Moderate-to-severe Vipera berus envenoming requiring ViperaTAb antivenom therapy in the UK

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ABSTRACT

\textbf{Background:} Bites by the European adder (Vipera berus) in the UK are uncommon but potentially life threatening, and can be associated with marked limb swelling and disability. Following an interruption in Zagreb Imunološki zavod antivenom supply around 2012, the UK changed its national choice of antivenom for Vipera berus to ViperaTAb, an ovine Fab monospecific antivenom. In the absence of randomised controlled trials, we established an audit to review its use in clinical practice.

\textbf{Methods:} A prospective audit of ViperaTAb use was conducted from March 2016 until November 2020 by the UK National Poison Information Service (NPIS). Users of the NPIS online toxicology database, TOXBASE, considering the use of antivenom for V. berus envenoming were invited to discuss the case with the on-call clinical toxicology consultant. Information was collected prospectively on indications, administration, adverse reactions and outcome of patients administered ViperaTAb antivenom.

\textbf{Results:} One hundred and seventy patients were administered ViperaTAb antivenom over five years. One hundred and thirty-two were adults and 38 children (median age and range: 38, 2–87 years). Bites occurred across the UK, but most commonly in coastal regions of Wales and of South-West and East England. Median time to presentation was 2.1 (IQR 1.5–4.0) h and to antivenom administration from presentation was 2.0 (IQR 0.9–3.6) h. A minority of patients presented to hospital more than 12 h after being bitten (n = 19, 11.2%) or received antivenom more than 12 h after presenting to hospital (n = 17, 10.0%). Features of systemic envenoming were present in 64/170 (37.6%) patients, including 23 (13.5%) with anaphylaxis and 26 (15.3%) with hypotension (nine with both). Clinician assessment considered the initial antivenom to have been effective in 122/169 (72.2%) patients. Repeated dosing was common, occurring in 55/169 (32.5%), predominantly due to persisting or worsening local effects (46/51, 90.2%). There were three cases of probable early adverse reaction. No deaths occurred during the study. Complications of envenoming were rare but included four patients that underwent surgery, three patients each with acute kidney injury, mild coagulopathy, or thrombocytopenia (one severe). The median duration of hospital stay was 43.7 (IQR 22.5–66.5) h, longer for children than adults (52.5 vs 41.3 h).

\textbf{Conclusion:} ViperaTAb antivenom appears to be effective and safe and should be administered as soon as possible for patients meeting clinical criteria. Patients require close observation following antivenom to detect adverse reactions and progression or recurrence of envenoming. Close collaboration with expert NPIS consultant advice can help optimise antivenom timing, ensure repeated dosing is given appropriately, and avoid unnecessary surgical intervention. All hospitals, particularly those located in areas of relatively high incidence, should stock sufficient antivenom available at short notice, 24 h a day.

Introduction

The European adder (Vipera berus) is the only species of venomous snake native to the UK. Bites are uncommon but potentially life-threatening, with an estimated 50–100 cases occurring annually in the UK [1,2]. Of these, 20–60 cases are treated with antivenom. Envenoming is characterised by bite site pain, bruising, and swelling that is often pronounced and may affect the entire limb and trunk [3]. A minority of patients develop signs (or features) of systemic envenoming, the most concerning of which is a non-immune anaphylactic reaction to venom which develops soon after the bite with angioedema,
urticaria, hypotension and collapse. Other features of systemic envenoming include vomiting, diarrhoea, and less commonly cardiotoxicity, acute kidney injury, thrombocytopenia, and coagulopathy [3,4].

The Zagreb Imunološki zavod antivenom, raised against Croatian Vipera ammodytes venom [5] was used in the UK for many years [1] before an interruption in production around 2012 [6]. This resulted in a change in choice of national antivenom from Zagreb to ViperaTAb (MicroPharm) antivenom for moderate-severe Vipera berus envenoming.

ViperaTAb is a monospecific ovine Fab antivenom raised against whole V berus venom (sourced from Russia) manufactured by MicroPharm Ltd, UK. Fab antigen binding complexes are smaller than F(ab’)2 or whole IgG fragments providing the theoretical advantage of enhanced tissue penetration and fewer adverse reactions. However, the increased volume of distribution and shorter half-life of Fab fragments increases the risk of recurrent local and systemic envenoming after antivenom administration [7]. Recurrent venom antigenaemia with coagulopathy after treatment with an Fab antivenom was first demonstrated in Echis ocellatus/romani envenoming in Nigeria [8] and subsequently in a comparative trial of Fab and F(ab’)2 antivenin for Crotalinae envenoming in the USA [9].

As is the case for all antivenoms, the same initial dose of ViperaTAb is given to children and adults. One dose consists of two vials (8 mL, 200 mg), diluted in normal saline and infused intravenously (IV) over 30 min. The preclinical evaluation and 50% binding value for ViperaTAb is superior to that of Zagreb antivenom for V. berus venom [10].

There have been no randomised controlled trials of ViperaTAb [5]. Published observational clinical data from 645 patients in Scandinavia, where ViperaTAb has been in use since 1991 [11–18], suggest efficacy with quick resolution in features of envenoming, comparatively short length of hospital stay, and fewer adverse reactions than with Zagreb antivenom [5,19].

Following the change in national antivenom supply, the UK’s National Poison Information Service (NPIS) prospectively audited the indications, administration, adverse reactions, and outcomes of patients receiving ViperaTAb in the UK over a period of five years.

Methods

A prospective audit was conducted by the NPIS, the UK’s national poison control centre, of all V. berus snakebites from 1 March 2016 until the last case in 2020 (06 November 2020). Records of calls to NPIS are routinely collected into a common database, the UK Poisons Information Database (UKPID).

Clinicians accessing the UK toxicology database TOXBASE for advice on the treatment of V. berus envenoming were prompted with a Pop-Up box to telephone the NPIS and discuss management with the toxicology consultant on call if they were considering antivenom. Demographic data and envenoming details were routinely recorded during these conversations. Advice on antivenom administration (performed under close observation due to the possibility of severe adverse reactions) was provided based on the clinical criteria set out in TOXBASE (Box 1). Formal gradation was not undertaken. Timing of bite, presentation to hospital, antivenom administration and discharge were used to ascertain time to antivenom administration and length of hospital stay. Follow-up telephone calls to support management were made by one consultant. Clinicians were asked whether they considered the antivenom to have been effective i.e., whether it resulted in rapid resolution of systemic features and/or cessation in spread of local oedema and whether any early adverse reactions had occurred.

Box 1. UK national guidelines for antivenom administration for Vipera berus envenoming (TOXBASE, NPIS).
Antivenom should be given if the patient has any of the following features:
1. Early anaphylaxis-like reactions to the venom.
2. Hypotension persisting for more than 10 minutes, with or without features of shock.
3. Systemic features including abdominal pain or diarrhoea.
4. Definite leucocytosis (especially if over 20 × 109/l).
5. ECG abnormalities (e.g., bradycardia or widespread ischaemia).
6. Metabolic acidosis.
7. Elevated creatine kinase.
8. Severe local envenoming (even in the absence of systemic features) i.e., swelling spreading beyond the next major joint.
9. Any other evidence of systemic envenoming e.g., spontaneous haemorrhage, pulmonary oedema.

Anonymous data were extracted from data recorded in UKPID. Statistical analysis was performed on GraphPad Prism, Version 8. Data were described using median and interquartile range (IQR). Categorical data were compared using non-parametric statistical tests with a p value of <0.05 considered significant.

This audit did not require approval by a UK Research Ethics Committee as it used information collected routinely as part of usual clinical care, with information provided for analysis in a fully anonymized format.

Role of the funding source

The funders of the study had no role in study design, data collection, data analysis, data interpretation, or writing of the report. The corresponding authors had full access to all the data in the study and had final responsibility for the decision to submit for publication.

Results

One hundred and seventy patients were identified as having received ViperaTAb for snake envenoming during the study period (average of 34 cases/year; range 31–40).

A further 57 cases (seven children) of envenoming who presented to hospital (2 in 2016, 9 in 2017, 14 in 2018, 17 in 2019, 15 in 2020) and were discussed with NPIS did not receive antivenom (because they had only mild local effects not requiring antivenom [36/57, 63.2%], presented too late.
usually after 48 h \([17/57, 29.8\%]\), had an unclear diagnosis on initial presentation \([3/57, 5.3\%]\), or because antivenom was not available \([1/57, 1.8\%]\)]. Of note, five of the patients presenting late, and one of the patients presenting with mild envenoming, were adults who had had early systemic features but had gone to bed rather than coming to hospital. They presented after 1–3 days when their features of their systemic envenoming had settled, and local signs did not require antivenom. These cases are not further discussed here.

**Bite occurrence**

Envenoming occurred predominantly over the summer months with the first case on 26 February and the last case on 06 November (Figure 1). Envenoming occurred most frequently on weekends between the hours of 12.00 and 18.00 h (Figure 1).

Cases had a predominantly coastal distribution with clusters occurring in West Wales and in South-West and East England. Bites often occurred on coastal paths or beaches, or in forests/heaths. The hospitals that treated most patients with antivenom were located in Bangor \((n = 11)\), Bournemouth/Poole \((n = 11)\), Truro \((n = 10)\), Swansea \((n = 9)\) and Exeter \((n = 8)\) (Figure 2). Two patients were envenomed in Sweden but travelled to the UK before seeking medical attention. The snakes were not always seen, sometimes delaying the presentation to hospital and/or the diagnosis.

**Patients and clinical features on presentation**

The median age of patients was 38.0 years (IQR 17.5–53.5) with a male predominance \((108 \text{ male}, 62 \text{ female})\) (Figure 3). There were 38 children \((\text{range} 2–15 \text{ years})\), 13 of whom were less than five years old.

There was a marked difference in bite site for women and men. In men, 74 bites \((68.5\%)\) were on upper limbs, particularly the right hand, while 33 \((30.6\%)\) occurred on the lower limbs. By contrast, only ten \((16.1\%)\) occurred on the upper limbs of women, while 51 \((82.3\%)\) occurred on the lower limbs \((p < 0.0001)\) (Figure 4). There was one bite on the face and one bite on the abdomen.

Patients generally arrived quickly at hospital – median time to presentation was 2.1 (IQR 1.5–4.0) h, although 19 cases \((11.2\%)\) presented to hospital more than 12 h after the bite. At least four patients were transferred by air ambulance, either from rural areas or from the west Scottish islands to the mainland.

![Figure 1](image1.png)  
*Figure 1.* Time of day (median and IQR), day of the week, and monthly distribution of 170 bites requiring antivenom.

![Figure 2](image2.png)  
*Figure 2.* Location of hospitals treating envenomed patients.
All but two patients presented with localized swelling, with or without pain. Twenty-three (13.5%, including two children) presented with a non-immune anaphylactic reaction, attributable directly to *V. berus* venom characterized by angioedema, urticaria, and wheeze (Table 1). Twenty-six (15.3%, four children) had hypotension, nine with other features of anaphylaxis and 17 without. Of the 41 patients with anaphylaxis and/or hypotension, 11 required bolus intramuscular (IM) or IV adrenaline, usually during transfer, before antivenom could be accessed. Other features of systemic envenoming included vomiting (32) and diarrhoea (14) (Table 1). A total of 64 patients (37.6%, 14 children) presented with systemic features. The incidence of systemic features including anaphylaxis and hypotension was similar for children, adults aged 16–64 years old and adults aged >64 years old (Table 1).

Two patients were intubated and ventilated: due to bronchospasm and hypoxia in an asthmatic adult while antivenom was being located and prepared, and due to airway compromise following a facial bite to a child. Six patients were noted to have transient ECG abnormalities attributed to the envenoming (ST changes [2], left axis deviation [1], prolonged PR interval [1], prolonged QT interval [1] and unspecified abnormality [1]). Although laboratory values were not systematically documented, leucocytosis and elevated D-Dimer were commonly reported.

**Antivenom administration**

Median time to antivenom administration from admission was 2.0 (IQR 0.9–3.6) h – a median of 4.6 (2.8–9.4) h after the bite (Figure 5). However, seventeen cases (10.0%) received their first dose of antivenom more than 12 h after presenting to hospital. In six patients, antivenom was either delayed or staggered due to insufficient antivenom stock at the initial hospital/health care centre presentation. Envenoming progressed while in the hospital for other patients, leading to the need for antivenom.

The appropriate initial dose of two vials (200 mg) was given to 168 patients. One patient, an 8-year-old girl, received half a dose (1 vial) with the second vial administered 5 h later; the second patient, a 45-year-old woman, received half a dose (1 vial), responded sufficiently and did not require further dosing. Of the 169 (99.4%) patients for whom follow up was completed, 43 (25.4%) received one additional dose and 12 (7.1%) received two additional doses. Overall, 55/169 (32.5%) received additional antivenom dosing, giving a total of 235 administrations.

The reason for repeated dosing of antivenom was identified for 51/55 patients. Persistent systemic features were responsible in five patients (coagulopathy (n = 2), arrhythmia and cardiovascular instability (1), thrombocytopenia (1) and hypotension (1)). Forty-six patients received additional dose(s) due to worsening or persistent local effects (swelling, less often haematoma, bite site necrosis). For just two patients, the repeated dose was not clinically indicated according to NPIS consultant review.

Median time to the second dose from the first dose was 15.2 (IQR 6.1–24.5, range 0.8–70.2) h and from presentation was 16.7 (8.0–26.9, range 1.5–70.2) h (Figure 5). Patients administered a second dose of antivenom for persistent or worsening local envenoming <6 h after the first dose (n = 16) had a shorter length of hospital stay (IQR 52.6 (47.1–69.0)) h than those who received a second dose >6 h after the first dose (n = 29) (71.0 (46.9–98.4)) h (p = 0.08).

The latest a patient received antivenom was 67.0 h after the bite and 47.5 h after presentation. The reasons for the very late antivenom administrations (>36 h after the bite) were delays to presentation to hospital (n = 3), misdiagnosis as infection (1), initial discharge followed by evolution of cytotoxicity (1) and self-discharge and re-presentation (1).

Assessment of effectiveness was attempted by asking clinicians whether they considered the antivenom to have been associated with improvement in their patients. They deemed the initial antivenom to have been effective in 122/169 patients (72.2%), generally with resolution of systemic features and/or a rapid halt in the spread of swelling up the limb. Nineteen of the 122 cases (15.6%) in which an initial response was judged to have been favourable subsequently required at least one additional dose of antivenom. One patient required redosing of antivenom for systemic envenoming (hypotension), 6 h after the initial dose. The remaining eighteen patients received an additional dose of antivenom due to worsening or persistent local effects at a median of time of 22.8 (IQR 7.1–29.1) h after the initial dose.

**Safety of ViperaTab**

There were three cases of probable early adverse reactions to antivenom: two of hypotension and one of urticaria tracking up.
the line of the arm vein receiving the antivenom. The hypoten-
sion resolved without treatment while the patient with urticaria
was given adrenaline and hydrocortisone. Each of these
patients received the dose of antivenom in full and at least one
further dose of antivenom without reaction.

There were three possible early reactions to antivenom: one
case of worsening facial oedema (already present before anti-
venom therapy), one case of profuse diarrhoea, and one case
of abdominal pain and retching. Each resolved without spe-
cific therapy.

Three patients were noted to develop thrombocytopenia
(nadir platelet counts of 145, 88 and $5 \times 10^9$/L). The lowest
count occurred in a child with a platelet count of 35 on
admission, before antivenom administration. The first two
cases were managed conservatively without platelet infusion;
the final case was treated with 1 unit of platelets and two

| Table 1. Systemic features related to age in patients envenomed by V berus subsequently treated with antivenom. |
|---------------------------------------------------------------|
|                                                                 |
| Number of cases | 38 | 119 | 13 | Fisher exact between child/adult < 65 |
| Anaphylaxis     | 2 (5.3%) | 20 (16.8%) | 1 (7.7%) | $p = 0.11$ |
| Hypotension     | 4 (10.5%) | 21 (17.6%) | 1 (7.7%) | $p = 0.44$ |
| Diarrhoea       | 2 (5.3%) | 11 (9.2%) | 1 (7.7%) | $p = 0.73$ |
| Vomiting        | 11 (28.9%) | 20 (16.8%) | 1 (7.7%) | $p = 0.10$ |
| Any systemic feature(s) | 14 (36.8%) | 46 (38.6%) | 4 (30.8%) | $p = >0.99$ |

NB. Each patient may have had more than one clinical feature of systemic envenoming.
further doses of antivenom as a decision was made that the thrombocytopenia was likely due to the venom, and not the antivenom.

**Complications of envenoming**

No deaths occurred during the study. Complications of envenoming are listed in Table 2. Four patients underwent surgery (necrotic digital ulcers debridement \( n = 2 \), incision and drainage of a haematoma in a patient on an anticoagulant when envenomed \([1]\), and fasciotomy for suspected compartment syndrome (healthy muscle was found at operation) \([1]\)). Five developed metabolic acidosis, three acute kidney injury, three mildly deranged clotting function, and two mild transaminitis. Two patients were noted to have microscopic haematuria on dipstick urinalysis, while three (noted above) had thrombocytopenia that may have been due to venom. All laboratory abnormalities improved prior to discharge from hospital.

**Length of hospital stay**

Median duration of hospital stay \( (n = 169, \text{99.4\%}) \) was 43.7 (IQR 22.5–66.5) h. It was longer for patients aged >64 yrs (67.2 [IQR 26.2–122.3] h), compared to children aged <16 yrs (52.5 [IQR 39.4–91.0] h), and adults aged 16–64 years old (38.8 [IQR 20.7–53.3] h) (Kruskal–Wallis \( p = 0.004 \) (Figure 6).

Eight patients remained in hospital for seven or more days (Figure 6). The reasons for these prolonged admissions were rehabilitation due to extensive local effects \( (n = 4) \), suspected compartment syndrome managed conservatively \( (2) \), incision and drainage of haematoma formed at the bite site in a patient on rivaroxaban \( (1) \), and fasciotomy for suspected compartment syndrome \( (1) \).

Table 3 presents five cases of envenoming that illustrate a spectrum of severe local and systemic envenoming, the importance of early antivenom and importance of compartmental pressure assessment. The inappropriate prioritisation of surgical assessment over antivenom therapy was observed in several patients.

**Discussion**

This prospective study of ViperaTab supports Scandinavian experience that the antivenom is safe and appears effective for *V. berus* envenoming. The median length of hospital stay (a useful marker of antivenom efficacy given the predominant local effects of envenoming) of 43.7 h was similar to that in a study of *V. berus* and *Vipera aspis* bites treated with a French viper antivenom, ViperFAV, and shorter
than reported for cases receiving the Zagreb antivenom \[5,19,20\].

Patients requiring antivenom typically presented to hospital, were assessed, and received antivenom in a timely manner, often within 4 h of the bite. However, there were delays in presentation and antivenom administration in a few cases. Delays to antivenom after arrival at hospital were sometimes explained by evolving clinical features. However, on other occasions antivenom was delayed due to the limited experience of the clinicians involved, with a sense that orthopaedic review rather than antivenom was the priority. There were delays to antivenom administration in six patients due to limited stock at the presenting hospital. Delays to antivenom therapy for \textit{V. berus} and \textit{V. aspis} envenoming has been found to be associated with increased risk of haematoma formation and prolonged functional impairment \[21,22\].

Seventeen patients delayed their presentation to hospital until it was considered too late (after 48 h) for benefit from antivenom. Scanning of newspaper articles revealed other patients who chose to not attend hospital, with clear consequences for the functional status over the coming weeks.

Table 3. Illustrative cases of severe envenoming and learning points.

| Case | Time from bite to ViperaTAb (h) | Time to discharge from presentation (h) | Illustrative point | Clinical details |
|------|-------------------------------|----------------------------------------|--------------------|-----------------|
| 1    | 21.0 and 66.9                 | 254                                    | Importance of early antivenom. | A female patient developed a sharp pain on her foot whilst walking (unwitnessed bite). She presented to hospital with a diffusely tender, swollen leg, leucocytosis (25 × 10⁹/mL) and metabolic acidosis. Snakebite was considered as a differential diagnosis in the emergency department; however, the patient was given IV antibiotics and referred to orthopaedics for assessment of possible necrotizing fasciitis and/or compartment syndrome. No surgical intervention was required. Antivenom was administered 17.5 h after presentation resulting in an improvement in clinical features and biochemistry; however, due to the substantial local effects, the patient required a prolonged period of rehabilitation. |
| 2    | 4.8                           | 212.7                                  | Importance of compartmental pressure assessment. | A male child presented following a bite to the right foot. Antivenom was given for local effects with no evidence of systemic features. It was adjudged to have been effective and no further doses were administered. The following day, further swelling of the affected limb was noted. He underwent fasciotomy for suspected compartment syndrome without prior radiological or compartmental pressure assessment. In theatre, the muscle was found to be swollen but pink and healthy. |
| 3    | 3.2, 11.7 and 24.0            | 242.9                                  | Severe systemic envenoming with anaphylaxis. Importance of compartmental pressure assessment. | Following a bite to the left hand, a male patient presented with perioral paraesthesiae, lip swelling, a sensation of “tight throat” and swelling to the elbow. He was administered IM adrenaline and ViperaTAb, resulting in resolution of systemic features. Over the next 24 h, oedema progressed to the upper arm and onto the thorax despite a further dose of antivenom. Due to clinical concern about compartment syndrome, compartmental pressures were measured and found to be normal (9–10 mmHg). He received a final (third) dose of ViperaTAb and gradually improved with conservative management. |
| 4    | 7.8, 22.3 and 27.9            | 142.6                                  | Systemic envenoming with cardiotoxicity. Envenoming associated thrombocytopenia. | A male child presented 3.4 h after sustaining a bite on the right ankle in a rural area. At presentation he was lethargic, complained of abdominal pain, and had ST changes and sinus tachycardia on ECG. Adrenaline and fluids were given as urgent treatment in the emergency department. Bloods revealed a thrombocytopenia (35 × 10⁹/L), raised lactate (2.3 mmol/L) and metabolic acidosis (pH 7.32). Despite delays due to a lack of antivenom stock at the hospital, his symptoms improved following antivenom administration. The following day the thrombocytopenia had worsened and remained low for a further five days (nadir of 5 × 10⁹/L). Two further doses of antivenom and 1 unit of platelets were administered and the patient observed until sustained improvement in platelets count (80 × 10⁹/L at discharge). |
| 5    | 3.2 and 5.7                   | 67.3                                   | Severe systemic envenoming. Importance of rapid assessment and treatment. | Within 15 min of a bite to his right hand, the male patient vomited, felt unwell, became drowsy and experienced swelling of his mouth and throat. He was treated at the scene with IM adrenaline before transfer to hospital by air ambulance. Antivenom was administered within 30 min of presentation to hospital. Following antivenom, systemic features resolved. A second dose of antivenom was administered for progressive swelling to the envenomed limb. The patient made a good recovery and was discharged 2 days later. |
and months [23]. Unfortunately, a radio and newspaper campaign in 2017 [24,25] failed to reduce the number of people presenting late to hospital.

The frequency of repeated dosing of ViperaTAb in this study (33.1%) is greater than previously reported in two large case series in Sweden (15.7% and 20.0%) [11,17] and a systematic review of all European antivenom (15.8%) [5]. This may be because of frequent contact between poison centre staff and clinicians looking after the patients, with careful discussion about need for additional antivenom. The common indication for repeated doses in this study was persistence or worsening of local envenoming, in particular swelling and increasing pain. On several occasions, clear benefit became apparent shortly after antivenom administration. This is supported by the shorter length of hospital duration for patients administered early (<6 h) repeated dosing of antivenom for persistent or increasing local envenoming and the illustrative cases 1 and 5 in Table 3. However, several French papers on V. berus and V. aspis envenoming report limited beneficial effect on local oedema and no reduction in length of hospital stay for patients administered multiple doses of ViperFAV for persisting oedema [20–22]. As a result, the practice of repeated dosing is reserved for patients with increasing Audebert envenoming severity score [26] or recrudescence of clinical features, following advice from French Poison Control Centres [20].

The need for repeated antivenom dosing because of persisting/recurrent systemic envenoming was uncommon in this study (only 2.9%), similar to that for patients treated with ViperaTAb by the Swedish Poisons Information Service (13/415, 3.1%) [17]. This contrasts with a French study which showed that 4/23 (17.4%) and 7/64 (10.9%) of patients with V. berus and V. aspis envenoming, respectively, treated with ViperaTAb developed persisting systemic envenoming after the initial dose of antivenom [20]. This apparent difference in effectiveness between UK/Scandinavia and France warrants further investigation. A comparative randomized control trial would help determine comparative effectiveness and the clinical implications of Fab and F(\(ab\))\(_2\) pharmacokinetics in Vipera genera envenoming [9,27].

Despite promising in vitro cross-reactivity of ViperaTAb across the Vipera genus [10], further evidence is required to demonstrate clinical effectiveness, especially in light of its relatively poor performance against Vipera ammodytes envenoming [23]. Both ViperaTAb and ViperFAV antivenoms are manufactured using V. berus venom sourced from Russia (Ian Cameron, personal communication). Consequently, the possibility of intraspecies geographical variation in venom composition should be considered as this might cause variable effectiveness of antivenom in different geographical locations [28,29].

ViperaTAb was well tolerated, with only three probable and three possible adverse reactions detected. With the exception of one patient who was treated cautiously with adrenaline and corticosteroids, the reported adverse reactions resolved spontaneously. Four of the six patients went on to receive a further dose of antivenom without any additional reaction. The lack of recurrent adverse reactions in repeatedly dosed individuals is suggestive of an alternative mechanism to classical Type-1 IgE mediated hypersensitivity [30]. In addition, it supports the advice that intradermal testing should not be used to predict patients susceptible to adverse reactions to antivenom [31]. It seems likely that the three observed cases of thrombocytopenia, in one case before antivenom treatment and in two cases after antivenom treatment, represent a manifestation of envenoming as commonly described in literature rather than an adverse reaction to antivenom [3,12,22].

Although rare, the pronounced early anaphylactic reaction to venom is a medical emergency and necessitates immediate management with adrenaline while antivenom is prepared. In this cohort, all 23 patients with early anaphylactic reactions – some of whom required multiple doses of adrenaline – responded to antivenom with rapid resolution of symptoms. In one patient, angioedema returned following antivenom administration and a further dose of antivenom was given with good effect. The frequency of systemic features of envenoming was similar for children, adults, and older adults aged >64 yrs, consistent with previous reports of V. berus and V. aspis envenoming [21,22]. Despite this, children more frequently received multiple dosing of antivenom and had modestly longer durations of hospital stay. One possible explanation of this discrepancy is that children experience greater bite site cytotoxicity with more pronounced oedema and haematoma formation due to their higher venom dose per body weight [21].

Four patients underwent surgery during their hospital admission resulting in an increased length of hospital stay (median of 6 days). One case underwent a fasciotomy without prior assessment of compartmental pressure (Table 3). The resulting surgery revealed healthy muscle. A further case was due to undergo imminent fasciotomy before advice to reconsider altered the plan and a conservative approach was taken. The patient with severe local envenoming with marked oedema of the thorax, after a bite on the hand, was effectively treated with antivenom only, after measurement of compartmental pressures showed normal pressures (Table 3). Local cytotoxicity from V. berus envenoming can result in marked soft tissue oedema but seldom if ever results in compartment syndrome. Both ultrasound and compartmental pressure assessment have been strongly advocated to distinguish extra fascial from subfascial oedema, thus limiting surgical intervention where possible [32–34].

There was marked gender difference in bite site location, with male subjects more likely to be bitten on the upper limbs, especially on their hands (Figure 5), and females more likely to be bitten on the lower limbs. This tendency was also observed in snakebites in the USA, where 56% of bites in males were on the upper extremity, and 67% resulted from handling the snake [35]. Unlike many tropical countries where envenoming is associated with agricultural work, envenoming in the UK typically occurs during leisure activities. As a result, a significant number of upper limbs bites resulted from deliberate interaction with the snake and
demonstrates the need for educational messaging, especially to men, to prevent such envenoming in future.

Limitations

The rarity of adder bite and envenoming in the UK necessitates the use of national poison centre records for prospective data collection. Efforts were made to overcome the inherent limitations of this method by using single caller follow up of cases and objective assessments of outcome (duration of hospital stay, adverse events and need for recurrent antivenom administration) in addition to an independent subjective assessment of effectiveness by the attending clinician. However, limitations are acknowledged with incomplete data collection for three patients, a lack of systematic collection of laboratory findings, a lack of comprehensive ancillary medications used, and no routine follow up arranged for discharge.

Conclusion

V. berus bites necessitating antivenom use are rare but remain a potentially life-threatening emergency in some circumstances as shown by the occasional need for adrenaline or intubation. ViperaTAb antivenom appears to be effective and safe and should be administered as soon as possible for patients meeting clinical criteria outlined in Box 1. Patients require close observation following antivenom for adverse reactions and recrudescence of swelling and pain in the limbs. Close collaboration with expert consultant advice from the NPIS can help optimise antivenom timing, ensure repeated dosing is given appropriately, and avoid unnecessary surgical intervention. All hospitals, particularly those located in areas of relatively high incidence, should stock sufficient antivenom available at short notice, 24 h a day. Increased community awareness of the dangers of snake handling may help avoid some unnecessary upper limb bites.

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