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Sodium azide poisoning: a narrative review

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**ABSTRACT**

**Context:** Sodium azide is a highly toxic chemical. Its production has increased dramatically over the last 30 years due to its widespread use in vehicular airbags, and it is available for purchase online. Thus, accidental exposure to azide or use as a homicidal or suicidal agent could be on the rise, and secondary exposure to medical personnel can occur. No antidote exists for azide poisoning. We conducted a systematic review of azide poisoning to assess recent poisoning reports, exposure scenarios, clinical presentations, and treatment strategies.

**Methods:** We searched both medical and newspaper databases to review the literature between 01/01/2000 and 12/31/2020, pairing the controlled vocabulary and keyword terms “sodium azide” or “hydrazoic acid” with terms relating to exposures and outcomes, such as “ingestion,” “inhalation,” “exposure,” “poisoning,” and “death.” We included all peer-reviewed papers and news articles describing human azide poisoning cases from English and non-English publications that could be identified using English keywords. Data abstracted included the number, age, and gender of cases, mode of exposure, exposure setting, azide dose and route of exposure, symptoms, outcome, and treatment modalities.

**Results:** We identified 663 peer-reviewed papers and 303 newspaper articles. After removing duplicated and non-qualifying sources, 54 publications were reviewed describing 156 cases, yielding an average of 7.8 reported azide poisoning cases per year. This rate is three times higher than in a previous review covering the period of 1927 to 1999. Poisoning occurred most commonly in laboratory workers, during secondary exposure of medical personnel, or from a ripped airbag. Hypotension occurred commonly, in some cases requiring vasoppressors and one patient received an intra-aortic balloon pump. Gastric lavage and/or activated charcoal were used for oral azide ingestion, and sodium nitrite, sodium thiosulfate, and/or hydroxocobalamin were used in severely poisoned patients.

**Conclusions:** Recent increases in azide poisoning reports may stem from greater commercial use and availability. Treatment of systemic poisoning may require aggressive hemodynamic support due to profound hypotension. Based on mechanistic considerations, hydroxocobalamin is a rational choice for treating azide poisoning.

**Keywords:** Sodium azide (NaN\textsubscript{3}); hydrazoic acid (HN\textsubscript{3}); poisoning; treatment; review

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**Introduction**

Sodium azide (NaN\textsubscript{3}) and its conjugate acid, hydrazoic acid (HN\textsubscript{3}), are toxic compounds. We refer to both agents generically as “azide.” Fortunately, azide poisoning is relatively rare, but this means most medical personnel may have not encountered a case and possess limited knowledge of azide poisoning. This may prove detrimental in a mass casualty event, for example, an industrial accident or terrorist attack. The latter is possible, because sodium azide is easily available through online retailers and NaN\textsubscript{3} is listed as a potential weapon of mass destruction [1,2]. Indeed, azide has been used in several planned and/or executed terrorist attacks [3–7].

Azide has been most notably used as a propellant in vehicular airbags and airplane safety chutes. Following an automobile crash, an igniter generates high temperatures that rapidly decompose NaN\textsubscript{3} into sodium metal and nitrogen gas. Due mainly to its use in airbags, NaN\textsubscript{3} production surged beginning in 1990, and at least 1,000 tons are produced annually [8,9]. The fate of azide pellets in old airbags is generally unknown, posing a potential environmental problem [10–12]. Azide is also used in chemical laboratories to facilitate synthetic reactions, and in biomedical laboratories to inhibit microbial growth.

Within a year of its discovery, azide was shown to be toxic to plants and animals [13]. The mouse LD\textsubscript{50} is 19 mg/kg by intravenous injection, and the human lethal dose is estimated to be \(\geq 700\) mg total or \(\sim 10\) mg/kg [14,15]. Humans can be exposed to azide through three major routes: ingestion, transdermal or transmucosal absorption, or inhalation of hydrazoic acid vapors or sodium azide dust particles. At low doses, azide causes dizziness, nausea, vomiting, and restlessness. At high doses, it causes seizures, hypotension,
metabolic acidosis, coma, and respiratory failure. Symptoms occur within minutes of exposure.

Azide has several mechanisms of toxicity. At the cellular level, it inhibits mitochondrial cytochrome C oxidase and catalase [16,17]. The former enzyme is part of complex IV in the mitochondrial electron transport chain and the latter enzyme detoxifies hydrogen peroxide to water and oxygen. Thus, azide can reduce ATP synthesis and cause oxidative stress, the latter due to mitochondrial electron leakage and reduced catabolism of reactive oxygen species. Cyanide also inhibits cytochrome C oxidase, and azide and cyanide are especially toxic to cells with high respiratory rates, such as neurons and cardiomyocytes. At the organismal level, azide is a potent vasodilator and inhibits platelet aggregation, likely via conversion to nitric oxide. Azide generates nitric oxide in vitro in erythrocytes, platelets, and isolated blood vessels, and recently, nitrosyl-hemoglobin was found in the blood of mice that had received azide [18]. Cytochrome C oxidase inhibition and nitric oxide generation likely underlie the hypotension, myocardial and respiratory failure, and metabolic acidosis that occur in azide poisoning.

A previous systematic review examined human azide poisonings between 1927 and 1999 [14]. Due to the marked increase in azide production over the last 30 years and the potential of new treatments, we thought it timely to perform an updated review of human azide poisoning, concentrating on contemporary exposure settings, clinical presentations, and treatment.

Methods

Search strategy

We conducted the study in accordance with the Preferred Reporting Item for Systematic Reviews and Meta-Analyses (PRISMA) guidelines. We searched all publications from January 1, 2000 to December 31, 2020 that described human cases of azide exposure and toxicity. We conducted separate searches of peer-reviewed papers and newspaper articles.

KH led the literature searches. For peer-reviewed papers, we searched PubMed (pubmed.gov), Embase (embase.com), and Academic Search Complete (ProQuest) using “sodium azide” and “hydrazoic acid” as controlled vocabulary and keyword terms, pairing them with terms relating to exposure, setting, or outcomes, such as “ingestion,” “inhalation,” “exposure,” “poisoning,” and “death.” For news articles, we searched Access World News and U.S. Major Dailies using a similar search strategy and keywords, without a controlled vocabulary. Full search strategies for all databases are in Appendix 1. Supplemental Google searches were conducted for both peer-reviewed papers and newspaper articles using the same search terms. All records were imported into EndNote and de-duplication followed the Wichor Bramer process [19].

Inclusion and exclusion criteria

All articles describing human azide exposure were included. Reports of physical trauma due to airbag deployment alone were excluded as were articles on azides other than sodium, since NaN₃ is the principal azide species produced and it is available for purchase online [2,20,21]. Finally, we excluded articles that were duplicated cases, had insufficient data, or irrelevant account, such as no discussion of human azide poisoning.

JT and SS evaluated the titles and abstracts, and full-text of the articles, with GRB arbitrating conflicts. To increase the international scope and relevance, non-English publications were included, when English language search terms led to their identification. If an English translation was unavailable, native language speakers translated appropriate sections of the papers.

Data collection

JT and SS abstracted the following data, when available: the number, age, and gender of cases, mode of exposure (accidental, suicidal, or homicidal), exposure setting (automobile accident, or industrial, laboratory, or secondary exposure of medical personnel), azide dose received and route of exposure (ingestion, inhalation, or dermal contact), clinical symptoms, outcome including hospitalization and survival of the victim(s), and treatment modalities. The level of evidence for each article was determined. If a case was reported in both a peer-reviewed paper and a news article, only the peer-reviewed paper is included.

Results

For peer-reviewed papers, the initial search identified 663 publications. Based on the Wichor Bramer process, 175 duplications were removed. Of the remaining 488 papers, we excluded 432 after screening the titles and abstracts and finding the same case published in multiple journals, or absence of reference to human exposure. We performed full-text review of the remaining 56 eligible papers, excluding an additional 19 papers based on grounds of duplication, irrelevant account, and/or insufficient data. Thus, 37 papers met our inclusion criteria and were reviewed (Figure 1). Among the 37 papers, a native Chinese speaker reviewed one Chinese paper, and a native Japanese speaker (SS) reviewed two Japanese papers.

For news articles, the initial search identified 303 articles. Applying the same screening methods as for the peer-reviewed papers, we removed 133 duplications, and then excluded 107 articles after screening the titles and abstracts. Of the remaining 63 articles eligible for full-text review, 46 articles were excluded, producing 17 news publications that met our inclusion criteria (Figure 2).

Cases are grouped based on the mode of azide exposure, accidental, intentional (suicidal or homicidal), secondary, or unknown, and sub-grouped based on exposure setting. We found a total of 156 unique cases: 106 cases in the 37 peer-reviewed papers (Table 1), and 50 cases in the 17 news articles (Table 2). Of the 37 peer-reviewed papers, 36 were case reports that met level VII evidence: opinions of authorities and/or reports of expert committees, and, one was an
observational study that met level IV evidence [22,23]. We did not perform quantitative bias analysis on these publications, because no quantitative data were available in either the peer-reviewed papers or the news articles that would allow us to correlate outcome to exposure location, route, or mode or treatment modalities. The median age of the patients was 32 with a range of 18 to 73 years old. The gender distribution was about 17% females and 31% males, with 52% unknown.

**Accidental exposure to azide**

In 77 cases, victims were accidentally exposed to NaN₃, either from airbag deployment after a vehicular accident, or in an industrial, laboratory, or medical setting. Three victims died.

**Airbag deployment: peer-reviewed papers**

In 13 patients, azide exposure occurred from airbag deployment (Table 1(A-1) [9,24–33]). The patients developed symptoms after either inhaling or having dermal or ocular contact with undecomposed NaN₃ released when an airbag ripped during a vehicular accident. Seven people had chemical burns and one person was diagnosed with contact dermatitis [24–27,31,33]. Five patients without prior history of respiratory disease developed pulmonary symptoms such as dyspnea or stridor, and two were diagnosed with chemical pneumonitis [9,28–30,32]. Three patients sustained ocular injury [9,31,32].

**Industrial exposure: peer-reviewed papers**

Three papers described 43 patients who were accidentally exposed to azide at work, presenting symptoms ranging...
Table 1. Review of 37 peer-reviewed papers yielded 106 cases.

### 1A-1. Accidental exposure due to airbag deployment

| Publication | Level of evidence | Number of cases | Gender | Age | Exposure route | Clinical features | Hospitalization | Treatment | Survival |
|-------------|-------------------|-----------------|--------|-----|----------------|-------------------|-----------------|-----------|----------|
| Corazza et al. [24] | VII | 1 | Male | 35 | Dermal contact | First-degree burns<br>Irritant contact dermatitis | Yes | Topical steroids | Yes |
| Foley and Helm [25] | VII | 1 | Female | 32 | Dermal contact | First-degree burns | Yes | Topical antibiotics | Yes |
| Wu et al. [26] | VII | 1 | Female | 29 | Dermal contact | Irritant contact dermatitis | Yes | Topical and oral steroids | Yes |
| Suhr and Kreusch [27] | VII | 1 | Female | 73 | Dermal contact | Second-degree burns | Yes | Topical antibiotics | Yes |
| Hambrook and Fink [28] | VII | 1 | Male | 47 | Inhalation | Asthma<br>Chest tightness<br>Cough | Yes | β-agonists<br>Inhaled steroids | Yes |
| Caudle et al. [29] | VII | 1 | Male | 18 | Inhalation | Chemical pneumonitis<br>Dyspnea<br>Hemoptysis | Yes | Oral steroids<br>Oxygen | Yes |
| Belhadj-Tahar et al. [32] | VII | 1 | Male | 37 | Inhalation and dermal contact | Burning eyes<br>Dyspnea | Yes | Ocular irrigation<br>Supplemental oxygen | Yes |
| Francis et al. [9] | VII | 1 | Male | 22 | Inhalation and dermal contact | Alkaline ocular injury<br>Erythematous supraglottis<br>Stridor<br>Tachypnea | Yes | Mechanical ventilation<br>Ocular irrigation<br>Stress | Yes |
| Govindarajan et al. [30] | VII | 1 | Male | 56 | Inhalation | Chemical pneumonitis | Yes | N/A | Yes |
| Sever et al. [31] | VII | 3 | Female/Male | 21/39/33 | Dermal contact | All three had second-degree burns<br>One patient had alkaline ocular injury | Yes | Standard burn care<br>Victim with ocular burn received additional ophthalmic care | Yes |

### 1A-2. Accidental exposure due to industrial work

| Publication | Level of evidence | Number of cases | Gender | Age | Exposure route | Clinical features | Hospitalization | Treatment | Survival |
|-------------|-------------------|-----------------|--------|-----|----------------|-------------------|-----------------|-----------|----------|
| Pham et al. [36] | VII | 1 | Male | 29 | Inhalation and dermal contact | Bradycardia<br>Hypotension<br>Metabolic acidosis<br>Third-degree burns | Yes | Intravenous fluids<br>Mechanical ventilation<br>Sodium bicarbonate<br>Vasopressors | No |
| Miljours and Braun [34] | IV | 41 | Unknown | Median age not available | Inhalation | Burning eyes<br>Dizziness<br>Headache<br>Palpitations | No | Unknown | Yes |

(continued)
Table 1. Continued.

| Publication          | Level of evidence | Number of cases | Gender | Age | Exposure route | Clinical features                        | Hospitalization | Treatment                          | Survival |
|----------------------|-------------------|-----------------|--------|-----|----------------|------------------------------------------|-----------------|-----------------------------------|----------|
| Fang et al. [35]     | VII               | 1               | Male   | 32  | Inhalation and dermal contact | Diplopia, Dizziness, Paresthesias, Reduced muscle strength | Yes             | Hyperbaric oxygen, Steroids, Traditional Chinese medicines, Vitamins B₁, B₁₂, and C | Yes      |
| Angelotti et al. [39]| VII               | 1               | Male   | 33  | Dermal contact (13 g)  | Burns, Hypotension, Lacerations           | Yes             | Amputation, Mechanical ventilation, Vasopressors, Wound care | Yes      |
| Watanabe et al. [47] | VII               | 1               | Female | 53  | Ingestion (1 g)        | Generalized seizures, Hypotension, Metabolic acidosis | Yes             | Hemodialysis, Gastric lavage with activated charcoal, Intra-aortic balloon pump, Intravenous steroids, Mechanical ventilation, Vasopressors | Yes      |
| Dermican et al. [46] | VII               | 1               | Female | 25  | Ingestion (0.1 g)      | Generalized seizures, Headache, Vomiting | Yes             | Gastric lavage                     | Yes      |
| Spadafora et al. [40]| VII               | 1               | Female | Unknown | Ingestion (>10 g)      | Bradycardia, Coma, Hypoxia, Metabolic acidosis | Yes             | Unknown                           | No       |
| Senda et al. [50]    | VII               | 1               | Female | 25  | Ingestion             | Acute respiratory distress syndrome, Arrhythmia, Coma, Cardiac arrest, Metabolic acidosis | Yes             | Gastric lavage, Intravenous steroids, Mechanical ventilation, Sodium bicarbonate | No       |
| Fuyuno and Cyranoski [51]| VII              | 1               | Male   | 42  | Ingestion             | N/A                                      | No (found dead) | N/A                               | No       |

(continued)
### 1B-1. Suicide in a laboratory setting

| Publication                  | Level of evidence | Number of cases | Gender | Age  | Exposure route | Clinical features                     | Hospitalization | Treatment                                 | Survival |
|------------------------------|-------------------|-----------------|--------|------|----------------|--------------------------------------|-----------------|------------------------------------------|----------|
| Łopiński et al. [52]         | VII               | 2               | Male   | 30   | Ingestion (>0.18 g) | Dizziness Metabolic acidosis Tachycardia Amythia Cardiac arrest Coma Metabolic acidosis | Yes             | Inhaled amyl nitrite Intravenous sodium nitrite Intravenous sodium thiosulfate | Yes      |
| Meatherall and Palatnick [53]| VII               | 1               | Male   | 59   | Ingestion | Cardiac arrest Coma Metabolic acidosis | Yes             | Exchange transfusion Inhaled amyl nitrite Intravenous sodium nitrite Sodium bicarbonate Mechanical ventilation Vasopressors | No       |
| French et al. [54]           | VII               | 1               | Male   | 28   | Ingestion (0.1 g) | Bradycardia Hypotension Metabolic acidosis | Yes             | Intravenous fluids Gastric lavage Sodium bicarbonate | Yes      |
| Kostek et al. [55]           | VII               | 1               | Female | 55   | Ingestion (0.6 g) | Metabolic acidosis | Yes             | Gastric lavage | Yes                                     |          |
| Le Blanc-Louvry et al. [56]  | VII               | 1               | Male   | 35   | Ingestion (6 g) | N/A                       | No              | N/A | N/A | No                                     |          |
| Barteka-Mino et al. [57]     | VII               | 1               | Female | 25   | Ingestion | Coma Hypotension Metabolic acidosis | Yes             | Hydroxocobalamin Vasopressors | Yes      |
| Downes et al. [58]           | VII               | 1               | Male   | 32   | Ingestion | Hypotension Metabolic acidosis | Yes             | Intravenous fluids Vasopressors | No       |
| Gao et al. [60]              | VII               | 1               | Male   | 23   | Ingestion (1.38 g) | Chest pain Hypotension Metabolic acidosis Nausea and vomiting | Yes             | Hemodialysis Intradavenous crystalloid Norepinephrine | Yes      |
| Muvalia et al. [59]          | VII               | 1               | Female | 19   | Ingestion (30 g) | Coma Hypotension Metabolic acidosis Nausea and vomiting | Yes             | Mechanical ventilation Vasopressors | No       |

### 1B-2. Suicide in a non-laboratory setting

| Publication                  | Level of evidence | Number of cases | Gender | Age  | Exposure route | Clinical features | Hospitalization | Treatment | Survival |
|------------------------------|-------------------|-----------------|--------|------|----------------|-------------------|-----------------|----------|----------|
| Wierowski et al. [61]        | VII               | 1               | Male   | 19   | Ingestion (~20 g) | N/A                       | No              | N/A | No       |
| Meatherall and Oleschuk [53] | VII               | 1               | Male   | 35   | Ingestion | N/A                       | No              | N/A | No       |
### Table 1. Continued.

#### 1B. Suicide in a non-laboratory setting

| Publication          | Level of evidence | Number of cases | Gender | Age | Exposure route | Clinical features | Hospitalization | Treatment | Survival |
|----------------------|-------------------|-----------------|--------|-----|----------------|-------------------|----------------|-----------|----------|
| Overtchouk et al. [63] | VII               | 1               | Female | 69  | Ingestion (15 g) | Myocardial dysfunction, Metabolic acidosis | Yes            | Gastric lavage, Intravenous fluids | Yes       |
| Rojek et al. [64]     | VII               | 1               | Female | 50  | Ingestion      | Bradycardia, Coma, Hypotension               | Yes            | Mechanical ventilation, Vasopressors | No        |
| Ciesla et al. [65]    | VII               | 1               | Male   | 24  | Ingestion      | Coma                                           | Yes            | Unknown                     | No        |
| Leonard et al. [2]    | VII               | 1               | Male   | 22  | Ingestion (40 g) | Arrhythmia, Cardiac arrest, Hypotension, Metabolic acidosis | Yes            | Mechanical ventilation         | No        |

#### 1C. Secondary exposure of emergency medical personnel

| Publication          | Level of evidence | Number of cases | Gender | Age (median) | Exposure route                          | Clinical features       | Hospitalization | Treatment                              | Survival |
|----------------------|-------------------|-----------------|--------|--------------|-----------------------------------------|-------------------------|----------------|----------------------------------------|----------|
| Hirose et al. [73]²  | VII               | 6               | Unknown| Unknown      | Inhalation                              | Burning eyes, Dizziness, Dyspnea, Headache | No             | Unknown                               | Yes      |
| Downes et al. [58]²* | VII               | 10              | 5 Males| 39           | Inhalation and dermal contact           | 1 case reported fatigue, 1 case reported stress | No             | Time off from work for the two cases | Yes      |

#### 1D. Unknown exposure

| Publication          | Level of evidence | Number of cases | Gender | Age | Exposure route | Clinical features | Hospitalization | Treatment                               | Survival |
|----------------------|-------------------|-----------------|--------|-----|----------------|-------------------|----------------|------------------------------------------|----------|
| Hirose et al. [73]²  | VII               | 7               | 1 Female| 22  | Ingestion      | Dizziness, Hypotension, Palpitations, Paresthesias, Syncope | Yes            | Gastric lavage, Inhaled amyl nitrite, Intravenous sodium nitrite, Intravenous sodium thiosulfate, Vasopressors | Yes      |
| Schwarz et al. [74]  | VII               | 5               | 3 Females| Median age not available | Ingestion | Arrhythmia, Headache, Hypotension | Yes            | Intravenous fluids, Oral antiemetics    | Yes      |

They are grouped based on mode of azide exposure (i.e., accidental, suicidal, secondary, and unknown, letters A through D, respectively) and sub-grouped based on exposure setting (numbers 1 through 4). Unless indicated, an entry is a primary patient.

²Downes et al. [58] reported a total of 11 patients. The primary victim committed suicide by azide ingestion and is described in Table 1(B-1). Ten medical personnel who treated the patient are described in Table 1(C).

*Although not in the abstract, we contacted the corresponding author who confirmed the patient ingested azide to commit suicide.

Ten patients were exposed to azide and hospitalized. Hirose worked at the hospital where seven of the patients were treated. Hence, the paper provided data on only those seven victims; they are summarized in Table 1(D) (information was not provided on whether poisoning was intentional or accidental). Six medical personnel who treated the seven azide-poisoned patients presented symptoms consistent with low-dose azide exposure, likely via inhaling HN₃ gas released when performing gastric lavage on the patients. They are described in Table 1(C).
Table 2. Review of 17 news articles yielded 50 cases.

2A-1. Accidental exposure due to industrial work

| Publication | Number of cases | Gender | Age | Exposure route | Clinical features | Hospitalization | Survival |
|-------------|-----------------|--------|-----|----------------|------------------|-----------------|----------|
| Perkins [37] | 1 Male | 45 | Dermal contact | Burned >15% of body | Yes | No |

2A-2. Accidental exposure in a laboratory

| Publication | Number of cases | Gender | Age | Exposure route | Clinical features | Hospitalization | Survival |
|-------------|-----------------|--------|-----|----------------|------------------|-----------------|----------|
| Green [41] | 1 Male | Unknown | Dermal Contact (65 g) | Burns | Yes | Yes |
| Author unknown [44] | 1 Male | Unknown | Ingestion | Unknown | Yes | Yes |
| Lillington [40] | 11 Unknown | Unknown | Inhalation | Unknown | Yes | Yes |
| Crabbe [42] | 1 Male | 27 | Dermal contact | Facial burns Glass embedded in chest and abdomen Minor lacerations | Yes | Yes |
| Kemsley [43] | 1 Male | Unknown | Dermal contact (200 g) | Second-degree burns Minor lacerations | Yes | Yes |
| Author unknown [45] | 1 Male | Unknown | Dermal contact | Minor lacerations | Yes | Yes |

2A-3. Accidental exposure in a medical facility

| Publication | Number of cases | Gender | Age | Exposure route | Clinical features | Hospitalization | Survival |
|-------------|-----------------|--------|-----|----------------|------------------|-----------------|----------|
| Author unknown [48] | 1 Male | 66 | Ingestion (1.5 g) | Rapid deterioration Vomiting | Yes | No |

2B. Suicide in a non-laboratory setting

| Publication | Number of cases | Gender | Age | Exposure route | Clinical features | Hospitalization | Survival |
|-------------|-----------------|--------|-----|----------------|------------------|-----------------|----------|
| Demare [67] | 1 Female | 32 | Ingestion | Unknown | No | Yes |
| Stout [68] | 1 Female | 25 | Ingestion | Unknown | Yes | No |
| Bender [66] | 1 Female | 71 | Ingestion | N/A | No | No |
| Singh [69] | 1 Male | 21 | Ingestion | Unknown | Yes | No |
| Hicks [70] | 1 Male | 27 | Unknown | N/A | No | No |

2C. Homicide

| Publication | Number of cases | Gender | Age | Exposure route | Clinical features | Hospitalization | Survival |
|-------------|-----------------|--------|-----|----------------|------------------|-----------------|----------|
| Author unknown [48] | 3 Unknown | Unknown | Ingestion (20 g) | Unknown | Yes | Yes |
| State of Arizona, Appellee, v. Wendi Elizabeth Adriano, Appellant [72] | 1 Male | 33 | Ingestion (possibly 21 g) | N/A | No | No |

2D. Secondary occupational exposure of medical personnel

| Publication | Number of cases | Gender | Age | Exposure route | Clinical features | Hospitalization | Survival |
|-------------|-----------------|--------|-----|----------------|------------------|-----------------|----------|
| Demare [67] | 5 Unknown | Unknown | Unknown | Minor respiratory symptoms | Yes | Yes |

(continued)
Table 2. Continued.

2D. Secondary occupational exposure of medical personnel

| Publication | Number of cases | Gender | Age | Exposure route | Clinical features | Hospitalization | Survival |
|-------------|----------------|--------|-----|----------------|------------------|-----------------|----------|
| Crabbe [42] | 1 Male         | 25     | Dermal contact | Facial and corneal burns | Yes | Yes |
| Stout [68]  | 6 Unknown Unknown Unknown Unknown Unknown Yes Yes |
| Hicks [70]  | 1 Unknown Unknown Unknown Unknown Unknown Yes Yes |

2E. Unknown exposure

| Publication | Number of cases | Gender | Age | Exposure route | Clinical features | Hospitalization | Survival |
|-------------|----------------|--------|-----|----------------|------------------|-----------------|----------|
| Author unknown [75] | 6 | 1 Male | Unknown Ingestion | Dizziness Syncope Tinnitus | Yes | Yes |
| Author unknown [76] | 4 | Unknown Unknown Ingestion | Dizziness Lightheadedness | Yes | Yes |

They are grouped based on mode of azide exposure (i.e., accidental, suicidal, homicidal, secondary, and unknown, letters A through E, respectively) and sub-grouped based on exposure setting (numbers 1 through 3). Unless indicated, an entry is a primary patient.

*Crabbe [42] reported two patients. The primary victim was exposed to azide in a laboratory explosion and is described Table 2(A-2). A firefighter was exposed to azide while responding to the primary victim and is described in Table 2(D).

*DeMare [67] reported six patients. The primary victim committed suicide by ingesting azide and is described Table 2(B). Five emergency response personnel who came into contact with the primary victim are described in Table 2(D).

*Stout [68] reported seven patients. The primary victim committed suicide by ingesting azide and is described Table 2(B). Six emergency response personnel who came into contact with the primary victim are described in Table 2(D).

*Hicks [70] reported two patients. The primary victim committed suicide with azide via an unknown route and is described Table 2(B). Among personnel who responded to the victim, one officer was hospitalized for observation and is described in Table 2(D).
from mild to severe, including one death (Table 1(A-2) [34,35,36]). One paper was a case-control survey of 41 workers who consistently inhaled more than the legal limit of 0.3 mg/m$^3$ of NaN$_3$ over a five-year period. The case subjects reported more events of burning eyes, dizziness, headache, and palpitations than control subjects [34]. In the second paper, a patient working in a poorly ventilated NaN$_3$ packing factory was hospitalized for diplopia, dizziness, paresthesias, and severely reduced muscle strength that immobilized him for three months [35]. He partially recovered muscle strength, but was left with moderate neurological dysfunction. The last paper described a patient who sustained third-degree burns when his forklift accidently struck a 50-gallon barrel containing NaN$_3$ waste, resulting in an explosion [36]. He subsequently developed hypotension and metabolic acidosis, and died, likely from both the azide exposure and the severe burns.

**Industrial exposure: news articles**

A worker died in an industrial explosion. He was cleaning out a filter drum used to produce NaN$_3$ when the drum exploded, causing burns to >15% of his body (Table 2(A-1) [37]). Less than five years earlier, another azide-induced explosion at the same plant injured four workers, one of whom died [38].

**Laboratory exposure: peer-reviewed paper**

A chemist was exposed to the equivalent of 13 g (185 mg/kg) of azide while performing an experiment using NaN$_3$ that led to an explosion (Table 1(A-3) [39]). He suffered thermal and chemical burns and lacerations. On hospitalization, he rapidly deteriorated with profound hypotension. He survived, but his left hand required amputation. Again, his clinical presentation was likely secondary to the combination of azide poisoning and severe burns.

**Laboratory exposure: news articles**

Six articles described 16 cases of accidental exposure among laboratory workers (Table 2(A-2) [40–45]). A clinical
immunology technician dropped a 200-mL bottle containing a weak solution of NaN₃ that generated volatile HN₃. Out of concern, 11 workers were hospitalized but subsequently returned to work without complications [40]. The other five cases involved chemists and were less benign. A graduate student did not follow proper safety procedures, and ingested an azide solution; he became ill, but survived [44]. In four cases, chemical reactions with NaN₃ resulted in explosions, leaving the operators with superficial burns and minor lacerations. Three explosions were caused by negligence using azide amounts that exceeded safety limits [41,43,45].

**Exposure in a medical facility: peer-reviewed papers**

Two papers described accidental azide ingestion in a medical facility (Table 1(A-4)). In one case, a dentist ingested ~10 mg NaN₃. At this low dose, she experienced generalized seizures, headache, and vomiting within minutes [46]. In the second case, a hospitalized patient drank a solution containing 1 g NaN₃ that was meant as a urine preservative [47]. She quickly developed generalized seizures, hypotension, and metabolic acidosis. Both patients survived, likely due to a relatively low azide dose in the first case and a rapid and aggressive medical response in the second case.

**Exposure in a medical facility: news article**

A patient died after swallowing an azide tablet thought to be a painkiller (Table 2(A-3) [48]). He started vomiting and rapidly deteriorated 30 min after ingesting the tablet, and died the next day. The source of the azide tablet was unknown.

**Intentional exposure to azide**

We found 24 cases of suicidal ingestion of NaN₃ (Tables 1(B-1), 1(B-2), and 2(B)). Thirteen cases occurred inside a laboratory (Table 1(B-1)). We also found four cases of homicidal exposure to NaN₃ (Table 2(C)). Of these 28 cases, 19 of the patients died.

**Suicide in a laboratory setting: peer-reviewed papers**

Of 13 people who intentionally ingested NaN₃, two were found dead, and the remaining 11 were hospitalized, of whom six later died, leaving five survivors (Table 1(B-1) [49–60]). Many of the patients were hypotensive with metabolic acidosis, consistent with high-dose azide exposure. When the amount of ingested azide could be identified, the highest survivable dose was 1.38 g (patient's weight was unknown) [60].

**Suicide in a non-laboratory setting: peer-reviewed papers**

Of six cases of suicidal ingestion of NaN₃, two of the patients were found dead (Table 1(B-2) [61–65]). The remaining four patients were hospitalized, and three of whom two were hypotensive and two had metabolic acidosis. The one surviving patient was particularly notable: a 69-year-old female who ingested ~15 g of NaN₃, about 20-fold more than the estimated lethal human dose, was not hypotensive but had reduced myocardial contraction with a left ventricular ejection fraction of 30% that returned to normal three weeks later [63].

**Suicide in a non-laboratory setting: news articles**

Five people who ingested NaN₃ in a non-laboratory setting all died, three were found dead, while the other two died after hospitalization (Table 2(B) [66–70]). In one case, NaN₃ was found in the victim's car, but it was unclear if he died due to ingesting azide or inhaling hydrazoic acid, which could have been created by mixing NaN₃ with water [70].

**Homicides: news articles**

Two articles described four homicidal cases (Table 2(C)). A chemist hoping to cause workplace havoc poisoned three of his colleagues’ coffee with ~20 g of NaN₃ [71]. Due to an unusual smell, the victims immediately spat out the coffee, but two of them became mildly ill and the third fell unconscious. In the fourth case, a woman bought NaN₃ to poison her husband to claim his life insurance policy [72]. She also stabbed and bludgeoned him. It was unclear how much azide contributed to his death, but it was found in his blood.

**Secondary occupational exposure of medical personnel**

We found 29 cases of medical personnel becoming secondarily poisoned while treating azide-poisoned victims. All of the personnel lived.

**Secondary exposure of medical personnel: peer-reviewed papers**

Ten medical workers were likely exposed to azide from a suicidal patient who was not decontaminated prior to hospitalization (Table 1(C) [58]). Three non-hospital staff who either provided prehospital care or transported the patient to the hospital were admitted to the emergency room for evaluation and were subsequently discharged, presumably without symptoms. But, of seven hospital personnel who were in direct care of the patient, two required time off. One person took a day off on supervisory advice to rest due to fatigue. The second person required several weeks off due to psychological stress from exposure to a toxic chemical. In a separate incident, six hospital personnel likely inhaled HN₃ gas liberated while performing gastric lavage on patients who had drank an azide-poisoned beverage at work (Table 1(C) [73]). They developed ocular irritation, dizziness, dyspnea, and headache, symptoms consistent with low-dose azide exposure, but promptly recovered.

**Secondary exposure of medical personnel: news articles**

Thirteen medical workers were secondarily exposed (Table 2(D) [42,67,68,70]). One case was particularly notable: a firefighter had to be treated for facial and eye burns after responding to an azide-related university laboratory explosion [42]. Although he was wearing a protective face mask, it
was postulated that azide powder contacted his face above the mask, where it reacted with sweat that then dripped down onto his eye.

### Unknown exposure

#### Peer-reviewed papers

Seven workers had to be hospitalized after becoming ill from drinking tea or coffee made from azide-contaminated water, and five people at a restaurant drank azide-tainted iced-tea (Table 1(D) [73,74]). Symptoms occurred within minutes, which included arrhythmias, headache, and hypotension. There were no fatalities.

#### News articles

A total of 10 people at two different medical schools drank from an azide-tainted coffee pot or coffee machine (Table 2(E) [75,76]). All of the victims survived, although they reported symptoms consistent with low-dose azide exposure.

### Treatment

No specific azide antidote currently exists. Therefore, treatments were largely supportive and in response to symptoms. Gastric lavage, with or without activated charcoal, was used in 13 patients, one of whom died [46,47,50,54,55,63,73]. The NaN₃ dose in these cases ranged from 0.1 g to 1.38 g, with the outlier being the one patient who survived 15 g [63].

Twenty-three patients presented with hypotension [2,36,39,47,53,54,57–60,64,73,74]. They received intravenous fluids and/or vasopressors, with some patients requiring large amounts of fluids, e.g., up to 32 L over 12 h [36]. The hypotension was presumably secondary to azide conversion to nitric oxide. Fifteen patients presented with metabolic acidosis, likely due to profound hypotension and anaerobic metabolism to compensate for mitochondrial inhibition [2,36,47,49,50,52–55,57–60,63]. In some cases, acidosis was severe enough that sodium bicarbonate was administered [36,50,53,54]. Two patients underwent hemodialysis, one of whom required an intra-aortic balloon pump to maintain blood pressure; both patients survived [47,60]. One patient who underwent exchange transfusion died [53].

Due to mechanistic similarities between azide and cyanide, many patients were given cyanide antidote(s). Ten patients were treated with sodium thiosulfate, amyl nitrite, and/or sodium nitrite, of whom two died [52,53,73]. One patient treated with hydroxocobalamin survived [57].

Persons who had respiratory distress due to inhaling hydrazoic acid or sodium azide dust were treated with corticosteroids and β-agonists [28,29]. People who had azide contact to skin or eyes received general treatment for a chemical exposure, which included steroids and antibiotics [9,24–27,31–33].

### Discussion

We found 156 cases in 37 peer-reviewed papers and 17 news articles over the 20-year period between 2000 and 2020, providing a mean of 7.8 cases per year. For comparison, Chang and Lamm found 185 cases over a 72-year period, or an average of 2.6 cases per year. Our rate could be higher due to better reporting and/or more cases, the latter presumably because of greater azide production in the last 30 years and/or the ease of purchasing sodium azide online.

In the Chang and Lamm review, three people were exposed to sodium azide in vehicular accidents. We found 13 people exposed to sodium azide during airbag deployment. The increase in reported cases is likely because all automobiles sold in the U.S. after September 1, 1998 had to have airbags around the front seat. Five persons developed respiratory symptoms after airbag deployment. These five cases are consistent with reports that both sodium azide and hydrazoic acid can cause respiratory distress in animals [15]. Although azide was the likely cause of the respiratory problems, airbags contain other substances such as talcum powder, sodium carbonate, and sodium hydroxide that could cause pulmonary injury. Seven patients exposed to azide during airbag deployment had evidence of dermal burns and three of the patients had evidence of ocular injury. Even though many airbag manufacturers have stopped using azide, it still persists in airbags in many cars; thus, physicians need to be aware of potential respiratory, cutaneous, and ocular injuries due to azide exposure from airbag deployment.

Accidental industrial exposure to azide is almost inevitable, since numerous factories either produce sodium azide, or use it in manufacturing processes. Two of the 44 cases of industrial exposure died from azide-related explosions. Azide is shock-sensitive, and vibrations occurring when azide-containing drums were moved may have led to the explosions. Several patients developed neurological symptoms, and neurotoxicity secondary to azide exposure has occurred in other human cases and in animal models [77,78]. Care should be taken to minimize azide inhalation through use of face coverings and working in well-ventilated spaces, as well as minimizing mechanical friction that could trigger azide explosions.

Azide-related laboratory accidents occurred in 17 cases. Several cases seem to have been the result of safety negligence. While no death occurred, several victims sustained serious injury, highlighting the need to be vigilant when working with azide.

Three cases of accidental azide ingestion occurred in a medical facility; one of whom died. Sodium azide is an ordinary white powder that when dissolved in water, becomes a clear, odorless solution. It is thus possible that sodium azide could be confused as medicine and that azide solutions could be mistaken for drinking water. While accidental ingestion of azide in a medical facility is rare, azide is commonly used as a preservative, and its use must be monitored closely.

Chang and Lamm reported azide was used as a suicidal agent by 13 people, all of whom died. We found 24 suicidal cases, of whom 18 died. Death due to azide is dependent on
dose and the time between exposure and treatment onset. The latter parameter was impossible to ascertain in many cases. Thirteen of the 24 cases were scientific workers who ingested azide in their laboratories and likely had direct access to the chemical. This, in conjunction with 17 cases of accidental laboratory exposure to azide, indicates laboratory workers constitute a high-risk group with regards to azide poisoning.

We found four attempted homicides using azide. Three cases were laboratory workers who did not sustain serious injury. In another incident, the victim was poisoned by his wife who purchased azide online. These cases highlight the danger of azide in the hands of those with malicious intentions.

Our findings reveal medical personnel are at high risk for azide exposure, 29 people became secondarily poisoned while caring for azide-poisoned patients. Although the risk of serious injury is low, several providers reported dizziness, dyspnea, headache, ocular irritation, and fatigue, symptoms compatible with low dose azide exposure. Medical response personnel need to be aware of the consequences of azide poisoning, both in caring for azide-poisoned victims and to protect themselves from secondary exposure.

Twenty-two people were poisoned after drinking from communal beverage sources. While no long-term sequelae occurred due to low exposure dose and rapid care, it was unclear if the poisoning events were accidental or intentional. The latter highlights the potential threat of large-scale azide exposure due to a terrorist attack, especially considering azide’s online availability and ease of synthesis.

Treatment varied, depending on the scenario. Gastric lavage with or without activated charcoal may not be effective, because it is not known how long sodium azide, which is water soluble and generally ingested as a solution, would remain in the stomach, and it is unclear if azide actually binds to charcoal [79,80].

Since azide can cause profound hypotension, supportive care with vasopressors and large amounts of intravenous fluids is frequently needed. Two patients underwent hemodialysis, one of whom also received an intra-aortic balloon pump to maintain blood pressure [47,60]. The former treatment may be useful, because NaN₃ is a small molecule (65 kDa).

Although both azide and cyanide inhibit cytochrome C oxidase, it seems unlikely that sodium thiosulfate, amyl nitrite, or sodium nitrite would be useful against azide poisoning. Sodium thiosulfate detoxifies cyanide by converting it to thiocyanate; it seems unlikely that sodium thiosulfate would react with azide. Amyl nitrite and sodium nitrite generate methemoglobin, which scavenges cyanide, but methemoglobin binds azide only weakly (log $K_{observed} = 5.427$ at pH 7.02) [81]. Moreover, nitrites could be detrimental, because they generate nitric oxide, which could exacerbate azide-induced hypotension [82,83].

Hydroxocobalamin appears to be a reasonable treatment for azide poisoning, because cobalamin binds nitric oxide, and high doses of cobalamin raise blood pressure in control subjects, likely via reducing plasma nitric oxide concentrations [84,85]. Thus, cobalamin could potentially raise the blood pressure in azide-poisoned hypotensive patients.

**Limitations**

While two-thirds of the sources we included were peer-reviewed, all but one paper were case reports, which are prone to publication bias and therefore limited generalizability. The single case-control study relied on a work-medical history questionnaire, which is susceptible to recall bias.

As in other reviews of clinical cases, we were limited by whether someone chose to publish a case and whether a diagnosis was made appropriately. The former is exemplified in Hirose et al. where data on only seven of 10 azide-poisoned victims were provided [73]. The latter may be particularly problematic for azide poisoning, since it can be mistaken for cyanide poisoning due to mechanistic and symptomatic similarities. Taken together, the actual incidence of azide poisoning is likely to be greater than that reported. The cases reported in newspaper articles generally did not describe treatment strategies, limiting our ability to tie treatment to outcome.

**Conclusions**

Multiple routes of azide exposure exist, resulting in a variety of symptoms and range of disease severity. Treatment depends largely on the type, site, and degree of injury, and success can be ambiguous. Preventing or minimizing exposure, especially among high-risk persons, is the best strategy.

Laboratory workers should perform risk assessments to identify potential areas of concern prior to working with azide, and consideration should be given to limiting access to azide to prevent accidental or intentional ingestion. Concentrated azide solutions should not be exposed to shock or friction, or come into contact with heavy metals due to the possibility of explosion.

For medical personnel, the most likely exposure routes are inhalation of hydrazoic acid expired by (or lavaged from) a patient, dermal contact from sodium azide dust left on patients or their clothing, or eye exposure to hydrazoic acid or sodium azide dust. First responders should therefore wear safety goggles and gloves, and use CPR barrier devices where azide poisoning is possible. Moreover, any person suspected of azide poisoning should be decontaminated prior to hospitalization according to standard Hazmat protocols.

For severe systemic exposures to azide, hemodynamic support is clearly important, with some patients requiring aggressive therapy. Quantitative information on appropriate drugs is lacking, but, hydroxocobalamin is a rational choice based on its mechanism of action and relatively good safety profile.

**Author contributions**

JT studied under GRB in conceiving the study. JT coordinated the study and GRB supervised the research. KH conducted the literature search and generated the library. JT and SS conducted the literature review and analyzed the articles. KH, RBP, and BAG provided organizational advice. JT and GRB wrote the manuscript with inputs from co-authors. All authors reviewed and approved the final manuscript.
Disclosure statement

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