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Abstract
Naloxone is a well-established essential medicine for the treatment of life-threatening heroin/opioid overdose in emergency medicine. Over two decades, the concept of ‘take-home naloxone’ has evolved, comprising pre-provision of an emergency supply to laypersons likely to witness an opioid overdose (e.g. peers and family members of people who use opioids as well as non-medical personnel), with the recommendation to administer the naloxone to the overdose victim as interim care while awaiting an ambulance. There is an urgent need for more widespread naloxone access considering the growing problem of opioid overdose deaths, accounting for more than 100,000 deaths worldwide annually. Rises in mortality are particularly sharp in North America, where the ongoing prescription opioid problem is now overlaid with a rapid growth in overdose deaths from heroin and illicit fentanyl. Using opioids alone is dangerous, and the mortality risk is clustered at certain times and contexts, including on prison release and discharge from hospital and residential care. The provision of take-home naloxone has required the introduction of new legislation and new naloxone products. These include pre-filled syringes and auto-injectors and, crucially, new concentrated nasal sprays (four formulations recently approved in different countries) with speed of onset comparable to intramuscular naloxone and relative bioavailability of approximately 40–50%. Choosing the right naloxone dose in the fentanyl era is a matter of ongoing debate, but the safety margin of the approved nasal sprays is superior to improvised nasal kits. New legislation in different countries permits over-the-counter sales or other prescription-free methods of provision. However, access remains uneven with take-home naloxone still not provided in many countries and communities, and with ongoing barriers contributing to implementation inertia. Take-home naloxone is an important component of the response to the global overdose problem, but greater commitment to implementation will be essential, alongside improved affordable products, if a greater impact is to be achieved.

Key Points
- Take-home naloxone is an effective public health intervention to prevent deaths and organ damage from opioid overdose.
- Four naloxone nasal spray products that have been developed for layperson use are now approved, all with approximately 40–50% bioavailability relative to parenteral references. They are increasingly available in clinical practice in a growing number of countries.
- Ongoing implementation challenges include naloxone cost as well as politico-social (e.g. stigma) and legal barriers (e.g. prescription status), although prescription-free distribution is now permitted in several countries (e.g. Australia, Canada, Italy, the UK).
1 Introduction

With an increasing global problem of opioid use and dependence, the mortality rate from opioid overdose continues to rise, and there are more than 100,000 deaths globally per annum [1]. Heroin is the most common drug involved in opioid overdose in much of the world, although prescription drugs and illicitly manufactured fentanyl (IMF) and analogues are increasingly implicated, particularly in North America. In the USA, 20,000 deaths from prescription opioids and 13,000 deaths from heroin were registered in 2015 alone [2].

These deaths occur primarily in the community and, while sometimes the patient is alone, he/she often in the presence of others (especially with heroin overdose) [3–6]. As such, these deaths are potentially preventable with timely detection and administration of naloxone, along with wider resuscitation measures, by non-medical members of the general public.

Naloxone is a remarkable antidote that is opioid specific and actively displaces heroin and other opioids from the mu-opioid receptor (MOR). It is a long-established medicine that is essential in the hospital emergency department and the ambulance medication kit. Naloxone administration reverses heroin/opioid overdose within minutes with rapid re-establishment of independent breathing and return of consciousness (and, in individuals who are dependent on opioids, risk of precipitation of an opioid withdrawal syndrome).

‘Take-home naloxone’ (THN) for the reversal of heroin/opioid overdose involves the pre-provision of an emergency naloxone kit, to non-medical persons along with training in basic overdose management, naloxone administration and after-care. The concept of THN constitutes an example of technology transfer, in that it tells us to take the solution (emergency naloxone kit) from the hospital into the community where the emergency (overdose) occurs, with the aim to reduce any harm (i.e. potential death) that could occur during the time delay while awaiting the arrival of an ambulance. This follows the example of technology transfer for other medical emergencies, such as diabetic coma or severe anaphylactic reactions, where potentially life-saving medicines (adrenalin/epinephrine, glucagon, snake anti-venom) are also pre-provided to the at-risk patient or individual, along with instructions to family members.

‘Take-home naloxone’ may be provided as part of comprehensive clinical care of patients in treatment for opioid use disorder in primary or specialty care or through dedicated THN schemes. Researchers have found that both peers and family members are highly willing to act as first responders providing interim emergency care whilst awaiting the arrival of ambulance (or other emergency medical care) [7, 8].

Support for THN has increased greatly in recent years, including guidelines from the World Health Organization [9] and endorsement from the United Nations [10], as well as from various national governments. Nevertheless, intervention inertia around THN continues to exist, with countries, services and clinicians uncertain what they can, or cannot, provide. In the meantime, lives continue to be lost, including in situations where the overdose victim was still alive at the point of discovery (and therefore death was likely preventable).

In this review, we provide the reader with the evidence for THN as a public health response to opioid overdose as well as presenting evidence on naloxone’s properties and the recent development of novel naloxone formulations and devices for layperson use as well as exploring ongoing challenges for implementation. To achieve a balanced reflection of the wider developments in the field of THN, we have deliberately brought together a team of co-authors with different areas of expertise who have led on the sections of this review. This paper draws on peer-reviewed and grey literature identified through a desk-based review of THN and naloxone formulations. English-language peer-reviewed literature was identified through searches of MEDLINE, EMBASE (via the OVID platform) and PubMed. Given the wide scope of this review, search terms included opioids, opiates, overdose, mortality, prevention, naloxone, intranasal (IN) and nasal. This general search strategy was then adapted by our co-authors to meet the specific focus of their respective section(s). Key literature (including grey literature) cited within the retrieved material was also consulted, and as a general principle, more recent literature was preferred over older data (1960–2009). Data from published papers and reports were extracted and synthesised as narrative reviews. We also refer the interested reader to earlier reviews with a more limited focus, including THN programmes [11, 12] and naloxone delivery systems [13, 14].

1.1 Structure

Sections 2 and 3 discuss the prevalence of opioid mortality and situational risk factors for overdose. Section 4 describes the discovery of naloxone and history of its use in medical practice as background for Sect. 5, which covers the study of pharmacokinetics and development of non-injectable formulations of naloxone. This is followed by a discussion of the pharmacodynamics of naloxone and its efficacy and safety for the emergency management of opioid overdose in Sect. 6. In Sects. 7 and 8, we review the effectiveness of THN and ongoing implementation barriers, leading up to concluding remarks in Sect. 9.

2 Epidemiology of Opioid Overdose Deaths

To understand the levels and variations in opioid overdose deaths, it is important to understand one of the important drivers in high-income countries, namely, prescription
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opioid utilisation [15]. North America, Oceania and some western European countries account for more than 95% of the worldwide consumption of analgesic opioids [15], as illustrated in Fig. 1, despite accounting for only about 15% of the world’s population [15].

2.1 USA

2.1.1 Increasing Opioid Prescribing

The USA had a substantial increase in opioid prescribing for chronic non-cancer pain [16], with opioid analgesic use in North America rising from about 2.4 billion DDD (defined daily dose; i.e. “the assumed average maintenance dose per day for a drug used for its main indication in adults”\(^1\)) per annum in 2001–2003 to about 5.3 billion DDD per annum in 2011–2013 [15].

In recent years, a number of policies have been introduced to reduce the problem of excessive opioid prescribing [17]. These have included educating professionals and the general public about appropriate prescription opioid use, implementation of prescription drug monitoring programmes, attempts to reduce egregious prescribing (by so-called “pill mills”—clinics devoted entirely to opioid prescribing) and developing novel abuse-deterrent opioid formulations. Despite an overall reduction in per capita prescribing since 2010, the oral morphine equivalent amount prescribed in 2015 was still approximately three times as high as in 1999 [18]. The unintended consequences of these supply-side interventions need to be considered. With access reduced and demand still high, individuals began to turn to the black market [19].

2.1.2 Increasing Opioid-Related Overdoses

The rise in opioid analgesic use in the USA has been associated with substantial increases in non-fatal and fatal overdose. Since 2000, there has been a steady increase in the rate of opioid overdose deaths (see Fig. 2). From 1999 to 2015, more than 183,000 people died from overdoses related to prescription opioids. In 2016 alone, there were over 63,000 drug-related deaths, of which 66% involved opioids (either prescription or illicit). In fact, the escalation in opioid overdose deaths has been a significant factor in reduced US life expectancy in 2015 and 2016, particularly for white Americans [20].

2.1.3 Re-emergence and Increase of Heroin Use

Despite concerted public health efforts to reduce opioid prescribing (see above), opioid-related deaths continued to increase by 16% from 2014 to 2015 [18]. These significant increases in mortality were largely driven by opioids other than methadone, predominantly IMF (see below) and heroin [21].

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\(^1\) https://www.whocc.no/ddd/definition_and_general_considera/.
A population-based study reported that the prevalence of heroin use increased from 0.3% in 2001–2002 to 1.6% in 2012–2013 [22], and heroin-related overdose increased from 1842 deaths in 2000 to 10,574 deaths in 2014 [17] (see Fig. 2). The increased availability of high-purity heroin combined with its low price (compared with diverted prescription opioids) have been major drivers of the upward trends in heroin use and overdose deaths [21]. Nonmedical use of prescription opioids is considered a significant risk factor for heroin use [17].

### 2.1.4 Increasing Availability of Potent Illicit Opioids

In recent years, deaths from illicit synthetic opioids have outstripped deaths due to heroin and prescribed opioids in the USA. In 2015, the Drug Enforcement Administration and Centers for Disease Control and Prevention issued nationwide alerts identifying IMF as a threat [23].

Illicitly manufactured fentanyl and its analogues are significantly more potent than morphine: carfentanil, for example, is approximately 10,000 times more potent, gram for gram. Fentanyl is more likely to lead to overdose than other opioids and is thought to have reduced cross-tolerance to other opioids [24]. A much smaller dose of fentanyl than heroin is required to achieve the same drug effect: thus errors at the level of the illicit manufacture and distribution, as well as errors at the level of the end user, result in frequent accidental errors of dosage whose implications are far more profound than errors where the dose alteration is less severe [24]. The high potency of fentanyl makes it often both cheaper and easier to trade (and access) than heroin given its much smaller bulk [19, 25]. Illicitly manufactured fentanyl is also increasingly found as adulterant in non-opioid drugs (e.g. cocaine) and considered a key factor for the more recent increases in opioid overdose mortality [21, 25, 26] (see Fig. 2).

In Massachusetts (2014–2016), 36% of fentanyl deaths had evidence of an overdose occurring within seconds to minutes after drug use, and 90% of fentanyl overdose decedents were pulseless upon emergency medical services (EMS) arrival [27]. It has been speculated whether massive overdoses are able to benefit from naloxone at all [28].

Fentanyl and its analogues differ from many other opioids by their propensity to induce muscle rigidity [29], including chest wall rigidity (sometimes called ‘wooden chest syndrome’), which makes assisted ventilation and breathing difficult [24] and may increase mortality risk and lead to rapid death [30]. Notably, muscle rigidity can be reversed by naloxone [30, 31]. Therefore, the same principles of response with THN initiatives apply to the prevention of deaths from fentanyl overdose, although early administration of naloxone is likely crucial.

### 2.2 Canada

Canada has the second highest level of prescribed opioid use globally after the USA, and consumption is increasing faster than in the USA, with a 203% increase from the 2-year period of 2000–2002 (8713 defined daily doses for statistical purposes) to 2008–2010 (26,380 defined daily doses [2]).

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https://www.incb.org/incb/en/narcotic-drugs/Availability/availability.html.
for statistical purposes) [15]. From 1991 to 2004, annual prescriptions for opioids increased from 458 to 591 per 1000 individuals [32], with opioid-related deaths doubling, from 13.7 to 27.2 per million in 2004. There are also recent indications of the use of IMF and analogues and associated overdoses [33], localised to certain provinces, such as British Columbia and Alberta [33].

### 2.3 Europe

Although pharmaceutical opioid use is increasing gradually in Western Europe, the prevalence is four times lower than in North America [34]. In general, heroin remains the most prevalent illicit opioid in Europe. However, in some countries, such as Estonia and Finland, where the heroin market plummeted in the early 2000s, heroin has been entirely displaced by fentanyl (mostly illicit) and buprenorphine, respectively [35].

In 2015, there were 8440 drug overdose deaths (any substance; occurring most commonly in the UK, followed by Germany and Sweden) in the European Union (plus Turkey and Norway), a total that continues to rise [36]. Most drug-related deaths (79%) involved opioids, predominantly heroin, though potent synthetic opioids such as fentanyl and its newer derivatives seem to play an increasing role [37]. For example, in Estonia there was a spike in overdose deaths in 2012 that was mainly attributable to fentanyl. On the European boundary, there have also been reports of fentanyl use in parts of Russia owing to heroin shortages. Of the 9263 drug-related deaths reported in Russia in 2010, 6324 were attributed to opioid use [38]. (N.B.: Comparisons across different countries are hampered by differences in data collection [39]).

### 2.4 Australia

In Australia, annual opioid analgesic use increased from 23 million DDD in 2001–2003 to 106 million DDD in 2011–2013 [15]. In 2016, there were 1045 opioid-related deaths in Australia, accounting for approximately 75% of all drug-related deaths [40]. Whilst the rate of heroin-related deaths has remained stable, overdose deaths from prescription opioids are now more common, having increased by 127% from 2006 to 2016 [40]. Deaths related to synthetic opioids (fentanyl, tramadol and pethidine, excluding methadone) have also been increasing, from approximately 5% (1999) to 17% (2016) of opioid-related deaths [40].

### 3 Clusters of Deaths and Crucial At-Risk Populations

Various factors are associated with increased risk of overdose death in people who use opioids (PWUO). These include individual and behavioural risk factors, such as male sex, older age, intravenous (IV) use, co-administration of other sedative drugs (e.g. benzodiazepines, alcohol), and irregular patterns of use (probably due to variability of dose) [41]. Sadly, many overdose deaths conform to Hans Fallada’s book title “Every Man Dies Alone”, as a significant number (approximately 30%) of opioid overdose fatalities happens when the drug is taken in the absence of other people [42]. It is also a serious problem that those around the individual often do not understand that the victim is not sleeping, but in imminent danger from an overdose [27].

Different types of opioid overdose victims die at different times after injection. In heroin overdose, the time course of opioid metabolites in post-mortem cases indicated that most victims stayed alive for more than 30 min, indicating a window for intervention within at least this time frame [43, 44]. However, in 44% of the 145 fatal cases, signs of rapid-onset collapse were also observed. The time frame for intervention may nevertheless be shorter with fentanyl overdoses, as anecdotal reports of immediate deaths in 14 subjects with needle/tourniquet in place (along with characteristic fentanyl/metabolite ratios) suggest [45]. In the case of immediate deaths from fentanyl or heroin, it is possible that these may not be secondary to respiratory depression, but rather the direct effect of primary cardiovascular collapse in susceptible undernourished and dehydrated individuals.

Situational factors are important, with the prison release population identified as a particularly high-risk group. For prisoners with a previous history of heroin use by injection, one in 200 will be dead within a fortnight of their release from prison, with most deaths being caused by opioid overdose [46–48]. These deaths likely occur during post-release deliberate intoxication, where, following a period of abstinence and consequent loss of tolerance, a previously regular dose may now lead to fatal overdose.

A similar clustering of deaths is seen after discharge from hospital or abstinence-based residential rehabilitation [49, 50], and the explanation is likely similar. In addition, those in abstinence-based outpatient treatment are at increased risk if they then relapse [51].

The importance of this clustering characteristic is that it should direct both policymakers and clinicians to points of transition in care and setting that would benefit from preventive measures and emergency interventions, including THN. The target population for THN crucially also includes PWUO who are outside the treatment system. We need to remember that treatment markedly reduces the risk of overdose death.

A different type of clustering is seen with outbreaks of overdose deaths in communities, which occur with the presumed arrival and distribution of a particularly potent batch of heroin or contaminated supply (e.g. with added fentanyl). Certainly, this accounts for some of the clusters of deaths, but it is unclear why this does not then lead to modification of drug-taking
behaviour (e.g. avoiding this supply, or taking a smaller dose), and it is likely that other factors, as yet unidentified, lie behind some of these time-limited geographical clusters.

4 History of Naloxone for the Emergency Management of Opioid Overdose

4.1 Initial Development and Testing (1960s)

Naloxone (N-allylnoroxymorphone) hydrochloride (CAS Number: 357-08-4) was first synthesised from thebaine in New York in 1960 by Dr. Jack Fishman and Dr. Mozes J. Lewenstein, whose patent described naloxone as a “more potent antagonist to the respiratory depressive effects of potent analgesics than the antagonists hitherto known” [52, 53]. Its molecular formula is C₁₉H₂₁NO₄HCl (molecular weight 363). Naloxone hydrochloride is a white-to-slightly-off-white powder and is soluble in water, dilute acids and strong alkali. It is slightly soluble in alcohol and practically insoluble in ether and chloroform. Naloxone has a fat-water partition coefficient 20 times that of morphine and similar to meperidine. Its pKa³ is 7.94 [54]. The chemical structure of naloxone is shown in Fig. 3.

4.2 Entry into Clinical Practice (1970s Onwards)

The US Food and Drug Administration (FDA) approved naloxone in 1971 as a prescription-only medication for intravenous (IV), intramuscular (IM) and subcutaneous administration for reversing the effects of opioids. Naloxone entered international clinical practice in the following years and was included as a specific antidote (i.e. in the formulation of 0.4 mg in 1-mL ampoules) in the World Health Organization’s Model List of Essential Medicines in 1983 [55, 56]. Owing to its unique effectiveness and safety profile, naloxone has become the treatment of choice for reversing opioid overdose in hospital emergency departments and ambulance services.

4.3 Community-Based Naloxone (1990s)

The first serious consideration of THN occurred in a 1996 BMJ editorial [57], which described how emergency naloxone kits could be pre-provided for emergency use to the following groups:

1. individuals at high risk of overdose, e.g. those leaving emergency care following overdose and those who lost tolerance as a result of detoxification, incarceration or abstinence-based treatment;
2. patients enrolled in treatment programmes; and
3. active users.

People who use opioids were described as the primary target group for THN because they are at risk of future overdose themselves and highly likely to be in a position to intervene in someone else’s overdose. Indeed, PWUO voiced strong support of THN provision. A London-based survey of PWUO [7] estimated that two thirds of witnessed overdose deaths could have been avoided with THN, and nearly 90% of respondents who had witnessed an overdose stated that they would have used the medication had it been available. Subsequent studies identified the willingness of PWUO [58, 59] and family members [8] to be trained in overdose management and naloxone administration.

Early implementation of THN was made possible through user advocates working with physicians willing to prescribe naloxone despite medicolegal uncertainty or explicit barriers. Take-home naloxone provision first occurred in the late 1990s and early 2000s, in the USA (Chicago, San Francisco), Germany (Berlin), the UK (Jersey) and Italy (Turin, Bologna, Padua) [60–69]. The identification of legal pathways for THN provision from the mid-2000s onwards facilitated the introduction of first national and state-wide programmes in parts of Europe and USA.

5 Pharmacokinetics and the Development of New Non-injectable Naloxone

5.1 Notion of Non-injectable Formulations: Potential for more Acceptable Implementation

Because injectable naloxone was not developed for layperson use, non-injectable naloxone formulations have been considered to have several implementation advantages for THN programmes. First, injectable medications are

³ pKa is defined as the negative base-10 logarithm of the acid dissociation constant (Kₐ) of a solution. The lower the pKₐ value, the stronger the acid.
intimidating for laypersons to use [70] and present logistical barriers: they require product assembly (e.g. needle and syringe) and training in administration. Surprisingly, even among experienced PWUO, nearly three out of four (74%) preferred naloxone nasal spray to injectable devices [71]. Second, with use of naloxone by injection, there is the risk of needle-stick injury. Third, non-injectable naloxone could likely overcome regulatory obstacles (e.g. prescription-only status for injectable medications) and be provided to a wider workforce (e.g. hostel staff, outreach workers, police).

5.2 Routes of Administration

Naloxone was originally developed and FDA approved as an injectable formulation for use by medical professionals (see above). The IV route requires skilled personnel and takes time to establish. Intramuscular is now the widely preferred parenteral route because IV carries a greater risk of exposure to contaminated blood as well as risks of withdrawals and potential aggression. Intranasal is the simplest and fastest procedure, and it circumvents the contamination hazard. While IV provides by far the most rapid and reliable response when measured from the time of administration, IM comes as a good second, followed closely by IN (see Table 1). However, when time from contact with the patient is the starting point in the emergency situation, IN [72] and subcutaneous [73] have been found to be as fast as IV. A potential shortcoming with the IN route is that effective uptake may be reduced by septal abnormalities, bleeding, nasal mucous, trauma and use of nasal vasoconstrictors [74].

These ‘improvised’ kits were used in many successful overdose reversals, and reports of problems using the kits were uncommon [75, 76]. For instance, in Norway, just over 2000 ‘improvised’ THN kits were distributed in 2014–2015, with 277 reports of successful reversals [77, 78]. However, concern was expressed about the reliance on such improvised nasal sprays because the naloxone concentration in the spray was dilute (1 mg/mL) [79]. Specifically, given the existence of licensed injectable products, the question was raised [79] whether it was acceptable for clinicians to supply unlicensed improvised nasal naloxone kits for take-home use, where no back-up of injectable naloxone would be available in the case of non-response (unlike the ambulance or hospital setting). This stirred a challenging debate among international experts in the field [80, 81].

Subsequent examination of patent records located data confirming only poor bioavailability (approximately 10% of dose administered) [82] and recent direct pharmacokinetic comparison of the improvised kits vs. the new concentrated naloxone nasal spray formulations found bioavailability of only approximately 20% with the improvised kits vs. more than 40% with the new concentrated naloxone sprays [83]. Nevertheless, the evident successful reversals of opioid overdoses with these improvised kits [75–77] should raise questions about the dose necessary for layperson reversal which, in these instances, appears to have been achieved with much lower absorbed concentrations of naloxone.

5.3 Improvised Nasal Naloxone Kits

In the absence of licensed non-injectable naloxone formulations, ‘improvised’ nasal kits (i.e. a 2-mg/2-mL pre-filled syringe with a nasal mucosal atomizer device [MAD] attached) were being distributed by some North American THN programmes from 2006 onwards (and later also in parts of Scandinavia and Scotland), despite not having been formally tested for safety or efficacy. This followed their use by EMS as well as fire fighters and police officers. (The safety of use of the improvised kits in medical practice is discussed in more detail in Sect. 6.6.)

In response to rising overdose mortality rates, the US FDA, Centers for Disease Control and Prevention, National Institute on Drug Abuse, and Office of the Assistant Secretary for Health and Human Services sponsored a stakeholder meeting in 2012 to encourage the development of non-injectable naloxone products suitable for layperson use in community-based settings. Following the meeting, the National Institute on Drug Abuse announced that it would provide funding for the development of user-friendly naloxone devices (i.e. IN rather than injection) [84].

The key regulatory criteria for any New Drug Application [85, 86] involved that naloxone would need to be absorbed into

| Route | Skills | Time to establish | Time to onset | Reliability | Risk for withdrawal |
|-------|--------|-------------------|---------------|-------------|---------------------|
| IV    | High   | Medium +          | Rapid         | High        | High                |
| IM    | Some   | Medium            | Medium +      | Medium to high | Moderate            |
| IN    | Little | Short             | Medium        | Unknown     | Moderate            |

IM intramuscular, IN intranasal, IV intravenous

*a Comparable doses*
the bloodstream rapidly, given the emergency situation, and in a quantity sufficient to effect quick reversal of opioid-induced respiratory depression. The reference for any candidate non-injectable product would be injectable naloxone, administered by the licensed IM, IV and subcutaneous routes [9].

5.4.1 Identification of Candidate Routes of Administration for Non-injectable Products

Naloxone has a very low oral and rectal bioavailability of 1–2% and 15%, which makes these routes unsuitable for emergency naloxone [87]. Strang et al. [88] thus explored candidate routes of administration for THN use. Only the transmucosal routes IN, buccal and sublingual met the required characteristics. Transmucosal uptake bypasses first-pass hepatic elimination. Among these three routes, approved products only exist for IN administration (see below). In the following section, we describe the pharmacokinetics of injectable naloxone as the reference for the development of non-injectable products.

5.5 Naloxone Pharmacokinetics

Naloxone is rapidly and extensively metabolised mainly by glucuronidation to the inactive compound naloxon-3-glucononide (N3G) thought to be primarily by the liver at that time [89]. Ngai et al. [90] and Berkowitz [91] reported initial distribution and terminal half-lives of 4 and 64 min. Recent pharmacokinetic studies [92–95] confirmed terminal elimination half-lives of 60–120 min after IV administration of 0.4–1.0 mg (see Table 2). It should be noted that arterial and venous serum concentrations are similar [95]. Older studies have shown a volume of distribution of 200 L and a total clearance of about 2 L/min [89, 96]. However, Yassen et al. [97] and Skulberg et al. [94, 98] reported volumes of distribution of about 320 L and clearances between 3 and 4 L/min. This is two to three times higher than the maximal liver clearance; consequently, naloxone must be exposed to significant extrahepatic metabolism.

For IM naloxone (see Table 4), the mean time to maximum concentration and mean dose-corrected maximum serum concentration (\(C_{\text{max}}\)) varied from 8 to 24 min and 1.55–4.66 ng/mL, respectively. The latter is far lower than those of IV 14.1–18.6 ng/mL (Table 2). However, dose-corrected AUCs were as expected rather similar for injectable naloxone (see Tables 2, 4), although AUCs were both AUC$_{\text{last}}$ and AUC$_{0-\text{t}}$ for IM.

5.5.1 Intranasal Route of Administration

The nose is readily available and presents the advantage that laypeople are already familiar with the use of nasal sprays. People who use opioids generally preferred nasal spray over injectable naloxone [71]. The mucosa of the nose is extensively perfused. Its endothelial lining is very open to the external environment. Moreover, its mucosa is constantly cleared by mucociliary transport, requiring that uptake takes place within about 15 min. However, the nose cannot accommodate spray volumes greater than 0.15 mL, as excess volume will be lost as nasal drip or post-nasally [99]. For this reason, many THN programmes have moved from using ‘improvised’ nasal kits (i.e. 2-mL syringes with attached MAD) to distributing approved nasal spray products, which deliver a volume of 0.1 mL.

### Table 2 Pharmacokinetics of intravenous naloxone

| References         | Dose (mg) | \(N\) | \(C_{\text{max}}\) (ng/mL) | Dose-corrected \(C_{\text{max}}\) | \(T_{\text{max}}\) (min) | AUC$_{\text{last}}$ (ng × h/mL) | Dose-corrected AUC | \(T_{1/2}\) (min) |
|--------------------|-----------|-------|-----------------------------|-------------------------------|-------------------------|-------------------------------|---------------------|-----------------|
| Skulberg et al. [94] | 0.4       | 22    | 7.44 (9.67)\(^a\)           | 18.6                          | 3.5 (4.2)\(^a\)         | 1.84 (1.49)\(^b\)            | 4.6                 | 74.3 (32.1)\(^a\) |
| Tylleskar et al. [95] | 1.0       | 12    | 14.1 (9.98–18.2)\(^b\)      | 14.1                          | Not reported            | 3.65 (3.05–4.27)\(^b\)       | 3.65                | Not reported   |
| McDonald et al. [92] | 0.4       | 34    | 5.94 (92.9)\(^c\)           | 14.9                          | 2 (1–5)\(^c\)           | 2.05 (0.4)\(^d\)             | 5.13                | 75 (13)\(^b\)  |
| Tylleskar et al. [93] | 1.0       | 12    | 14.2 (9.13–19.2)\(^b\)      | 14.2                          | 2.25 (1.70–2.80)\(^b\)   | 4 (3.45–4.55)\(^b\)          | 4                   | 70.1 (60.1–78.7)\(^b\) |
| Gufford et al. [103] | 2.0       | 6     | –                            | –                             | –                       | 10.8 (8.92–13.2)\(^d\)       | 5.4                 | 91 (64–130)\(^d\) |

\(AUC\) area under the curve, \(AUC_{\text{last}}\) maximum serum concentration, \(min\) minutes, \(T_{1/2}\) half-life, \(T_{\text{max}}\) time to maximum concentration

\(^a\)Arithmetic mean (standard deviation)  
\(^b\)Arithmetic mean (95% confidence interval)  
\(^c\)Median (minimum, maximum)  
\(^d\)Geometric mean (90% confidence interval)  
\(^e\)Geometric mean (% coefficient of variation)  
\(^f\)Partly under opioid exposure
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5.5.2 Approved Nasal Spray Products

Four high-concentrate and approved products, Narcan® [100], Nyxoid® [92], Nalscue® (in France), and Ventizolve® (12 European countries) [94] utilise the Aptar Pharma (hereafter referred to as ‘Aptar’) unit dose spray (0.1 mL, see Table 3). Narcan® by Adapt Pharma (hereafter referred to as “Adapt”) was the first approved nasal spray product, having received FDA approval in November 2015 [101] and Health Canada approval in October 2016. Adapt subsequently also received regulatory approval for a 2-mg/0.1-mL version of

| Product name | Narcan® | Nalscue® | Nyxoid® | Ventizolve® |
|--------------|---------|----------|---------|-------------|
| Manufacturer | Adapt®  | Indivior | Mundipharma | Den norske Eterfabrikk (DnE) |
| Regulatory status: licensed in | USA (November 2015) | France: July 2016 (temp.), July 2017 (marketing authorisation) | Europe (EMA, September 2017) | 12 European countries (May 2018) |
| Formulation | 4 mg/0.1 mL, (2 mg/0.1 mL)a | 0.9 mg/0.1 mL | 2 mg/0.1 mLb | 1.4 mg/0.1 mL |
| Relative bioavailability | 46% | 37% | 47% | n/a |
| Absolute bioavailability | 47% | 52–54% |

*EMA European Medicines Agency, FDA US Food and Drug Administration, n/a, NR, OTC, temp.*

*a A 2-mg/0.1-mL version of Narcan® by Adapt was approved in North America but, to our knowledge, has never been introduced in clinical practice. Kreiter et al also report, in their 2019 paper, that they had been informed by Adapt Pharma, the manufacturer, that the 2mg product had not been marketed and that there were no current plans to launch it.

*b All formulations in this table are reported as naloxone hydrochloride, with the 1.8 mg of naloxone listed in Nyxoid being equivalent to 2 mg of naloxone hydrochloride.

*c A generic version of the 4-mg/0.1-mL Narcan nasal spray by Teva Pharmaceuticals received FDA approval in April 2019 (https://www.fda.gov/drugs/developmentapprovalprocess/howdrugsaredevelopedandapproved/drugandbiologicalapprovalreports/andagenericdrugapprovals/)

### Table 3 Comparison of nasal sprays approved in different countries

| Product name  | Manufacturer | Regulatory status: licensed in | Formulation | Relative bioavailability | Absolute bioavailability |
|---------------|--------------|--------------------------------|-------------|-------------------------|--------------------------|
| Narcan®       | Adapt®       | USA (November 2015)            | 4 mg/0.1 mL, (2 mg/0.1 mL)a | 46%                     | 47%                      |
| Nalscue®      | Indivior     | France: July 2016 (temp.), July 2017 (marketing authorisation) | 0.9 mg/0.1 mL | 37%                     |                         |
| Nyxoid®       | Mundipharma  | Europe (EMA, September 2017)   | 2 mg/0.1 mLb | 47%                     | 52–54%                   |
| Ventizolve®   | Den norske Eterfabrikk (DnE) | 12 European countries (May 2018) | 1.4 mg/0.1 mL | n/a                     |                         |

| Product name  | Manufacturer | Regulatory status: licensed in | Formulation | Relative bioavailability | Absolute bioavailability |
|---------------|--------------|--------------------------------|-------------|-------------------------|--------------------------|
| Narcan®       | Adapt®       | USA (November 2015)            | 4 mg/0.1 mL, (2 mg/0.1 mL)a | 46%                     | 47%                      |
| Nalscue®      | Indivior     | France: July 2016 (temp.), July 2017 (marketing authorisation) | 0.9 mg/0.1 mL | 37%                     |                         |
| Nyxoid®       | Mundipharma  | Europe (EMA, September 2017)   | 2 mg/0.1 mLb | 47%                     | 52–54%                   |
| Ventizolve®   | Den norske Eterfabrikk (DnE) | 12 European countries (May 2018) | 1.4 mg/0.1 mL | n/a                     |                         |

### Table 4 Pharmacokinetics of intramuscular naloxone

| References          | Dose (mg) | $N$  | $C_{max}$ (ng/mL)c | Dose-corrected $C_{max}$ | $T_{max}$ (min)d | $AUC_{int}$ (ng × h/mL)c | Dose-corrected $AUC$ | $T_{1/2}$ (min)e |
|---------------------|-----------|------|--------------------|--------------------------|----------------|--------------------------|---------------------|----------------|
| Skulberg et al. [94]| 0.8       | 22   | 3.73 (3.34)a       | 4.66                     | 13.6 (15.4)a   | 3.43 (0.66)a             | 4.29                | 84.8 (26.5)a    |
| Skulberg et al. [98] | 0.8       | 12   | 3.62 (2.64–4.60)b  | 4.53                     | 7.75 (5.01–10.5)b| 4.07 (3.28–4.87)b      | 5.08                | 69.7 (59.5–79.8)b|
| McDonald et al. [92] | 0.4       | 34   | 1.27 (55.8)        | 3.18                     | 10 (4–90)      | 2.01 (17.7)              | 5.03                | 81 (16)       |
| Gufford et al. [103]| 2.0       | 6    | 3.1 (2.3–4.2)c     | 1.55                     | 22.5 (10–60)d  | 7.23 (6.43–8.12)c       | 3.61                | 100 (89–111)c   |
| Evzio (Kaleo, 2016)| 0.4       | 30   | 1.2 (51.4)         | 3                        | 15.0 (4.80–73.8)| 1.9 (23.4)              | 4.75                | 78 (38)       |
|                    | 0.4 × 2   | 24   | 2.2 (47.4)         | 2.75                     | 12.6 (5.40–51.0)| 3.8 (19.1)              | 4.75                | 90 (23.7)     |
|                    | 2         | 24   | 7.9 (45.8)         | 3.95                     | 15.0 (7.8–40.2)| 10.3 (15.2)             | 5.15                | 90 (23.7)     |
| Krieter et al. [100]| 0.4       | 29   | 0.90 (31.2)        | 2.25                     | 24.0 (6.0–126) | 1.8 (22.7)              | 4.5                 | 78 (27.8)c     |
| Gufford et al. [103]| 2.0       | 6    | 3.1 (2.3–4.2)c     | 1.55                     | 22.5 (10–60)d  | 7.23 (6.43–8.12)c       | 3.61                | 100 (89–111)c   |

AUC area under the curve, $AUC_{int}$, min minutes, $AUC_{last}$, $C_{max}$, maximum serum concentration, opioid partly under opioid exposure, $T_{1/2}$ half-life, $T_{max}$ time to maximum concentration

a Arithmetic mean (standard deviation)
b Arithmetic mean (95% confidence interval)
c Geometric mean (% coefficient of variation)
d Median (minimum, maximum)
e Geometric mean (90% confidence interval)
f $AUC_{last}$
the spray, but this product is not commercially available at the time of writing.4

A competitor product by Indivior (Nalscue®, 0.9-mg/0.1-mL formulation, also in the Aptar device) failed to receive FDA approval because the nasal spray was found not to be absorbed sufficiently rapidly relative to IM naloxone [102]. Nonetheless, this product secured a marketing authorisation in France in July 2017.

In Europe, Mundipharma received European Medicines Agency approval in November 2017 for a 2-mg/0.1-mL naloxone hydrochloride dihydrate (i.e. 1.8-mg naloxone base; Nyxoid®) concentrated nasal spray for delivery by the same Aptar device as the other new nasal sprays, and has subsequently been rolled out progressively across much of Europe. The pharmacokinetics of the spray formulation have been published [92].

In June 2018, the Norwegian company AS Den Norske Eterfabrikk received regulatory approval in 12 European countries for a 1.26-mg base (1.4 mg hydrochloride) naloxone nasal spray (Ventizolve®); its pharmacokinetics has been published [94].

5.5.3 Intranasal Naloxone: Pharmacokinetics

As part of the development of these high-concentrate naloxone spray products (and their application for regulatory approval), a number of recent studies have explored the pharmacokinetics of IN naloxone (see Table 4). Nasal naloxone doses in these studies ranged from 0.8 to 8 mg. (In the small study [n = 6] by Gufford et al. [103], 0.1- and 0.2-mL volumes were used. Edwards et al. [74] also studied the MAD delivering 2.0 mL).

The high-concentrate IN formulations all have reasonably rapid uptake with a mean time to maximum concentration of 15–30 min, which is somewhat slower than the 8–24 min for the IM formulations (Tables 3, 4; see also Fig. 4). The mean dose-corrected C max for IN varied from 1.29 to 2.04 ng/mL, which is lower than for IM (mean 1.55–4.66 ng/mL). A dose–serum concentration relationship was repeatedly reported. Dose-corrected AUCs were far lower (1.40–3.34 ng/mL) than for injectable formulations. This conforms with the relative bioavailability of IN to IM of about 44–54% for the high-concentrate formulations, while only 10–15% was found for the dilute formulations [74]. However, Skulberg et al. [98] found a far higher bioavailability (75%) in subjects under exposure to the opioid remifentanil for 2 hours of the 6-hour study. Whether this is a specific remifentanil or generic opioid effect remains unknown, but a potential higher bioavailability under an opioid may have implications for future regulatory studies of nasal administration.

Absolute bioavailability for concentrate nasal sprays varied from 47 to 54% [92, 93]. Most importantly, these high-concentrate sprays deliver therapeutic doses (0.4–2.0 mg) in a single 0.1-mL spray, and a second spray gives a proportionate rise in serum concentrations. In contrast, dilute nasal spray (2 mg/2 mL) administered via the MAD only had 11–20% absolute bioavailability, implying that a subtherapeutic dose of approximately 0.2–0.4 mg was delivered [74, 82, 83]. A recent study confirmed that, even after two administrations, dilute nasal spray (2 mg/2 mL, administered via a MAD) failed to achieve naloxone plasma concentrations comparable to concentrate nasal sprays (2 mg/0.1 mL, 4 mg/0.1 mL) at any time [83].

4 https://www.accessdata.fda.gov/scripts/cder/daf/index.cfm?event=overview.process&ApplNo=208411.
When comparing IV with IM and IN, it is evident that a comparable dose given IV generates immediate and far higher initial serum concentrations. It is also important to acknowledge that the $C_{\text{max}}$ for the 2- to 4-mg doses of IM (e.g., Evzio®) and IN are far higher than those of the 0.4-mg IM comparator. This is relevant for the discussion below on the risk of withdrawal symptoms.

Skulberg et al. [94] used pharmacokinetic modelling to illustrate the relationship between first-responder nasal naloxone dosing and subsequent IM administrations by ambulance paramedics 10 min later. First-responder dosing of 1.4 mg IN produced plasma concentrations that were higher than 0.4 mg IM 10 min later all the time up to 20 min but lower than for 0.8 mg IM in the first 15 min. However, the combined 1.4 mg IN and 0.4 mg IM 10 min later were higher than 0.8 mg IM alone at the same time. It was concluded that it was beneficial to administer IN for up to 2 min before an ambulance paramedic delivered a 0.4-mg IM dose.

### 6 Pharmacodynamics, and Efficacy and Safety of Naloxone for the Emergency Management of Opioid Overdose

#### 6.1 Pharmacodynamics

Naloxone is essentially a pure competitive MOR antagonist. Its affinity compares well with the opioid agonists such as morphine, methadone and fentanyl, thus being capable of reversing their actions (see Table 5) [104]. It is estimated that a dose of 0.9 mg/70 kg occupies about 50% of brain receptors in opioid-naïve subjects [105].

Animal testing characterised naloxone as a ‘potent, rapid-acting, and relatively pure narcotic antagonist’, which counteracted the effects of a range of opioid agonists, including morphine and methadone, without agonist activity of its own [106, 107]. In humans, it was found that 0.35 mg/70 kg of naloxone had a greater effect on opioid-induced respiratory depression than 150 and 20 mg/kg of nalorphine and levallorphan, respectively [108]. Naloxone did not, in contrast to the two others, induce respiratory depression in subjects not given an opioid. Therefore, it was recommended that naloxone should be the opioid antagonist of choice for clinical use [108].

The partial agonist buprenorphine is different (see Table 6). It has a high affinity to MOR and dissociates slowly, in contrast to the much faster dissociation of naloxone [109]. Thus, naloxone reversal of the action of buprenorphine is slow and far larger doses are required [110].

Naloxone equilibrates rapidly with the site of action in the brain as shown by its blood-effect site equilibration half-life of 6.5 min [97], a little slower than that of fentanyl [111]. It is speculated whether an active transport system is involved in this uptake process [112]. In a very recent, positron emission tomography study of nasal naloxone in healthy volunteers, carfentanil was used for initial characterisation of MOR availability. The MOR occupancy was slightly delayed to serum concentrations, and half of peak occupancy was reached at 10 min. The 4-mg IN dose gave up to 85% MOR occupancy [113]. These experimental findings concord with the clinically observed rapid onset of action of naloxone.

Berkowitz [91] showed in the rat that brain concentrations of subcutaneous naloxone rose as rapidly as serum concentrations and stayed steadily above these during a parallel decline. For the agonist morphine, both uptake and egress from the brain lagged its serum concentrations all the time.

Kaufman et al. [114] reported that naloxone in healthy men had a dose-dependent duration of about 1.5 hours on morphine-induced respiratory depression. Onset of action after about 0.35 mg/70 kg IV was 2–3 min, peaking at 25 min. A dose of about 0.33 mg/70 kg was required to reduce the effect of 12 mg of morphine to that of 4 mg.

Pupilometers were used by Gufford et al. [103] to study the attenuating effect of naloxone (2 mg IM and IN) on 4 mg of oral alfentanil (with or without grapefruit juice). However, only moderate increases (about 20%) of the pupil diameter were found. Skulberg et al. [98] used target-controlled infusion of remifentanil to create steady-state opioid agonism. Intramuscular naloxone (0.8 mg) reversed pupil size significantly and was superior to the same IN dose. In a subsequent paper from the same group [95], the duration of action of 1 mg of naloxone IV was 120 min. Moreover, a minimum effective concentration of naloxone in steady state of 0.5 ng/mL was established.

Middleton et al. [115] compared subjective effects and pupil diameter in ten non-dependent PWUO after giving 2 mg and 8 mg of buprenorphine, alone or in combination with 0.5 mg and 2 mg of naloxone. No statistically significant differences were found between formulations. Experimental studies of the effects of naloxone in opioid addicts are rare. Loimer et al. [116] provoked abstinence with naloxone for diagnostic and therapeutic purposes in dependent

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Table 5 Opioid receptor affinities (modified from [104])

| Drug          | Opioid receptors |   |   |   |
|---------------|-----------------|---|---|---|
|               | Mu (μ)           | Gamma (γ) | Kappa (κ) |
| Naloxone      | – – –            | – –        | – –        |
| Morphine      | +++              | No effect  | No effect  |
| Methadone     | +++              | No effect  | No effect  |
| Fentanyl      | +++              | No effect  | +          |
| Buprenorphine | P                | No effect  | – –        |

$P$ partial agonist action, + agonist, – antagonist
PWUO in Pakistan, assessing withdrawal using an Objective Opioid Withdrawal Scale. Peak onset of action was within a few minutes for IV, and about 15 min for IM and IN. Durations of actions exceeded 90 min for all administrations.

Overall, several factors affect the outcome of naloxone antagonism (see Table 7). Most of these factors are unknown in an overdose situation, where it is not clear what drugs were consumed. Therefore, naloxone should always be titrated to restore adequate respiration and cognitive function.

### 6.2 Clinical Effects

Naloxone can reverse serious effects of opioids such as respiratory depression and stupor. Naloxone does not cause physical or psychological dependence and has virtually no effect in a healthy non-dependent person. Naloxone doses

| Drug                        | Blood brain equilibration | Egress from brain | Dissociation rate | Terminal half-life |
|-----------------------------|---------------------------|-------------------|-------------------|-------------------|
| Naloxone                    | Rapid                     | Rapid             | Rapid             | Short             |
| Morphine                    | Slow                      | Slow              | ?                 | Medium            |
| Methadone                   | Rapid?                    | ?                 | ?                 | Long              |
| Fentanyl                    | Rapid                     | Medium            | ?                 | Medium            |
| Buprenorphine               | ?                         | ?                 | Slow              | Long              |

### Table 6 Pharmacokinetics of intranasal naloxone

| References                  | Dose (mg) | N  | \(C_{\text{max}}\) (ng/mL)\(^b\) | Dose-corrected \(C_{\text{max}}\) | \(T_{\text{max}}\) (min)\(^c\) | AUC\(_{\text{inf}}\) (ng × h/mL)\(^b\) | Dose-corrected AUC | \(T_{1/2}\) (min)\(^b\) | \(F_{\text{IM}}\)% |
|-----------------------------|-----------|----|---------------------------------|-------------------------------|-------------------------------|---------------------------------|------------------|-----------------|----------------|
| Skulberg et al. [94]        | 1.4       | 22 | 2.36 (0.68)\(^d\)               | 1.69                          | 20.2 (9.4)\(^d\)              | 2.84 (0.94)\(^d\)               | 2.03             | 73.0 (20.2)\(^d\) | 52             |
|                             | 2.8 (1.4 × 2) | 22 | 4.18 (1.53)\(^d\)               | 1.49                          | 20.7 (9.54)\(^d\)             | 5.47 (1.89)\(^d\)               | 1.95             | 69.8 (12.8)\(^d\) | 75\(^n\,\text{opioid}\) |
| Skulberg et al. [98]\(^n\,\text{opioid}\) | 0.8       | 12 | 1.63 (1.25–2.02)\(^c\)          | 2.04                          | 28.0 (22.0–34.0)\(^c\)        | 3.25\(^c\)                      | 3.34             | 63.7 (59.2–68.2)\(^c\) | 59.1           |
| Krieter et al. [100]        | 2         | 29 | 3.1 (51.4)                       | 1.55                          | 18 (18–126)                   | 4.7 (29.8)                      | 2.35             | 114 (34.6)      | 51.9           |
|                             | 4         | 29 | 5.3 (44.6)                       | 1.33                          | 30 (12–60)                    | 8.5 (39)                        | 2.13             | 132 (34.6)      | 46.2           |
|                             | 4 (2 × 2) | 29 | 6.5 (32.3)                       | 0.81                          | 18 (12–36)                    | 9.7 (26.7)                      | 2.43             | 144 (31.7)      | 53.5           |
|                             | 8 (4 × 2) | 29 | 10.3 (38.1)                      | 1.03                          | 18 (12–609)                   | 15.8 (23.1)                     | 1.98             | 132 (39.0)      | 43.9           |
| Tylleskar et al. [93]       | 0.8       | 12 | 1.45 (1.07–1.84)\(^c\)          | 1.81                          | 17.9 (11.4–24.5)\(^c\)        | 1.65 (1.28–1.72)\(^c\)         | 2.06             | 89.7 (76.8–103)\(^c\) | 54\(^c\) |
|                             | 1.6 (0.8 × 2) | 12 | 2.57 (1.49–3.66)\(^c\)          | 1.61                          | 18.6 (14.4–22.9)\(^c\)        | 3.08 (2.05–4.13)\(^c\)         | 1.93             | 79.0 (65.3–92.7)\(^c\) | 52\(^c\) |
| McDonald et al. [92]        | 1         | 32 | 1.51 (50.2)                      | 1.51                          | 15 (10–60)                    | 2.69 (40.5)                     | 2.69             | 80 (23)         | 50.2           |
|                             | 2         | 33 | 2.87 (49.6)                      | 1.44                          | 30 (8–60)                     | 4.97 (38.5)                     | 2.49             | 84 (30)         | 46.8           |
|                             | 4 (2 × 2) | 33 | 6.02 (54.5)                      | 1.51                          | 15 (10–60)                    | 10.07 (35.8)                    | 2.52             | 102 (28)        | 48.1           |
| Gufford et al. [103]        | 1.0       | 6  | 5.7 (3.3–10)\(^e\)              | –                             | 12.5 (5–15)\(^f\)             | 4.7 (3.33–6.65)\(^f\)          | 2.35             | 61 (53–72)      | 75             |
|                             | 2.0 (1 × 2) | 6  | 3.0 (1.7–5.3)\(^e\)             | 1.5                            | 5.0 (5–15)\(^f\)              | 2.8 (1.95–4.0)\(^f\)           | 1.40             | 80 (56–113)     | 44             |
| Evzio (Kaleo, 2016)         | 2/2 mL    | 36 | 1.3 (48.3)                       | 0.65                          | 15 (4.2–40.8)                 | 1.5 (31.6)                      | 0.71             | 90 (18.5)       | 14.6           |
|                             | 2/2 mL    | 36 | 0.7 (52)                        | 0.35                          | 19.8 (4.8–60)                 | 1.1 (35.1)                      | 0.54             | 90 (22.9)       | 11.1           |

\(AUC\) area under the curve, \(AUC_{\text{inf}}\), \(AUC_{\text{last}}\) \(C_{\text{max}}\) maximum serum concentration, \(F_{\text{IM}}\) min minutes, \(\text{opioid}\) partly under opioid exposure, \(T_{1/2}\) half-life, \(T_{\text{max}}\) time to maximum concentration

\(^a\)Arithmetic mean (95% confidence interval)

\(^b\)Geometric mean (% coefficient of variation)

\(^c\)Median (minimum, maximum)

\(^d\)Arithmetic mean (standard deviation)

\(^e\)Geometric mean (90% confidence interval)

\(^f\)\(AUC_{\text{last}}\)

\(^g\)Absolute bioavailability

### Table 7 Factors affecting opioid reversal by naloxone

| Drug                  | Blood brain equilibration | Egress from brain | Dissociation rate | Terminal half-life |
|-----------------------|---------------------------|-------------------|-------------------|-------------------|
| Naloxone              | Rapid                     | Rapid             | Rapid             | Short             |
| Morphine              | Slow                      | Slow              | ?                 | Medium            |
| Methadone             | Rapid?                    | ?                 | ?                 | Long              |
| Fentanyl              | Rapid                     | Medium            | ?                 | Medium            |
| Buprenorphine         | ?                         | ?                 | Slow              | Long              |
up to 1 mg/kg have been tolerated, while 4 mg/kg was found to provoke undesired behavioural effects [117]. As noted above, naloxone also reverses skeletal muscle rigidity from fentanyl and its analogues [30, 31].

The most rewarding use of naloxone is in opioid overdose. Opioid overdose is characterised by stupor, impaired respiration and pin-point (miotic) pupils. Respiratory failure may lead to hypercapnia (also called hypercarbia) and hypoxemia, which may eventually be fatal. The best predictor of a successful response to naloxone in overdose is the miotic pupil [118].

6.3 Recurrence of Toxicity

Naloxone is a short-acting drug compared to the duration of action of many of the opioids [119]. In overdose, the duration of action of methadone, extended-release morphine, buprenorphine, extended-release oxycodone and fentanyl may exceed 12 hours. Thus, the effect of naloxone may wane before the respiratory depression by these opioids has ended [120]. In such cases, hospitalisation and repeat doses or continuous infusion of naloxone may be required [121]. Fortunately, the mortality rate from re-intoxication is low (<1%) [122, 123].

6.4 Acute Opioid Withdrawal Symptoms

In opioid-dependent subjects, administration of naloxone may produce acute onset of withdrawal symptoms such as agitation, nausea, vomiting, piloerection, diarrhea, lacrimation, yawning and rhinorrhea [116]. Tachycardia and hypertension are potentially serious circulatory effects, while violent behaviour and drug craving are far from trivial. Opioid withdrawal is not considered life threatening [124].

Buajordet et al. [125] conducted a prospective observational study in 1192 episodes of opioid overdose. Suspected adverse events from naloxone were reported in 45% of cases. Thirty-three percent had opioid withdrawal (gastrointestinal disorders, aggressiveness, tachycardia shivering, sweating and tremor), and 15% had headaches and seizures. Only 0.3% were hospitalised for an adverse event. The initial naloxone doses administered were 0.4–0.8 mg IM combined with an IV dose of 0.4 mg.

6.5 Dosing Guidelines for Opioid Overdoses

Recently, a comprehensive review paper (recommended for the interested reader) on naloxone dosing for opioid reversal was published [28]. The initial doses of naloxone commonly recommended for overdose are 0.4–2.0 mg IV or IM [124], with doses surpassing 0.8 mg potentially increasing the risk of significant withdrawal symptoms [9]. The duration of action of naloxone is 20–90 min depending on dose and the situation of the patient [119]. As reported above, a duration of action of about 90–120 min of 1 mg of naloxone was found in healthy volunteers and opioid-dependent individuals [95, 116]. Doses should be titrated every 2–3 min according to response for a total dose of up to 10 mg.

6.5.1 Special Populations

Care should be taken in the elderly, patients with preexisting cardiovascular disease and in those receiving potentially cardiotoxic drugs. The limited available data on naloxone use in pregnant women are not sufficient to inform a drug-associated risk. However, there are risks to the foetus of the opioid-dependent mother with the use of naloxone (Evzio summary of product characteristics). Children (aged 5 years and below) require relatively higher doses such as 0.1 mg/kg because of more severe intoxications [119]. However, lower initial doses (0.01 mg/kg) are also recommended, possible because opioid withdrawal may be life threatening in neonates.

6.5.2 Dosing of ‘Take Home Naloxone’

The situation is different when it comes to THN. The advantage of THN is that naloxone can be administered at an earlier stage, prior to ambulance arrival, similar to lay-person use of defibrillators in cardiac arrest. At present, the naloxone dose required in this situation is controversial (see below).

6.5.3 Naloxone Dosing for Fentanyl Overdoses

Severe intoxications of long duration resulting from swallowing fentanyl-adulterated hydrocodone/paracetamol (acetaminophen) tablets were reported [121]. One of the 18 overdose victims died. Seventeen required naloxone boluses, and four needed a prolonged infusion (26–39 hours) of naloxone.

Examination of about 95,000 cases in the national EMS data has also documented an overall increase in the severity of US opioid overdoses [126]. The use of multiple naloxone doses increased from about 15% in 2012 to 18% in 2015 [126]. In contrast, Bell et al. [127] found that systemic dosing of 0.4 mg naloxone (IM) did not increase from 2013 to 2016, despite an increase in fentanyl-related deaths from 4 to 69% in the same period. More than 90% of the reversals were successful after two naloxone doses (0.8 mg).

As for the intoxications resulting in immediate deaths, only anecdotal reports are available on the effect of nasal naloxone on fentanyl overdoses. Somerville et al. [27]
maximum concentration is slower and dose-corrected given that for dilute IN formulations (≤ 1 mg/mL) the time to one was not inferior to IM. This is somewhat surprising,

paper (2-mg/2-mL formulation). The safety of IN nalox-

compared to 78% and 72% reported in the Kerr et al. [128] (2-mg/5-mL formulation; 82% vs. 63% successful reversals),

study by Kelly et al. [129], IM did somewhat better than IN

comes in these studies were similar, with adequate respira-

tion and Glasgow Coma Scale response at 8–10 min. In the

initial dose of 2 mg IM with a 2-mg IN dose. Primary out-

comes in these studies were similar, with adequate respira-

tion and Glasgow Coma Scale response at 8–10 min. In the

study by Kelly et al. [129], IM did somewhat better than IN
(2-mg/5-mL formulation; 82% vs. 63% successful reversals),

compared to 78% and 72% reported in the Kerr et al. [128] paper (2-mg/2-mL IN formulation). The safety of IN nalox-
one was not inferior to IM. This is somewhat surprising,
given that for dilute IN formulations (≤ 1 mg/mL) the time to
maximum concentration is slower and dose-corrected C_max/
AUCs are generally lower.

6.6.2 Observational Studies

A retrospective study from Boston, USA, examined the use
of nasal naloxone (2 mg/2 mL by MAD syringe) by basic life
support providers [130]. After dosing, patients were trans-
ported to an emergency department. Nasal naloxone reversed
overdose in 95% of 793 patients, and only 9% required addi-
tional naloxone. Ninety-seven percent of patients had taken heroin, 3% had taken other opioids.

In another retrospective study, 2 mg/2 mL of nasal nalox-
one (MAD syringe) was administered by law enforcement
officers and basic life supporters. The overdose victims

(\( n = 2166 \)) were subsequently managed by paramedics [131],
and 9% needed at least one additional naloxone dose.

Despite somewhat different follow-up procedures in these
studies, low rates for re-dosing (9%) were reported. This
indicates that nasal systemic doses equivalent to 0.2–0.4 mg
IV/IM [74, 82], which may be below the lowest World
Health Organization-recommended starting dose of 0.4 mg
[9], seem both safe and effective in most circumstances
when given in medical settings (in contrast to layperson
administration).

6.7 Safety of Approved Nasal Formulations

As the basis for approval of naloxone nasal sprays, the
FDA requires that any new formulation must prove that it
obtains a serum concentration similar to, or higher than the
0.4-mg IM reference in healthy volunteers [132]. It should
be noted that there is a discussion within the FDA whether
the comparator dose should be increased [133]. The focus
of the FDA is on the first 10–15 min after administra-
tion. To achieve this goal, owing to the somewhat slower
IN uptake, a nasal formulation must systemically deliver
relatively more naloxone than IM leading to a considerable
“overshoot” of the C_max and the AUC (i.e. amount entering
the systemic circulation). This carries a significant risk for
naloxone overdosing that may provoke withdrawal symp-
toms. Although opioid withdrawal is not considered fatal,
it is not trivial. Experiencing withdrawal is feared among
people who use opioids. Qualitative interviews in Scottish
and US cohorts of PWUO identified negative views on
naloxone administration in the emergency room and the
harm caused, such as acute withdrawal, aggression, self-
discharge and further drug seeking activity [134, 135].
These attitudes were missed by ordinary observational
studies. Indeed, these behaviours and further drug seeking
activity perhaps constitute behaviourally mediated toxic-
ity [136].

The relatively high success rates reported by first respond-
er using one or two sub-therapeutic doses of naloxone (sys-
temic equivalents of about 0.2–0.8 mg) raises the possi-
bility that many overdoses may be over-treated with the new
approved devices delivering naloxone in the upper recom-
mended range (e.g. to about 2 mg by injection). In a report
concerning Narcan® (4 mg), 74 of the 196 reported cases
of community use had events that may, but not necessarily,
conform to opioid withdrawal; none were life threatening
[137]. If an intervention turns out to be labelled “withdrawal
hazard” by the user community, this may well affect their
decision to use the product.

Moreover, in their review of new naloxone products, reg-
ulatory bodies did not consider that characteristics of the
nose may differ in the target population compared to healthy
volunteers or the unexpected observation by Skulberg et al.
of a higher nasal bioavailability under opioid infusion. However, we know that all the approved nasal sprays deliver far higher systemic doses than doses supplied by the unapproved dilute nasal sprays. This should secure a very broad safety margin. The nasal (systemic) doses given in the ambulance-based randomised controlled trial in Australia were far lower than the IM comparator [128], yet the two groups performed similarly, except that about one in five overdose cases needed “rescue naloxone” (i.e. a second naloxone dose, this time by injection) following non-response to improvised naloxone nasal spray (2 mg/mL). As a moderate “overtreatment” with naloxone is hard to document, it can be argued that the comparator dose was higher than required for overdose reversal. This complies well with the observational studies that demonstrated unexpected high success rates, also in fentanyl overdoses. Therefore, the new purpose-developed concentrated naloxone nasal sprays which are being approved (dose range 1.4–4.0 mg), all with an option for titration with an accompanying second dose, are likely to offer a higher margin of safety with opioid overdoses, even in the era of illicit fentanyl. Nevertheless, the suitability of the different products will likely depend on the patterns of opioid use in a community, with higher strength formulations potentially needed for the reversal of overdoses from potent synthetic opioids (e.g. fentanyl and its analogues). Last, ventilator support given as rescue breathing is the cornerstone of opioid-induced respiratory failure.

As stated above, there are basically no safety data on the approved products. Pharmacovigilance may answer some of the important questions related to the new products. There is an urgent need for high-quality randomised comparisons of IM vs. IN naloxone administration in overdose. Moreover, observational studies independent of industry examining the efficacy of the new potent nasal naloxone sprays in overdose are warranted. Any consideration of significantly higher initial doses (than those that worked in the past) should be based not on anecdotes but on hard facts.

7 Reviewing the Effectiveness of Take Home Naloxone: Application of the Bradford-Hill Criteria

Twenty years after the 1996 BMJ editorial in which the notion of THN was first mooted [57], it remained unclear whether THN programmes would reduce the rate of opioid overdose deaths. The absence of published data from randomised controlled trials meant that uncertainty regarding the potential public health benefit of THN programmes persisted, leading to implementation delays and lack of political support in many jurisdictions. A small meta-analysis [138] had found a significantly increased odds of recovery from overdose for naloxone administration by bystanders compared with no naloxone administration (odds ratio 8.58, 95% confidence interval 3.90–13.25), but this analysis was only based on nine studies and 66 overdose events, and hence likely not representative of the majority of THN programmes.

To address this evaluation gap, a systematic review [139] was carried out to assess the effectiveness of THN, in terms of impact of opioid overdose mortality. Because of a lack of randomised controlled trials and a wide range in the methodological quality of the research studies, a meta-analysis was dismissed in favour of an analysis using the Bradford Hill [140] criteria: [1] Strength of the Association, [2] Consistency, [3] Specificity, [4] Temporality, [5] Biological Gradient, [6] Plausibility, [7] Coherence, [8] Experiment and [9] Analogy.

Based on their analysis, the authors [139] concluded that THN programmes lead to improved survival rates among programme participants and to reduced overdose mortality rates in the community. A later publication, which also based its analysis on the Bradford Hill criteria, reached a close-to-identical conclusion [141].

8 Barriers to Wider Implementation

Successful implementation of THN would result in widespread acceptability and availability of the intervention, such that it is accessed by relevant target populations and embedded in usual practice in drug treatment and mainstream health services. A number of papers have explored legal [142] and workforce [143, 144] barriers. These have been country and region specific. To date, there has not been a consolidated analysis of barriers to effective THN implementation. The first reason is that there is not a singular model of THN. The second reason is that a comprehensive analysis of implementation barriers requires a conceptual framework.

For the purposes of the analysis here, we refer to two broad types of models of THN implementation. Peer-led (PL) models were initially developed by and for PWUO, to provide training and supply naloxone directly to people who are likely to witness overdoses [138, 145–147]. These PL models are provided through organisations oriented to the needs of PWUO. In contrast, mainstream healthcare practitioner (MHCP) models harness medical services (doctors and pharmacies) to supply naloxone to patients and carers. Despite some overlap, they are very different approaches, involve different types of negotiation, and bring different advantages and disadvantages. They also vary in their suitability for different countries or states at different points in time.

Here, we use Greenhalgh et al.’s conceptual framework [148] to consider determinants of THN implementation.
across the two models. Based on the framework, achieving scaled-up implementation of THN involves considerations regarding the features of the innovation (such as efficacy, product, compatibility); the outer context (including socio-political climate, legal regime and policies, incentives and mandates); the system antecedents and readiness for change (such as organisational structures); the role of individual adopters (such as match with needs, motivation); communication and influence (such as social networks, champions); and the implementation process itself (formalised decision making, logistics and costs). This framework is not a prescription for successful diffusion of innovation, but an aid to considering the multiple aspects, behaviours, and determinants and their interactions.

8.1 Innovation Itself

Several determinants of successful implementation reside in the innovation itself, including the nature of the naloxone product. The injectable naloxone products present some barriers, with concerns including those relating to needle-stick injuries [76]. The newer IN formulations represent a significant advance in the product formulation and address previous concerns about IN bioavailability [11, 92, 149]. With recent reports of synthetic fentanyl not being responsive to usual doses of naloxone [150], we should also not be complacent about efficacy issues (please consult the THN safety discussion, Sect. 6.7).

The prescription status of THN is a significant feature impacting on implementation. Prescription products almost always require medical scripting and pharmacy dispensing and a trained, resourced and supported workforce (see below). Availability via OTC [151] and standing orders [152] address some barriers but not without other barriers arising such as workforce, the potential cost to patients (in some countries) and access [152, 153]. Currently, there are also major issues with stock and supply in many countries including the USA, UK and Australia [153, 154].

Implementation success is also subject to the compatibility of the intervention with the groups delivering it. In theory, THN is highly compatible with the values, norms, and needs of peers and carers (saving friends and loved ones), as well as healthcare professionals (reduced mortality). Yet for both PL and MHCP, there may be some incompatibility with goals. For the PL models, a recently published qualitative study of the views of PWUO highlighted the complex judgement about when to use naloxone given concern about “ruining someone’s high” (p. 34) [155]. There was no consensus on defining the line between a good high and an overdose [155]. For MHCP, there are concerns that providing such a life-saving intervention may encourage less safety in drug use [156–158]. Heavey et al. [155] found that some participants reported that naloxone access changed the quantity and frequency of use: “chase a bigger high” (page 32; but see also [159]). A recent working paper [156] that suggested a ‘moral hazard’ problem with THN provoked a furor, although most commentators consider the analysis weak and conclude that the evident logic and benefit of THN remain clear [160, 161]. Nevertheless, the intensity of debate from this paper reveals that there is more work to do in ensuring that THN is seen as compatible with health caring roles, as well as managing the perceived risks associated with THN.

8.2 Outer Context

The second domain of implementation in Greenhalgh et al.’s framework is the “outer context” including the socio-political climate and policy context. The legal frameworks are important in introducing and sustaining THN, including laws concerned with indemnity for consumers (from being charged with offence), and medical personnel (prescribers/ dispensers) immunity from being sued [142, 162–165]. The legal barriers differ between PL and MHCP models. The extent to which these legal regimes are directly associated with increased uptake has yet to be empirically demonstrated [166]. Nonetheless, a conducive legal environment is a prerequisite for widespread implementation.

A key barrier within the outer context domain is the broad stigma and marginalisation of people who use opioids. The general community largely has negative attitudes towards PWUO [167, 168], and this is compounded by internalised stigma [169].

The outer context in the Greenhalgh et al. framework also covers system incentives and mandates. These are the formalised policies and procedures that regulate and support THN programmes. While international documents reference the importance of THN [9, 170], beyond these, there appear to be few formal organisational incentives, directives and mandates in place currently. Mandatory co-prescribing of naloxone with opioid prescriptions has been proposed to address prescription opioid overdoses, and pilot studies have tested this in clinical practice, finding it feasible and associated with reduced opioid-related adverse events [171].

8.3 System Antecedents and Readiness for Change

In Greenhalgh et al.’s framework, both system antecedents and readiness for change are key determinants of implementation success. A receptive context for change, with leadership and vision by the medical profession and by drug user organisations is a pre-requisite for success. Drug user organisations are often under-resourced, and specific funding to provide PL programmes is required. For the MHCP model, the systems and structures for prescribing and dispensing naloxone are certainly well developed. However,
barriers remain with respect to adoption by the workforce (see Sect. 8.4).

8.4 Adopters

There are three essential groups of “adopters” whose needs, motivations, values and goals need to align with the intervention to ensure effective widespread availability of THN: people with an opioid-overdose risk; peers and carers; and healthcare professionals. One barrier for potential adopters is stigma, including negative perceptions of PWUO by healthcare providers [172], as well as the stigma of being identified as a person who may use drugs [173]. Stigma reduction programmes targeted at the potential adopters (as well as the general community, see Sect. 8.3) may be required [174–176]. The universal supply of naloxone to all with a risk of opioid overdose as part of standard care (and with a patient ‘opt out’ model [177]) has been proposed to address stigma, and also address clinicians’ duty of care where overdose risk is identified. The cost effectiveness of universal supply models is unknown.

There are also significant barriers in relation to knowledge and skills including the need for education and training of potential healthcare providers to enable adoption of THN in the MHCP model [144, 178]. While PWUO often have considerable knowledge of overdose signs and symptoms, knowledge gaps exist amongst different populations with opioid-overdose risk [155, 173, 179].

8.5 Implementation Processes

The process of implementation presents two key barriers here for THN: the workflow logistics and cost. Healthcare professionals have limited time, and THN needs to be incorporated into their workflow, including how to get the THN kit to the patient and the timing of this [152]. There is an opportunity cost for time-pressured healthcare professionals, where training patients on naloxone use may leave less time for other needs. For general medical practitioners and pharmacists, this can be compounded by the systems that pay for their time. For PL models, significant resources are required to deliver training programmes, which peer organisations may not have.

A further substantial implementation barrier is cost, which relates to the medication itself and who is expected to pay. As a prescription medicine, the costs may be covered centrally as part of regular treatment, or subsidised through private health insurance plans, Medicare and Medicaid [142] and/or through grants to harm reduction programmes to purchase THN and then provide it free of charge to clients. In Australia, a script for THN will be affordable to a concession card holder, but OTC (i.e. not subsidised) supply significantly increases the cost to the consumer [153]. Thus, while new supply routes (such as OTC) and formulations (IN) reduce some barriers, they simultaneously present other barriers (in these cases, the cost).

8.6 Communication and Influence

At present, THN largely remains an isolated initiative led by individual champions, harm reduction advocates or health professionals experienced in drug dependence. For it to become a mainstay of usual care, much wider dissemination and diffusion of the technology need to take place. Here, the system variables discussed earlier become most important: widespread stigma against PWUO, the lack of systematic endorsement across all settings and the absence of wide access (such as OTC) that can be provided at low cost.

8.7 Understanding Barriers and Developing Integrated Responses

Across the analysis conducted here using one conceptual implementation framework, it becomes clear that THN implementation has barriers related to the intervention itself (notably the product and its prescription status), the outer context (stigma and a lack of formalised policies and procedures), the adopters (in particular attitudes, knowledge and skills of adopters), and in the practicalities of implementation, especially the cost and the logistics. These determine the extent of THN diffusion such that it becomes a part of regular practice.

9 Conclusions

9.1 Potentially Transformative Public Health Intervention

Naloxone is a well-established emergency medicine for the treatment of life-threatening heroin/opioid overdose. In recent years, with interest in technology transfer, the concept of ‘THN’ has evolved, comprising pre-provision of the supply of naloxone to individuals who may be in the position to intervene and provide essential interim care whilst awaiting the arrival of an ambulance.

9.2 Better Naloxone Products Required for More Effective Take-Home Naloxone

New naloxone products (auto-injector and nasal spray) have been developed in the last few years that have good bioavailability, can provide therapeutic doses in a single step, and hence are suitable for use by lay responders and other
non-medical personnel. Four concentrated nasal spray products have now reached the stage of regulatory approval, with several already introduced in some countries globally: Narcan® 4 mg (USA, Canada, approved and introduced in 2016–2017), Nalscue® 1 mg (approved and introduced in France in 2017), Nyxoid® 2 mg (European Medicines Agency approval in 2017, introduced across much of Europe from 2018 onwards) and Ventizolve® 1.4-mg spray (developed in Norway and approved for 12 European countries in June 2018).

9.3 Elephant in the Room: Cost and Affordability

When comparing the different naloxone devices, including ampoules and vials as the most basic, there is a clear trade-off between usability and cost. With all products containing the same medication (naloxone), this point is perhaps best illustrated with the example of the FDA-approved auto-injector (Evzio®). At approximately US$600–4500 per unit, the auto-injector is at risk of being considered almost irrelevant for community-wide provision. New concentrated nasal sprays, if costs are kept low, might be particularly suitable for much wider public pre-provision. The cost of twin packs of the new approved naloxone nasal sprays varies between countries from approximately US$30 (equivalent) to several hundred dollars, compared with less than a dollar for ampoules vs. several thousand dollars for auto-injectors. The affordability of new naloxone products will likely define their real-world availability and accessibility for the target population.

9.4 Over-the-Counter Status

As discussed above (see Sect. 8), re-scheduling of naloxone to OTC medication could remove several logistical and legal barriers. However, the issue of THN costs overlaps with the issue of OTC status as high cost could render the product unattainable, even if technically available OTC. By analogy, in the prevention of sexually transmitted infections, cost is a barrier to condom use [180], and free-of-charge condom distribution to target high-risk populations is recommended [181, 182]. A dual implementation model is likely optimal for naloxone access; potential OTC sales should best be considered as operating alongside free-of-charge THN programmes.

9.5 Opioid Overdose Prevention in Low- and Middle-Income Countries

The question of the cost of non-injectable naloxone (vs. injectable forms of the same medicine) becomes particularly relevant for global health. With support from the Global Fund, several countries currently procure naloxone ampoules at the low cost of approximately US$0.50 per ampoule (0.4 mg/mL). Local non-governmental organisations typically hand out two take-home ampoules per client, i.e. equivalent to a total medicine cost of US$1.00 per person. Given the cost disparity between the nasal spray and local ampoule supply, the purchase of naloxone nasal spray is likely not sustainable for low- and middle-income countries at present. However, bulk buying and the production of generics could lower costs in the long term, as has been achieved for second-line antiretroviral therapy for human immunodeficiency virus/acquired immune deficiency syndrome in low- and middle-income countries, where medication costs have been dramatically reduced.

9.6 Need for Continued Research

Surprisingly little attention has been paid to study these new naloxone products with participants from the relevant target population of individuals with substance use disorders, hence uncertainty remains. Will damage to nasal membranes from long-term drug use alter the speed and effectiveness of absorption? Are the findings from healthy volunteers generalisable to overdose victims with potentially very low body weight, near-universal chronic smoker status or varying degrees of hepatic impairment? Is nasal bioavailability of naloxone higher under opioid influence? Can dose titration be taught effectively? What is a safe starting dose, the dose that together with a similar follow-up dose within 3 min that reverse most respiratory failures without provoking too much withdrawals? Such studies may be challenging to devise and conduct, but they would ordinarily be considered necessary steps with the development of new interventions with other clinical populations. There is also a need for stronger study designs to guide further improvements in THN provision.

9.7 Final Remarks

Take-home naloxone has established itself as a major new strand of our response to opioid overdose deaths. This has required new policies and practices, alongside new naloxone formulations and products. These have been developed successfully, and continue to be refined for new challenges, new

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5 A generic version of the 4-mg/0.1-mL Narcan nasal spray by Teva Pharmaceuticals also received FDA approval in April 2019 in the US but has not yet been introduced; and an application in the US from InSys for an 8-mg/0.1-mL naloxone nasal spray was accepted for consideration by FDA in July 2019.

6 A 2-mg version of Narcan® was also approved in North America but was never introduced in clinical practice—see footnote to Table 3.

7 See footnote to Table 3 for explanation of dose of Nyxoid.
countries and contexts, and for new opioid drug problems. More research is, of course, needed—with stronger research designs as well as the need for new products and new methods of provision; but this new research must be conducted alongside the introduction of the best products and practices that already exist. With a large and growing problem of heroin/opioid overdose deaths, we have a responsibility to expect the best of science, policy and practice to guide our local, national and global responses.

Compliance with Ethical Standards

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Conflict of interest John Strang, through his university, is working with the pharmaceutical industry to identify new or improved treatments and his employer (King’s College London) has received grants, travel costs and/or consultancy payments; this includes investigation of new naloxone formulations and has included work with in the past 3 years, Martindale, Indivior and Mundipharma (all of whom have naloxone products). His employer (King’s College London) has also registered intellectual property on a novel buccal naloxone formulation, naming John Strang et al., and he was earlier named in a patent registration by a pharmaceutical company regarding a concentrated nasal naloxone spray. John Strang et al. have worked as consultants for the United Nations Office on Drugs and Crime, supporting them with a project introducing take-home naloxone to four central and western Asian countries as well as contributing to local take-home naloxone schemes. For a fuller account, see John Strang’s web-page at http://www.kcl.ac.uk/iopnp/depts/addictions/people/hod.aspx. Rebecca McDonald has undertaken an unpaid student industry placement with Mundipharma Research Ltd. and has received conference-related travel funding and an honorarium from IOTOD (Improving Opioid Outcomes in the Treatment of Opioid Dependence). Rebecca McDonald has been involved in the development of a tablet formulation on which the university (King’s College London) has registered intellectual property. Rebecca McDonald has worked as a consultant for the United Nations Office on Drugs and Crime. Gabrielle Campbell is supported by a National Health and Medical Research Council fellowship (No. 1119992). The National Drug and Alcohol Research Centre at the University of New South Wales is supported by funding from the Australian Government under the Substance Misuse Prevention and Service Improvements Grant Fund. Gabrielle Campbell has received investigator-initiated untied educational grants from Reckitt Benckiser/Indivior for a study of opioid substitution therapy uptake among patients with chronic non-cancer pain. None of these are directly relevant to the current article. Louisa Degenhardt is supported by National Drug and Alcohol Research Centre research fellowships (No. 1109366 and No. 1041472/No. 1135991) and by the National Institutes of Health grant NIDA R01DA1104470. The National Drug and Alcohol Research Centre at the University of New South Wales is supported by funding from the Australian Government under the Drug and Alcohol Program. Louisa Degenhardt has received investigator-initiated untied educational grants from Reckitt Benckiser/Indivior for studies of buprenorphine-naloxone, buprenorphine depot, naloxone, the development of an opioid-related behaviour scale and a study of opioid substitution therapy uptake among patients with chronic non-cancer pain. Louisa Degenhardt has received an untied educational grant from Seprius for a post-marketing study of tapentadol and an untied educational grant from Mundipharma for a post-marketing study of oxycodone. These bodies had no involvement in or knowledge of the current article. Suzanne Nielsen has been a named investigator on untied educational grants from Indivior and Seprius and has delivered training on opioid dependence for Indivior for which honoraria were paid to her institution. Suzanne Nielsen has participated in an advisory board meeting by Mundipharma relating to naloxone for which she declined a sitting fee. Suzanne Nielsen is a current recipient of a National Health and Medical Research Council Research Fellowship (No. 1163961). Alison Ritter is funded through a National Health and Medical Research Council Research Fellowship (GNT1136944). She has received no funding or untied educational grants from any pharmaceutical company in the last 15 years. This article covers (among others) a naloxone formulation that has been approved by 12 European countries from 16 June, 2018 under the name of VentiZolve® (Respinal® in Sweden), produced by Sanivo Pharma AS. The sponsor of this trial was AS Den Norske Eterfabrikk. Ola Dale was engaged by AS Den Norske Eterfabrikk as the principal investigator in this study, for which Ola Dale receives no personal honorarium. Ola Dale’s employer, the Norwegian University of Science and Technology and its subsidiary Technical Transfer Office have signed cooperation and licensing contracts with DNE PHARMA to seek commercialisation of this nasal naloxone formulation. This regulates potential royalties for Ola Dale through the Norwegian University of Science and Technology. DNE Pharma AS has compensated Ola Dale for business travel from Trondheim to Oslo.

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