel, but the particular combination of strategies employed here – using diagnoses and external cause of injury codes (E-codes) beyond the primary to make better use of all data; creating a static data set with all data over a period of time in one row per patient; and specifically applying these methods to the topic of sports and recreation injury – are novel in their use. We believe improved training of researchers and practitioners in these methods and their underlying theories will widen their use. For example, analysts need to understand that acquiring data can be much more complicated than just downloading from a website. Health data that is protected (e.g. contains information that could be used to directly identify an individual) in the US is subject to the Health Insurance Portability and Accountability Act (HIPAA) which dictates a set of rules for accessing and engaging this information. The particular data used in this study required multiple telephone and email conversations with AHCA to discuss the project and the feasibility using the data; a HIPAA waiver of authorisation from our institutional IRB; a separate application to the government of the State of Florida through AHCA that needed to be approved by multiple state officials; receipt of the data; and separate receipt of secure keys to access the data. This process took about three months. After data was received, any additions to who accessed the data needed to be confirmed with AHCA. At points where more data became available (e.g. a new year of data), the entire process started again and all analyses needed to be re-run because the masks for the SSNs change with every pull of the data. Both our institutional IRB and the State of Florida required annual renewals of approval. These were tedious though not impossible steps. Sometimes emails to generic agency email addresses (e.g. data@agency.gov) never get a response. We were lucky to have a known contact at the agency to work through our issues with and have fostered a long-standing relationship that we hope will be fruitful to make changes to how data is made accessible for all research and practice users and how we can make a greater number of identifiers available without compromising the integrity of the data. Future collaboration with AHCA and similar agencies is necessary to make previous linkages of this data available for all end users of the data without more barriers in place. We find that this is not impossible but the result of longstanding (though understandable) communication and trust issues between the data stewards and researchers. Secure, source hosted versions of tools such as GitHub and volunteer moderators to work with state agencies could be the next step to make this a reality.

As data becomes more and more costly to obtain and funding for new surveillance sources is at a minimum, we find it important to have completed this work to provide data for future injury policy and prevention. We also feel these methods, as explicitly laid out here, can be applied to a variety of data sources and across a wide variety of health topics. Most notably, these methods may be quite applicable to areas with limited resources or low prevalence such as rare diseases and disorders. This study was completed using data from the US, but as with many methods, we know it to be applicable across countries though with caveats of how data is captured and what identifiers are available to the public.

A goal of this work was to create steps that could be followed by people with some knowledge and expertise in data and statistical programming, yet also accessible to those without large financial resources. If the point of surveillance is to help identify health issues and places for improvement, we felt it was necessary to contribute to these achievements. Relational database languages such as SQL (structured query language) were considered for this project, but would have had significant limitations for our purposes. The data as obtained from AHCA was hierarchical in nature, meaning that there might be many pieces of data (known as ‘children’) but the data belonged to an individual (known as the ‘parent’). The main limitation we considered was the difficulty in SQL to use hierarchical data without specified depth (e.g. various numbers of diagnosis variables per person and per data source) without expending a tremendous amount of resources (e.g. server or computer space, time, money, very specific software) [60]. The complexity of the data structure and the amount of data that SQL would need to handle it would require exorbitant funds to scale the system appropriately to support the size and the performance necessary for even simple queries because SQL’s use of memory. In other words, every query would potentially add the need for more computing resources (think: hard drives and RAM). Performance is lost when data needs to be split up on too many servers or other computing machines. Newer additions to SQL since this work started have made dealing with hierarchical data easier but these additions are still dependent on the vendor for the SQL software [61]. Clinical organizations might be able to minimize these limitations, but more independent researchers, researchers at institutions with limited computing resources, and various government institutions such as health departments would not be able to consistently provide these types of funds or maintain these size systems long term. It would also be difficult to share data and methods as the data becomes even more complex. In the end, we decided to create methods that would benefit the many as opposed to the few. Users with the resources to use SQL or other relational database programs for data at scale may consider an alternative method more acceptable.

**Successes and future potential**

Some past studies of data linkage in sport and recreational injury, nearly all of them using data with many available identifiers, are linking data from an investigator led primary
data collection study with secondary data, or link data limited in scope. For example, Ackerman et al linked individual level data regarding knee injuries in Australia but achieved this by having the government agency link the data then provide it to the investigators [62]. In the present study, we obtained the data and performed the linkage. Both methods achieve an end goal, yet if researchers were able to obtain more unique information without overburdening data providers, it would be easier to achieve this type of work. The present study linked multiple data sets from the same agency together and linked data across sources (ED and hospitalisation). A specific goal was to only use secondary data which was achieved, but this manuscript did not demonstrate how to then take the linked data and combine it with data from other secondary data sources or with primary data as these very important tasks are outside the scope of this work. However, these are important aspects others have addressed and that need to be addressed as this project progresses.

Beauchamp et al used a stepwise approach in their study of de-identified records for hospital admissions in Victoria, Australia and compared the effectiveness of each of their methods [63]. Beauchamp had access to names, hospital names, full birth date, and zip codes (post codes) and were able to successfully link 87% of their cohort study population to hospitalisation data [63]. Clark et al matched ambulance data to emergency department records with no identifiers in England [64]. The process was time intensive (65 weeks), and required both a large team and external input [64], but resulting in an average of 81% data matched (range of 50–94% depending on the hospital trust) [64]. Though we matched approximately 70% of the data on a single identifier, the limitations of using state specific patient level data from Florida was that obtaining an identifier for linkage led to a vast amount of other identifiable data becoming unavailable for access for research purposes such as patient zip code, date of birth for more accurate age assessment, and zip code of the facility. The single identifier we had available, masked SSN, was not required to be recorded for patients. While residents of many other countries have national identification numbers, US citizens and residents do not. Had other variables been available along with the masked identifier, we could have potentially used probabilistic linkage or at least matched records with more certainty. It occurred to us that we potentially could have matched records using a combination of the masked identifier, sex, and race. However, discrepancies in how race and ethnicity were captured across time combined with how they are not self-identified in this data would have greatly reduced the match. Additional discrepancies for sex across time may not have occurred with the data we used in this project, but could be a factor in the future as people are able to more accurately identify sex and gender in medical records.

This project was achieved using SAS 9.4 software [53]. We have also begun to post equivalent examples using Python, specifically Python 3.10.2 [65]. One limitation to prior work by others in this space was to solely use a single software, often expensive software inaccessible to others around the world. We count using publicly available data for surveillance as a great success but to do it in a way that increases accessibility and applicability of these methods with multiple types of software as an even greater victory. This process certainly was not without a number of challenges that must be addressed as this project moves forward.

Using data only from more recent years with consistent categories or improved categories will alleviate the problem of needing to recode or recategorise information, but it causes the loss of data that provides a better longitudinal outlook. This is not an uncommon problem – as we know better in science, especially when it comes to the capture of sociodemographic data, we do better. Progress should not be held up solely for need of long-term comparison. We address these limitations and potential methods for minimising their effect in future work. Furthermore, while it is outside the scope of this work, we do encourage continued efforts such as those by Lucyk [66] to qualitatively examine the processes that come together to make administrative data such as the discharge data we used in an effort to improve them. The complexity of linking data together across time cannot simply be explained by the methods as we describe them here, but in all the hands and applications it takes to get the data from point A to point Z.

There is quite a bit of future potential in this work. We did not address the application of these methods to ICD-10 data, combining ICD-9 and ICD-10 data for surveillance purposes, demonstration of the applicability of the data resulting from this data, and the application of existing tools (e.g. Barell Matrix) and new tools (Baker-10 Matrix) here. We will address each of these points in soon to come manuscripts. Syntax for this manuscript and future manuscripts will all be made available for use, for critique, and for modification with a GNU GPLv3 license in both SAS and open software (i.e. Python and R). Upon request, we will translate this work to other programs such as Stata and SPSS to increase usability.

**Conclusion**

We hope that readers are left with an understanding of and use of discharge data for surveillance of sport and recreation injuries. The quality of the data used for linkage and surveillance is always dependent on the source that the discharge data originates from, usually a medical provider. Because of this, these sources may not be as in depth or explanatory as data collected specifically for the purpose of surveillance or research, but has a great benefit in being affordable and quickly accessed. Reformattting the data into a usable format for surveillance requires several considerations in advance of acquiring and combining the data. Whether the data comes from a government, an organization, or directly from clinical records, it is important to be clear of your purpose up front and imperative to work with data managers to clarify any abnormal findings in the data.

**Ethics statement**

Ethics approval was granted by Florida Agricultural and Mechanical University (IRB # 630088).

**Conflicts of interest**

The authors have no conflicts to disclose.
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References


33. University of Minnesota Center for Infectious Disease Research and Policy. Costs to develop and maintain a state biosurveillance system:


60. Špetič A, Gennick J. Transact-SQL Cookbook: O’Reilly; 2002.


