Unintentional poisoning hospitalisations among young children in Victoria

Julie L Hoy, Lesley M Day, James Tibballs, Joan Ozanne-Smith

Abstract

Objectives—To describe the epidemiology of unintentional childhood poisoning hospitalisation in Victoria, Australia, in order to monitor trends and identify areas for research and prevention.

Methods—For children under 5 years, all Victorian public hospital admissions, July 1987 to June 1995, due to unintentional poisoning by drugs, medicines, and other substances were analysed. Similar cases were also extracted from the database of the Royal Children’s Hospital intensive care unit, Melbourne for the years 1979–91. Log linear regression modelling was used for trend analyses.

Results—The annual average childhood unintentional poisoning rate was 210.7 per 100 000. Annual rates for males consistently exceeded those for females. The most common agents were those acting on the respiratory system and on smooth and skeletal muscles (muscle relaxants, cough and cold medicines, antihistamines), aromatic analgesics (paracetamol), and systemic agents (including antihistamines).

Further investigation is justified for cardiac agents, some respiratory agents, and asthma medications.

Conclusions—Childhood poisoning hospitalisation rates have not decreased in Victoria over recent years. A focused, agent specific approach, as well as a series of generic measures for the prevention of poisoning to children under 5 is advocated. The ongoing surveillance, collection and analysis of data, in addition to research on specific poisoning agents are essential components of any prevention strategy.

(Keywords: unintentional poisoning; hospitalisation; intensive care unit; epidemiology)

Significant gains were made in childhood poisoning prevention in Australia and other countries mainly due to the introduction of child resistant packaging. However, unintentional poisoning of children less than 5 years of age remains the second leading cause of hospital admission for childhood injury in the four Australian states and territories where 86% of the population resides. Consequently, it has been identified as a priority in the national health goals and targets for injury prevention.

A consistent observation in the literature is that childhood poisoning is a heterogeneous issue with many individual agents contributing to the overall problem. This suggests that further gains in the reduction of poisoning will require focus on a number of individual poisoning agents, in addition to the consideration of further generic measures. One aspect of childhood poisoning which has received little attention is the way in which children gain access to poisoning agents.

The Childhood Poisoning Research and Prevention Project was designed to address these issues by (1) describing the dimensions of childhood poisoning among children under 5 years of age, (2) identifying poisoning agents for further study on the basis of frequency, severity, and potential for prevention, (3) identifying the mechanisms of access for specific agents, and (4) identifying potential prevention strategies.

Method

This study was designed to provide descriptive data and trend analyses for all children under 5 years admitted to public hospitals as a result of unintentional poisoning for a period of eight years in Victoria, Australia.

Two databases were used, the first holding information on all hospital admissions, and the second on admissions to the intensive care unit of the largest paediatric hospital.

Monash University Accident Research Centre holds a subset of the Victorian Inpatient Minimum Database (VIMD) records selected by external cause of injury codes (E codes) from the International Classification of Diseases, ninth revision, clinical modification (ICD9-CM).

We extracted cases of children less than 5 years of age admitted to public hospitals by the ICD9-CM E code relating to unintentional poisoning by drugs, medicinal substances, biologicals, other solid and liquid substances, gases, and vapours. Data were extracted from July 1987 to June 1995. Readmissions to the same public hospital were excluded. Transfers between public hospitals could not be excluded, but previous estimates of these occurrences were shown to be insignificant. Variables included age, sex, agent of poisoning, length of hospital stay, and place where poisoning occurred.

The intensive care unit of the Royal Children’s Hospital, Melbourne, holds a database of admissions from which cases of unintentional poisoning among children less than 5 years of age were identified for the years 1979–91. Additional information was ex-
Table 2 Trend in and distribution of childhood (<5 years age) poisoning hospitalisation by external cause of injury code (E code), Victoria, 1987–88 to 1994–95

<table>
<thead>
<tr>
<th>E code</th>
<th>Description of agents covered by E code</th>
<th>Frequency</th>
<th>%</th>
<th>β (slope)</th>
<th>95% CI for β</th>
</tr>
</thead>
<tbody>
<tr>
<td>858.6</td>
<td>Respiratory system, smooth and skeletal muscle agents</td>
<td>496</td>
<td>9.3</td>
<td>0.05</td>
<td>0.01 to 0.1</td>
</tr>
<tr>
<td>850.4</td>
<td>Aromatic analgesics</td>
<td>482</td>
<td>9.1</td>
<td>0.15</td>
<td>0.11 to 0.19</td>
</tr>
<tr>
<td>858.1</td>
<td>Systemic agents</td>
<td>327</td>
<td>6.1</td>
<td>0.07</td>
<td>0.000 to 0.14</td>
</tr>
<tr>
<td>858.7</td>
<td>Skin and mucous membrane, ear, nose, throat and dental drugs</td>
<td>280</td>
<td>5.3</td>
<td>-0.09</td>
<td>-0.15 to -0.02</td>
</tr>
<tr>
<td>858.3</td>
<td>Cardiovascular system agents</td>
<td>274</td>
<td>5.2</td>
<td>0.02</td>
<td>0.06 to 0.09</td>
</tr>
<tr>
<td>855.2</td>
<td>Benzodiazepine based tranquillisers</td>
<td>259</td>
<td>4.9</td>
<td>-0.04</td>
<td>-0.14 to 0.05</td>
</tr>
<tr>
<td>855.5</td>
<td>Sympathomimetics (adrenergics)</td>
<td>209</td>
<td>3.9</td>
<td>0.03</td>
<td>-0.05 to 0.1</td>
</tr>
<tr>
<td>863.4</td>
<td>Unspecified insecticides</td>
<td>178</td>
<td>3.3</td>
<td>0.06</td>
<td>-0.01 to 0.12</td>
</tr>
<tr>
<td>855.0</td>
<td>Anticonvulsant and antiparkinsonism drugs</td>
<td>169</td>
<td>3.2</td>
<td>-0.07</td>
<td>-0.13 to -0.01</td>
</tr>
<tr>
<td>858.2</td>
<td>Agents affecting blood constituents</td>
<td>137</td>
<td>2.6</td>
<td>-0.04</td>
<td>-0.11 to 0.02</td>
</tr>
</tbody>
</table>

Poisoning agents
Seventy three per cent of poisoning hospitalisations were due to ingestions of drugs, medications, or biological substances. The remainder was mostly due to domestic chemicals.

Four digit E codes are used to describe the 10 agent groups most often involved in poisoning (table 2). Specific drugs cannot be identified from the VIMD, with the exception of paracetamol, the only drug currently available in Australia that is classified as E850.5. Other drugs that could be classified into this group have not been marketed in Australia for some time. Of the 10 major agent groups resulting in hospitalisation, evidence of an upward trend was found for three and a downward trend for two groups (table 2). Also of concern was the upward trend found for rodenticides (β=0.11, CI 0.03 to 0.19) and antidepressants (β=0.08 CI 0.02 to 0.15).

Severity of poisoning
Ninety seven per cent of admissions due to poisoning were less than three days’ duration. Eight agent groups resulting in length of stay greater than two days caused 44% of these longer admissions (table 3). No single E code group accounted for more than 9% of these longer stays.

Place of poisoning
In 61.4% of cases, poisoning occurred within the home. For a further 36.3% of cases the location was unknown.

INTENSIVE CARE ADMISSIONS, ROYAL CHILDREN’S HOSPITAL
Rates and frequencies
A total of 250 admissions to the intensive care unit was recorded between 1979–91, an annual average of 19. Poisoning accounted for between 1.1 and 3.3% of all intensive care unit admissions. The only fatality was a 2 year old child who died after ingesting carbamazepine.

Age and gender
Admission frequency to the intensive care unit was greater for boys (53.6%) than girls (45.6%). As with hospital admissions, the peak frequencies were at ages 1 and 2 years (34%, 32%).

Poisoning agents
Sixty per cent of intensive care unit admissions were a result of drug ingestions. Eight agent groups (based on four digit E codes) were...
Table 3 Main poisoning agents associated with length of hospital stay greater than two days, childhood poisoning (<5 years age), Royal Children’s Hospital, Victoria, 1979–91

<table>
<thead>
<tr>
<th>E code</th>
<th>Agent</th>
<th>Frequency</th>
<th>% of all longer* admissions</th>
</tr>
</thead>
<tbody>
<tr>
<td>853.2</td>
<td>Benzodiazepine based tranquillisers</td>
<td>14</td>
<td>8.7</td>
</tr>
<tr>
<td>855.0</td>
<td>Anticonvulsant, antiparkinsonism drugs</td>
<td>12</td>
<td>7.5</td>
</tr>
<tr>
<td>850.4</td>
<td>Aromatic analgesics</td>
<td>10</td>
<td>6.2</td>
</tr>
<tr>
<td>853.0</td>
<td>Phenothiazine based tranquillisers</td>
<td>10</td>
<td>6.2</td>
</tr>
<tr>
<td>853.1</td>
<td>Butyrophenone based tranquillisers</td>
<td>7</td>
<td>4.3</td>
</tr>
<tr>
<td>854.0</td>
<td>Antidepressants</td>
<td>6</td>
<td>3.7</td>
</tr>
<tr>
<td>858.3</td>
<td>Cardiovascular system agents</td>
<td>6</td>
<td>3.7</td>
</tr>
<tr>
<td>858.7</td>
<td>Skin and mucus membrane, ear, nose, throat and dental drugs</td>
<td>6</td>
<td>3.7</td>
</tr>
</tbody>
</table>

*Length of stay >2 days.

Table 4 Distribution of childhood (<5 years age) poisoning admissions to intensive care unit by external cause injury code (E code), Royal Children’s Hospital, Victoria, 1979–91

| E code | Agents | Frequency | %
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>858.6</td>
<td>Respiratory system, smooth and skeletal muscle agents</td>
<td>27</td>
<td>10.8</td>
</tr>
<tr>
<td>854.0</td>
<td>Antidepressants</td>
<td>24</td>
<td>9.6</td>
</tr>
<tr>
<td>855.0</td>
<td>Anticonvulsants</td>
<td>22</td>
<td>8.8</td>
</tr>
<tr>
<td>858.3</td>
<td>Cardiovascular system agents</td>
<td>19</td>
<td>7.6</td>
</tr>
<tr>
<td>858.4</td>
<td>Gastrointestinal system agents</td>
<td>10</td>
<td>4.0</td>
</tr>
<tr>
<td>853.0</td>
<td>Phenothiazine based tranquillisers</td>
<td>7</td>
<td>2.8</td>
</tr>
<tr>
<td>852.8</td>
<td>Specified sedatives, hypnotics</td>
<td>5</td>
<td>2.0</td>
</tr>
<tr>
<td>853.2</td>
<td>Benzodiazepine based tranquillisers</td>
<td>4</td>
<td>1.6</td>
</tr>
</tbody>
</table>

Access and place of poisoning
Poisoning occurred in the child’s own home in 87% of incidents, and in 64%, the patient had accessed the agent with no apparent help from any other person. Ten per cent were known to have climbed to get the agent.

Long term complications
Seven children experienced one or more long term complications. These were oesophageal strictures after ingesting corrosive substances (3), persistent drooling after ingesting disinfectant (1), visual impairment after ingesting quinine (1), epilepsy and spastic quadriplegia after poisoning by motor vehicle exhaust gas (1), and decreased motor coordination after ingesting calcium carbonate (1).

Discussion
his study was limited to hospital admissions which account for 17% of medically treated childhood poisonings in Victoria. Other sources of childhood poisoning data in Victoria include the Victorian Injury Surveillance System (VISS) and the Victorian Poisons Information Centre (VPIC). These data sources reflect generally less severe poisoning and relate to emergency department presentations at participating hospitals (VISS), and calls regarding suspected and actual ingestions (VPIC). Unfortunately, it is not currently possible to link these databases to provide a more complete overview of childhood poisoning. Analyses of these data sources for less severe cases have been conducted previously. For prevention purposes, a focus on hospitalisation data addresses the more severe end of the spectrum.

The current classification of injury due to poisoning by ICD9 has no firm guidelines. The potential for inconsistent classification exists. There is some potential for different interpretation of the E codes by the hospital medical records clerks coding for the VIMD. This is particularly applicable for poisoning as the clustering of drugs into functional groups in the E code framework creates a situation where the same drug or substance can feasibly be categorised into more than one E code. Accuracy studies of the VIMD indicate that the use of E codes for injury surveillance is feasible and reliable.

An additional limitation of broad E code categories is that the researcher is unable to identify specific drugs or substances within those categories. The development of a better coding system, or an extension to the ICD9 E codes, would enhance the ability to identify agents involved. When implemented, ICD10 will allow identification of an increased number of subclasses within the current broad E code categories, and will facilitate more consistent classification. However, identification of specific poisoning agents will still be limited.

Length of stay has limited use as a proxy for injury severity since it excludes those with adverse sequelae who are managed as outpatients, and includes cases of prolonged admission due to non-medical reasons. Other limitations of the VIMD have been documented...
The absence of private hospital data for the period of the study would result in minimal bias since an annual average of 11 children under 5 years are admitted to private hospitals in Victoria as a result of unintentional poisoning. This study confirms that unintentional poisoning in children under 5 years of age in Victoria conforms to the patterns typically found in Westernised countries. These features include the incidence peak of poisoning at 2 years of age, a generally higher incidence in males, most incidents occurring at home, similarity of agents, low mortality, and short length of stay. The broad range of poisoning agents identified here confirms what is reported in other studies, both in Australia and overseas. Of most concern are the drugs or drug groups that occur with high frequency, upward trends, result in longer duration of stay, or in intensive care unit admissions. It is these groups that should be the focus of intensive prevention strategies. However, to develop a focused approach to prevention, a means to identify individual agents within these groups is required.

We used intensive care data from this study, in addition to emergency department presentation data from some Victorian hospitals published elsewhere, to identify the agents from each of the groups where extra focus is required. Case series studies to identify the mechanisms of access have been conducted for paracetamol, eucalyptus oil, tricyclic antidepressants, benzodiazepine tranquillisers, and rodenticides. Similar studies are yet to be conducted for cardiovascular drugs, asthma medications, antispasmodics, and cough and cold preparations. The decreasing trend for anticonvulsants or antiparkinsonism drugs is of interest. Intensive care unit data identified all admissions in this group as carbamazepine, an anticonvulsant. A legislative requirement to have child resistant closures and storage for this agent is of interest. Intensive care unit data identified all admissions in this group as carbamazepine, an anticonvulsant. A legislative requirement to have child resistant closures and storage for this agent is of interest. Intensive care unit data identified all admissions in this group as carbamazepine, an anticonvulsant. A legislative requirement to have child resistant closures and storage for this agent is of interest. Intensive care unit data identified all admissions in this group as carbamazepine, an anticonvulsant. A legislative requirement to have child resistant closures and storage for this agent is of interest. Intensive care unit data identified all admissions in this group as carbamazepine, an anticonvulsant. A legislative requirement to have child resistant closures and storage for this agent is of interest. Intensive care unit data identified all admissions in this group as carbamazepine, an anticonvulsant. A legislative requirement to have child resistant closures and storage for this agent is of interest.

The use of measures to limit absorption were often applied beyond their limits of efficacy. These treatments may, therefore, represent unjustified levels of invasive treatment, increase morbidity, and prolong hospital stay.

**Implications for prevention**

The high number of poisoning incidents occurring within the home offers potential for prevention strategies. Efforts need to be aimed at both 1 and 2 year olds, rather than just 2 year olds, the commonly accepted peak age of poisoning.

There are a number of drugs without mandatory requirement for child resistant packaging that are significant in childhood poisoning, including benzodiazepines and other tranquillisers, some cardiac drugs, some asthma medications, and eucalyptus based inhalation solutions. Other drugs, covered by the legislation, still cause significant, and in some cases serious, morbidity. These include paracetamol, tricyclic antidepressants, anticonvulsants, pure eucalyptus oil, and some cardiac drugs. The correct use of child resistant closures and storage of medications in the home require further investigation. There may also be a need for improving the efficacy of current child resistant packaging.

We have identified a number of poisoning agents for which further investigation of access by children is justified but has not yet been conducted.

Previous studies have suggested that poisoning prevention is optimal when both safe storage and child resistant packaging are used. The development and implementation of safe benchtop storage, as suggested elsewhere, may lead to significant decreases in poisoning. Design of such storage would need to accommodate both bottle and blister/strip packaging.

Funding was provided to Monash University Accident Research Centre (MUARC) by the Victorian Health Promotion Foundation and the Victorian Department of Human Services; through the Public Health Training Scheme. Dr Malcolm Dobbin and Mr Barry Parsons provided valuable advice. Julie Valeri, Christina Leong, Voula Stobabas, Barb Fox, and Dr Holger Hagen supplied a high standard of technical assistance with data retrieval, coding, and analysis. Dr Frank Shann gave approval to use data from the Royal Children’s Hospital intensive care unit. This project was carried out at MUARC.

A young woman was burned after striking the top of an egg she had boiled in water in a microwave oven. To avoid this type of injury, manufacturers recommend piercing the shell first; ideally the yolk should be pierced as well. Most manuals warn that eggs should not be cooked with an intact shell in the instruction pages, but fail to reiterate this information in the recipe pages (Burns 1998;24:585–6).

Seventy three cases, including 28 deaths, of injuries resulting from television sets toppling onto children (principally aged 12–24 months) were reported to the US Consumer Product Safety Commission, 1990–97. As television sets become larger and stands smaller, manufacturers need to pay attention to securing this top heavy arrangement of equipment. Many sets toppled when children climbed onto furniture to manipulate the controls. A 36 inch 78 kg television set falling one metre onto a child is roughly equivalent to a 1 year old child weighing 10 kg falling from a 10 storey height (Pediatrics 1998;102:e32).

Equipment fitted with sharp rotating blades needs to be well designed to prevent injuries to foolhardy operators. Sixty two people aged 11–77 years sustained hand injuries from snowblowers (44 complete finger amputations and 42 partial amputations), most incidents occurring when the operators placed their hands into the exit chutes and contacted the blades while clearing the chutes, the majority of which were not guarded. Voluntary guidelines have been inadequate in the US to enforce safety requirements such as labelling, design, durability of shields and guards, and an automatic stopping switch for these products (American Journal of Orthopedics 1997;26:863–7).

Are electronic security devices effective in preventing child abductions or to keep children from gaining access to roads and swimming pools? Respondents to a survey in Fairfax, Virginia, believed that responsible parents would use security devices, but were not willing to pay much for them (Child: Care, Health and Development 1997;23:415–21).
Unintentional poisoning hospitalisations among young children in Victoria

Julie L Hoy, Lesley M Day, James Tibballs and Joan Ozanne-Smith

*Inj Prev* 1999 5: 31-35
do: 10.1136/ip.5.1.31

Updated information and services can be found at:
http://injuryprevention.bmj.com/content/5/1/31

**References**

This article cites 22 articles, 1 of which you can access for free at:
http://injuryprevention.bmj.com/content/5/1/31#BIBL

**Email alerting service**

Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

**Topic Collections**

Articles on similar topics can be found in the following collections

Poisoning/Ingestion (141)

**Notes**

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/