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Review

Animal toxins — Nature’s evolutionary-refined toolkit for basic research and drug discovery

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ABSTRACT

Venomous animals have evolved toxins that interfere with specific components of their victim's core physiological systems, thereby causing biological dysfunction that aids in prey capture, defense against predators, or other roles such as intraspecific competition. Many animal lineages evolved venom systems independently, highlighting the success of this strategy. Over the course of evolution, toxins with exceptional specificity and high potency for their intended molecular targets have prevailed, making venoms an invaluable and almost inexhaustible source of bioactive molecules, some of which have found use as pharmacological tools, human therapeutics, and bioinsecticides. Current biomedically-focused research on venoms is directed towards their use in delineating the physiological role of toxin molecular targets such as ion channels and receptors, studying or treating human diseases, targeting vectors of human diseases, and treating microbial and parasitic infections. We provide examples of each of these areas of venom research, highlighting the potential that venom molecules hold for basic research and drug development.

1. Introduction

Venomous animals have taken advantage of the inherent biological complexity of their victims by developing toxins that target one or all of the nervous, musculoskeletal, or cardiovascular systems in order to facilitate prey capture or defense against predators. For example, many predatory venoms contain neurotoxins that disrupt neuromuscular transmission in order to rapidly paralyse envenomated prey [1]. Due to the inherent specificity and potency of most venom molecules, venoms have become an invaluable source of modulators to study a wide range of molecular targets, especially ligand-activated [2–4] and voltage-gated ion channels [5–7]. While research on many potential drug targets is hampered by the lack of selective modulators, our increasing knowledge of venom pharmacology is beginning to shed light on some of these poorly studied targets (e.g. [8]). Here we provide examples of how venom compounds have been used to study diverse molecular targets and aid our understanding of their role in human disease. We use knowledge gained from approved venom-derived drugs [9,10] to illustrate how basic research can lead all the way from venom to a human therapeutic. Finally, we touch on other potential uses of toxins for human health, such as targeting vectors of human diseases, and development of antimicrobial and antiparasitic drugs (Fig. 1).

2. The molecular toolkit of bioactive venom compounds

During the course of evolution, venom has evolved independently at least 100 times across eight different phyla [11]. It has been estimated that there are > 220,000 venomous species on the planet, representing ~15% of extant animal biodiversity [12], spanning invertebrates such as annelids, arthropods, cnidarians, molluscs, nematodes, sea urchins, star fish, and vertebrates such as snakes, lizards, fish, shrews, and platypus [1,12–14]. Chemically, venoms are complex mixtures of salts, small molecules, peptides, and proteins [15,16], with an estimated 20 million compounds in spider venoms alone [17]. Chemically, venoms are complex mixtures of salts, small molecules, peptides, and proteins [15,16], with an estimated 20 million compounds in spider venoms alone [17]. Most of these compounds are toxins, which are defined as substances that cause dose-dependent pathophysiological injury to a living organism, thereby reducing their viability [18]. The high metabolic cost of venom production [19] has ensured that over the course of evolution only the most potent toxins have prevailed. Venoms can therefore be considered natural libraries of highly optimised bioactive compounds [20,21]. Given the vastly different phyletic origins of venomous species and the even greater diversity of their target organisms, it is expected that venom toxins modulate an extremely diverse range of molecular targets (Fig. 2).

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The dominant components of most venoms are small peptides. The high potency, selectivity and biological stability of many venom-derived peptides has made them particularly attractive from a drug discovery perspective. There are already two FDA-approved venom-derived peptides that have made them particularly attractive from a drug discovery perspective. There are already two FDA-approved venom-derived peptides that have been used to target human disease or disease-inducing organisms.

Venomous animals have evolved peptide toxins with a huge diversity of molecular frameworks, most of them with tertiary structures stabilised by disulfide bonds. The most well-known of these disulfide-rich frameworks is the ubiquitous inhibitor cystine knot (ICK) or knottin motif, which is defined as an antiparallel β-sheet stabilised by a cystine knot. A major advantage of ICK peptides for drug development is their resistance to proteases, high temperature, and harsh chemicals, which has been attributed to their knotted structure. However, despite our extensive knowledge about a limited number of disulfide frameworks present in animal venoms, there are many families of disulfide-rich venom peptides that remain completely unexplored. Further research on these novel disulfide-rich frameworks is therefore required to unleash any potential they might hold for drug development.

Recent advances in proteomics and transcriptomics have now made it possible to rapidly elucidate the venom proteome of even miniature venomous animals. While these developments have greatly accelerated our understanding of the structure, function and evolution of venom proteins, they have led to a general neglect of non-proteaceous venom molecules. A wide variety of small molecules have been found in venoms including salts, amino acids, hormones, nucleosides, neurotransmitters, and polyamines. In addition to playing important roles in envenomation, some of these molecules are valuable pharmacological tools.

3. Screening strategies — Finding the right assay

Early toxicological research focused on the study of venoms that adversely affect humans, with the aim of understanding their mechanism of action. The molecular targets of these venoms were typically delineated using low-throughput approaches such as in vivo and ex vivo tissue studies, manual patch-clamp recording, or biochemical approaches such as radioligand binding studies. These bioassays can be highly informative because they provide information on integrated tissue or cellular responses that can be used to provide target information. However, they require a priori knowledge about well-defined receptor populations and signalling pathways in target tissues, as well as the availability of selective agonists or antagonists.

In addition to providing fundamental insights into the molecular targets and mode of action of venoms and toxins, these traditional approaches yielded key insights into mammalian physiology, as exemplified below. However, these approaches neglected toxins that are of no consequence to human health, those that are present in low abundance in venom, and toxins that act on novel targets that are not represented in conventional bioassays. These neglected toxins may nonetheless offer fundamental insights into human disease, and some may even have therapeutic benefit. Accordingly, high-throughput screening approaches have been developed over the past decade to exploit the exceptional biodiversity of animal toxins for drug discovery. Since many venoms and toxins exert these biological effects through actions on cell membranes, receptors and ion channels, high-throughput techniques assessing changes in cellular signalling have proven particularly insightful. These include electrophysiological techniques, fluorescent imaging of Na⁺, Ca²⁺, K⁺ or Cl⁻ flux, enzyme-linked immunosorbent assays (ELISAs), as well as assays based on detection of bioluminescence, fluorescence polarisation, fluorescence-resonance energy transfer (FRET), bioluminescence resonance energy transfer (BRET), and scintillation proximity. Invariably, these assays are designed to determine pharmacological activity at specific targets of interest, although each technique has limitations with regard to assay flexibility, throughput, amount of toxin required, and specificity. Thus, the choice of assay will depend on whether the intention is to identify the biological target of venom components that exert effects on biological processes either at the level of the cell or organism, or whether the screen is aimed at discovery of novel bioactivity at specific molecular targets or processes of interest.

4. Venom compounds as tools for understanding human physiology

4.1. Venom toxins provide key insights into the structure and function of acetylcholine receptors

It has been known since the early 1960s that the specific and virtually irreversible blocking action of α-bungarotoxin, a venom peptide from the banded krait, is due to functional antagonism of the depolarising effects of acetylcholine. However, the identity of the presumed target of α-bungarotoxin, the cholinergic receptor protein, was unclear until a single toxin-bound membrane protein was isolated from electric eels of the marbled electric ray Torpedo marmorata and the electric eel Torpedo bicolor.
Electrophorus electricus [59]. It is now known that acetylcholine receptors (AChRs) can be classified into muscarinic (mAChR) and nicotinic (nAChR) subtypes, with the latter subtype being the target of α-bungarotoxin and numerous other venom peptides. mAChRs are metabotropic receptors whereas nAChRs are pentameric ligand-gated ion channels. The profound impact of α-neurotoxins on various biological processes highlighted the importance of nAChRs in both physiology and pathology. Early medical applications included the surgical use of tubocurarine, a plant-derived toxin that potently inhibits muscle nAChRs and is famous for its use as a paralysing agent in poison arrows.
employed by South American Indians [60]. The use of tubocurarine as a muscle relaxant allowed patients undergoing surgery to be paralysed without using dangerously high doses of general anaesthetics [61]. More recently, nAChRs have gained attention as putative therapeutic targets for a range of human disorders. For example, a number of nAChR subtypes are thought to play key roles in chronic pain and inflammation [62], and several α-conotoxins with good selectivity for these subtypes are being developed as drug leads [63,64].

The high affinity interaction of α-neurotoxins with nAChRs also provided key insights into the structure and mode of action of these receptors [65], and they remain important pharmacological tools due to their exquisite selectivity for specific AChR subtypes. For example, fluorescent derivatives of α-bungarotoxin are excellent probes for identifying α7 nAChRs [66], while subtype-selective α-conotoxins from the venom of marine cone snails can distinguish between nAChR subtypes containing common subunits [67,68]. Since ablation of nAChR subunits using gene editing approaches would lead to complete loss of all nAChR isoforms containing the deleted subunit, α-neurotoxins remain invaluable tools for dissecting the physiological roles of specific nAChR subtypes and their contribution to human disease.

4.2. Uncovering new analgesic targets using animal toxins

In addition to being a valuable source of molecules that modulate known drug targets, venoms can also be used to discover new therapeutically targets. For example, The Centre of Excellence for New Target Discovery in Brazil, which is co-funded by GlaxoSmithKline, is focused on using venoms to identify and validate new drug targets. One therapeutic area in which venom peptides have been successfully used to identify new drug targets is chronic pain [8,69]. Although the primary role of venom for most animals that possess a venom system is predation, most venomous animals also use venom for defense. Some animals, such as bees, caterpillars, stonefish, and stingrays, use venom exclusively for this purpose. In contrast to the diverse mechanisms by which predatory venoms debilitate prey, defensive venoms are largely algogenic, that is they are designed to cause intense pain (as most people that have been stung by a bee or wasp can testify).

Unlikely death, pain induces a lasting memory that can be conveyed to conspecific predators. Since pronociceptive venom toxins have evolved to target vertebrate sensory neurons they have the potential to uncover new components of mammalian pain signalling pathways and to serve as pharmacological tools for studying these components [69,70].

From a screen of venoms that activate mouse sensory neurons, Osteen et al. recently isolated two homologous peptides (Hm1a and Hm1b) from venom of the African tarantula Heteroscodra maculata that activated only a subset of sensory neurons [8]. The effect of these peptides was blocked by tetrodotoxin (TTX), a non-selective inhibitor of voltage-gated sodium (NaV) channels. This ultimately led to the discovery that these peptides are highly potent and selective agonists of NaV1.1 channels [8], which were not previously thought to be involved in pain signalling. NaV1.1 was shown to regulate the excitability of Aδ nerve fibres that transmit mechanical pain signals to the spinal cord, thereby highlighting this ion channel for the first time as a potential analgesic target [8]. Moreover, Nav1.1 was shown to be present in sensory neurons that innervate the gut, and its expression was upregulated in a mouse model of irritable bowel syndrome (IBS) [8]. This suggests that compounds that specifically inhibit NaV1.1 might be useful for treatment of IBS-related gut pain, and indeed NaV1.1 inhibitors were recently shown to reduce mechanical hypersensitivity in several models of chronic visceral pain [71].

An unexpected consequence of the discovery of Hm1a and Hm1b was their application to Dravet syndrome (DS) epilepsy [72,73], which is caused by heterozygous loss-of-function mutations in the gene encoding Nav1.1. DS is a pharmacoresistant epileptic encephalopathy characterised by childhood-onset polymorphic seizures, cognitive impairment, psychomotor regression, autistic traits, and increased risk of sudden death [72,73]. In the central nervous system, NaV1.1 is found in the axon initial segments of GABAergic inhibitory interneurons, and consequently epileptic seizures in DS are thought to result from the reduced inhibitory activity of these neurons [72]. Remarkably, in a mouse model of DS, intracerebroventricular infusion of Hm1a restored the function of inhibitory interneurons and led to almost complete abolition of seizures within 3 days, which rescued the mice from premature death [72].

In a similar approach to that described above, a screen of snake venoms on somatosensory neurons revealed robust activation of trigeminal ganglia by Texas coral snake venom [74]. This activity could only be reproduced when two toxin components were combined to form the high affinity heteromeric MitTx complex. Subsequent patch-clamp studies revealed the target of MitTx to be acid-sensing ion channels (ASICs). ASICs are unusual homo- or heterotrimERIC channels that are activated by protons and expressed throughout the nervous system [75,76]. Although MitTx promotes activation of several ASIC subtypes, it is ~100-fold more potent at the splice-isomers ASIC1a and ASIC1b compared with other subtypes. Texas coral snake envenomation causes intense pain in humans, and indeed MitTx elicited robust nociceptive behaviour when injected into the hindpaw of mice. This response was absent in pan-ASIC1 knockout mice, but present in an ASIC3 knockout, revealing the contribution of ASIC1 to peripheral pain sensation.

A subsequent study further delineated the specific roles of ASIC1a and ASIC1b in different pain pathways using mambalgin toxins isolated from black mamba venom [77]. Mambalgins were found to block ASICs expressed in the central (ASIC1a, ASIC1a/2α, andASIC1a/2β) and peripheral (ASIC1b and ASIC1a/1b) nervous system. Intrathecal and intracerebroventricular injection of mambalgin in mice produced potent analgesia that was naloxone resistant and dependent on the presence of ASIC1a, suggesting an opioid independent mode of action proposed to be via central inhibition of ASIC1a/2α channels. Strikingly, peripheral intraplantar injection of mambalgins in mice produced analgesic effects in wild-type and ASIC1a knockout mice. This demonstrated for the first time that ASIC1b-containing channels in rodent nociceptors, as opposed to the splice isoform ASIC1a, are important for peripheral analgesia. More recent work using an ASIC1b knockout animal confirmed the role of this subtype in peripheral pain sensing [78]. In this model, peripheral injection of mambalgin had no effect on acid-induced hyperalgesia in ASIC1b knockout mice. Thus, the combined use of genetic techniques and venom toxins (MitTx and mambalgins) enabled dissection of the roles of ASIC1 splice isoforms in pain signalling.

4.3. Using toxins to elucidate the mechanisms underlying cold pain

In addition to venom peptides, non-peptidic marine toxins have proven to be invaluable tools for elucidating diverse pain pathways. Ciguatoxins (CTX) are temperature-stable polyethers produced by various species of Gambierdiscus, a benthic dinoflagellate [79]. The consumption of fish with high levels of ciguatoxins present in their flesh and viscera, results of bioaccumulation, causes ciguatera fish poisoning [80]. These effects are due the action of CTX on ion channels including Nav channels [80]. The most potent congener P-CTX-1 has been used to pharmacologically probe neuropathic pain (Fig. 3). Ciguatera is characterised by both gastrointestinal and neurological symptoms including paresthesia, pruritus and cold allodynia, and it can even lead to death in severe cases [81]. The pathognomonic cold allodynia observed in ciguatera patients is also reported in a broad range of clinical indications including in chemotherapy patients, complex regional pain syndrome, and fibromyalgia [82–84]. Defined by sensitivity to innocuous cooling temperatures, cold allodynia is a poorly understood phenomenon. Subcutaneous injection of P-CTX-1 into the rodent hindpaw causes robust and concentration-dependent cold allodynia [85], and
therefore it is uniquely positioned to elucidate the molecular basis of cold allodynia.

The large family of thermosensitive transient receptor potential (TRP) channels plays a critical role in temperature sensation [86]. Interestingly, TRPM8, the canonical cold sensing channel that is sensitive to menthol, is not associated with P-CTX-1-induced cold allodynia, as TRPM8 null mice administered P-CTX-1 display robust pain behaviours in response to innocuous cooling [85]. Rather, the combined findings from ex vivo skin-saphenous nerve recordings, in vivo studies and in vitro assays highlighted a role for TRPA1 in P-CTX-1-induced cold allodynia [85]. However, TRPA1 is not directly activated by P-CTX-1. Both TRX-sensitive and TRX-resistant NaV channels were responsible for the altered neuronal excitability, which drives a TRPA1-dependent change in cold sensing [85]. Although the precise role of different TRP channel isoforms in cold sensing remains contentious, the use of this marine toxin has proved valuable in delineating these complex mechanisms.

P-CTX-1 was also utilised to assess a range of analgesic treatments for cold allodynia. Using a suite of selective pharmacological agents including venom-derivated peptides (such as analgesic conotoxins; recently reviewed in [87]), P-CTX-1-induced cold allodynia was found to be dependent on NaV1.6 and NaV1.8 [88]. Oxaliplatin-induced cold allodynia was completely blocked by μ-conotoxin GIIXA, a NaV1.6 inhibitor from cone snail venom [89], although intraplantar administration of Cn2, a NaV1.6-selective agonist from scorpion venom [90], elicited spontaneous pain and mechanical allodynia but did not enhance cold sensitivity. The mechanisms underlying neuronal cold sensitivity were further delineated using a class of presynaptic neurotoxins produced by mamba snakes (Dendroaspis sp.) (Fig. 3). Specifically, in trigeminal neurons, application of α-dendrotoxin-K, a blocker of Kv1.1 channels, elicited a reversible, concentration-dependent shift in the cold threshold [91].

4.4. Using scorpion toxins to study the function of peripheral sensory neurons

Subtype-selective NaV channel toxins have confirmed a crucial role for both NaV1.6 and NaV1.7 in pain signalling (Fig. 3). The NaV1.6 subtype is highly expressed in the central and peripheral nervous system, in particular at the nodes of Ranvier [92]. Due to the expression of NaV1.6 on motor neurons, NaV1.6 null mice are not viable beyond post-natal day 14 (P14) [93]. This greatly impeded the study of NaV1.6 in peripheral sensory neurons, where its role in excitability was less well defined. The subtype-selective scorpion scorpion toxin Cn2 was therefore used to probe the role of this channel in nociception. Cn2 causes NaV1.6 to open at hyperpolarised (more negative) potentials, and it enhances resurgent currents in large diameter sensory neurons in vitro and increases excitability of medium-diameter A fibres ex vivo [90,94]. Therefore, it follows that in vivo nociceptive behaviours elicited by Cn2 administration are primarily mediated by medium-diameter rather than smaller sensory neurons. Critically, this demonstrates how toxins can be used to pharmacologically isolate a target of interest (in this case NaV1.6) in the presence of highly similar homologs of the target (NaV1.1, NaV1.7, NaV1.8 and NaV1.9).

The toxin OD1, isolated from venom of the Iranian yellow scorpion Odontobuthus dorai, potently enhances the activity of NaV1.7 by inducing a hyperpolarising shift in the voltage-dependence of activation, inhibiting channel fast inactivation, and increasing peak current [95]. Injection of OD1 into the footpad of mice elicits pain behaviours including licking, lifting and flinching of the injected paw that can be reversed by local or systemic treatment with selective NaV1.7 inhibitors. Thus, the analgesic efficacy of NaV1.7 inhibitors can be rapidly profiled in vivo using OD1-induced pain behaviour, demonstrating that even toxins with no therapeutic potential can be effective tools for understanding the complexities of peripheral pain sensing [70] (Fig. 3).

4.5. Snake venom components act on almost every factor involved in hemostasis

Snakes are the most well-studied venomous animal, and hundreds of snake-venom proteins have been isolated over several decades. These venom components have improved our understanding of venom toxicity, significantly advanced our the knowledge of hemostatic disease mechanisms, and led to the development of diagnostic agents and life-saving drugs [96–98]. Due to the immense amount of research on snake venoms, we focus here on just a few examples where studies of snake venom have advanced our understanding of hemostasis and associated diseases.

4.5.1. Vasconstrictor peptides from stiletto snakes are orthogs of human endothelin

Studies into the lethal effects of envenomation by the burrowing asp Atractaspis engaddensis led to the discovery of a novel peptide named sarafotoxin (SFTX). SFTX is a potent vasconstrictor of coronary blood vessels that can induce cardiac arrest [99–101]. Several SFTX isoforms were soon isolated from other stiletto snakes [102,103], and it was subsequently discovered that the SFTXs are in fact orthologs of mammalian vasconstrictor peptides known as endothelins (ETs) [104]. However, a major difference between these peptides is that a protease cleavage site involved in ET inactivation is not present in the SFTXs [105,106], making them more stable in vivo. This allowed the SFTXs to be used as molecular tools to study vasocostruction and expanded the available pharmacological toolbox beyond the ETs. The SFTXs and ETs have both been used to characterise ET receptors and downstream signalling pathways [107,108].
4.5.2. The snake-venom protein botrocetin sheds light on von Willebrand factor function

von Willebrand factor (vWF) is important for platelet adhesion, and its deficiency in humans leads to a common blood coagulation disorder known as von Willebrand disease (vWD). Botrocetin is a heterodimeric protein found in venom of the South American pit viper Bothrops jararaca and it functions as a platelet-aggregating protein. It was initially named venom coagglutinin [109] as it leads to agglutination and aggregation of platelets by forming a complex with vWF and enhancing its affinity for the platelet receptor glycoprotein Ibα (GPIbα). This effect makes botrocetin an excellent diagnostic tool to assay for vWF binding to GPIbα in diseases such as vWD, or thrombopathies such as Bernard-Soulier disease [110,111]. Botrocetin has also been used as a tool to study the vWF-platelet interaction and its role in disease [112,113]. Unlike other vWF binding proteins such as the antibiotic ristocetin, botrocetin-induced platelet agglutination activity is independent of the vWF multimer size. This is advantageous as it facilitates quantitation of vWF and allows assessment of vWD severity. The combination of botrocetin and ristocetin has allowed identification of missense mutations that cause Type B vWD [114].

4.6. Viper venom promotes nerve growth

A fortuitous observation using viper venom to study neurite outgrowth in healthy and tumorous cells led to the Nobel Prize in 1986. In studies of the regulation of cell growth, a fraction from mouse sarcoma extracts was shown to induce nerve cell growth in chick embryos [115]. This fraction was termed nerve growth factor (NGF) [116], and it was shown to be composed of both nucleic acid and protein. Snake venom from the cottonmouth viper Agkistrodon piscivorus was known to contain phosphodiesterase activity, and thus it was proposed that this venom could degrade the nucleic acid component (which was thought to be a contaminant) and reveal the active ingredient. Surprisingly, minute amounts of crude snake venom strongly promoted nerve growth, approximately 3000–6000 times more potently than the NGF-containing sarcoma fraction [117]. Comparison of the sarcoma extract and venom contents ultimately led to isolation and characterisation of the proteinaceous NGF. NGF has since been found in several snake venoms from the Viperidae, Crotalidae, and Elapidae families and it has been used widely to study cell growth regulation [118,119]. The use of snake venom in this case not only led to a serendipitous finding with widespread importance to basic science, but NGF has also been important for elucidating the etiology of diseases such as developmental/growth deformations, tumours, and delayed wound healing.

4.7. Cobra venom elucidates complement system activity in innate immunity

Cobra venom factor (CVF) from the venom of cobra (Naja sp.) snakes has the intriguing property of disrupting activation of the complement component of the innate immune system. The complement system is comprised of a set of interacting proteins, and it can be activated in three different ways for it to play a role in pathogen clearance. The alternative pathway of activation involves spontaneous hydrolysis of complement factor C3 to C3b, which allows the plasma protein complement factors B and D to bind to C3b and transform it to C3-convertase on the external surface of an invading pathogen membrane. Subsequent recruitment of additional components of the complement pathway system leads to assembly of the membrane attack complex, which forms pores in the target cell membrane that kill or damage the pathogen [120,121].

CVF is a structural and functional analogue of the human complement factor C3, and it acts in a similar manner to human factors C3b and C3c to activate and deplete the complement cascade [122,123]. These properties of CVF have made it possible to independently study the role of complement in host defense, immune responses, and pathogenesis of disease. CVF was in fact pivotal to the discovery of complement factor B itself, which facilitated further characterisation of this complex pathway [120,124–127]. CVF is not only an immensely useful research tool, but it has recently been used as an immunosuppressant in tissue and organ transplants and cancer therapies [128,129], and CVF derivatives are being designed to activate the complement system in disease states where it has been depleted [130,131].

4.8. Spider toxins highlight the role of ASIC1a in stroke-induced brain damage

Stroke is the second leading cause of death worldwide [132], yet therapeutic options are limited to dissolution of blood clots with tissue plasminogen activator (tPA). Unfortunately, due to the risk of inducing an intracranial haemorrhage, tPA is used in < 5% of stroke patients [133], and consequently there is an unmet need for compounds that can protect the brain from stroke-induced injury. Clinical trials of numerous agents, especially glutamate receptor antagonists, have failed to provide effective neuroprotection in stroke patients, which has spurred interest in better understanding the etiology of ischemia-induced brain injury [133]. Spider-venom peptides have been crucial for uncovering the key role of ASICs in stroke-induced brain damage, and validating these channels as a target for neuroprotective drugs [134–137].

During ischemic conditions the brain switches from oxidative metabolism of glucose, its primary fuel, to anaerobic glycolysis, which results in lactic acidosis. The brain extracellular pH can fall from ~7.3 to as low as 6.0 in the ischemic core [134,138,139], which is sufficiently acidic to robustly activate proton-gated ASICs. The ASIC1a isoform is the dominant subtype in rodent and human brain [140,141] and it primarily conducts sodium ions, although it is the only ASIC subtype that is also permeable to calcium (PNa/PCa ~ 15) [75]. It has been proposed that ASIC1a activation contributes to toxic intracellular calcium overload during stroke via both direct calcium permeability and downstream activation of voltage-gated calcium (CaV) channels and NMDA receptor-gated channels via ASIC-induced membrane depolarisation [75]. However, the primary mechanism by which ASIC1a activation leads to neuronal cell death appears to be recruitment and activation of RIPK1 [142,143], a master regulator of necroptosis.

Although several small molecules and endogenous ligands of ASIC1a have been discovered to date, their lack of potency and selectivity limits their use as pharmacological tools for dissecting the role of ASICs in disease. The first potent and moderately selective ASIC1a ligand discovered is PcTx1, a 40-residue peptide from venom of the Trinidad chevron tarantula Psalmopoeus cambridgei [144]. PcTx1 inhibits homoseric ASIC1a (IC50 ~ 1 nM), ASIC1a/2a heteromers (IC50 ~ 3 nM), and ASIC1a/2a (35–85% inhibition at 50 nM) channels, and potentiates proton-activated ASIC1b currents by over 3-fold control (EC50 ~ 25 nM) [144–148].

The key role of ASIC1a in stroke-induced brain damage was first highlighted in a seminal 2004 study which showed that genetic ablation of the ASIC1a-encoding gene reduced brain infarct size by ~60% in a mouse model of ischemic stroke [134]. However, since genetic ablation can give rise to compensatory upregulation of related channels with unknown consequences [149,150], these researchers also used “PcTx1 venom” (i.e., crude P. cambridgei venom) as a pharmacological tool to validate the genetic studies. They showed that “PcTx1 venom” administered 30 min before and after stroke onset reduced infarct size by the same magnitude as genetic ablation of ASIC1a [134]. However, spider venoms are complex molecular cocktails that modulate an array of ion channels, and the venom of P. cambridgei is no exception. PcTx1 constitutes only ~0.4% of total P. cambridgei venom [136], and other components in this venom are known to target TRPV1 and voltage-gated potassium (Kv) channels [151,152]. This left open the question of whether the observed neuroprotective effects were induced by PcTx1, other components in the venom, or a combination of both. To address this question a more recent study used pure recombinant PcTx1 to...
confirm the involvement of ASIC1a in stroke [136]. Structure-activity relationship (SAR) studies of PcTx1 [153,154] facilitated design of a weakly active PcTx1 mutant, which was not neuroprotective in the same rodent stroke model, thereby strengthening the evidence that ASIC1a inhibition reduces stroke-induced brain damage.

More recently, the 76-residue peptide H1i1a from venom of the Australian funnel-web spider *Hadronyche infensa* proved to be highly neuroprotective in a focal model of ischemic stroke, even when administered 8 h after the stroke onset [137]. In comparison to PcTx1, H1i1a is a slightly more potent inhibitor of ASIC1a (IC₅₀ 0.5 nM) and it is more selective, with > 2000-fold higher potency for ASIC1a than homomeric ASIC1b, ASIC2a, and ASIC3 channels. H1i1a inhibition of ASIC1a is much less reversible than PcTx1 [137], and these venom peptides have a different mechanism of action [137,155]. The different properties of PcTx1 and H1i1a broaden the pharmacological toolkit for studying ASIC1a and ASIC1a-related pathologies. Together these two venom peptides have provided striking pharmacological evidence that ASIC1a is involved in ischemic injuries of not just the brain, but also the heart [156] and kidney [157].

5. Venom compounds as leads for novel anti-infectives

The evolution of drug resistance across bacteria, fungi and parasites has created an unmet need for novel anti-infectives. The 2019 United Nations report on antimicrobial resistance estimated that > 700,000 people die annually due to drug-resistant infections [158]. While most focus has been on human antibiotic resistance, control of parasitic nematodes in both livestock [159,160] and companion animals [161,162] is also seriously threatened by resistance. Despite the clear need for new anti-infectives, there is a diminishing pipeline of anti-infective drug candidates in both the human and veterinary spaces [163-165], which has created interest in venoms as a potential source of novel anti-infectives.

5.1. Antimicrobial toxins

Antimicrobial peptides (AMPs) are found in many animal venoms, and hundreds of examples have been reported with varying activity against fungi and both Gram-positive and Gram-negative bacteria [166-171]. Venom-derived AMPs (vAMPs) are typically small, linear peptides that are rich in cationic residues (Lys, His and Arg), which facilitates their interaction with anionic membranes [166,172]. They are often unstructured in aqueous solution but form ordered α-helices upon interaction with lipid membranes [172,173].

Many vAMPs exhibit broad-spectrum cytolytic effects. This can be a double-edged sword for antimicrobial development as it often confers broad-spectrum activity against a wide range of bacterial and fungal pathogens but also cytotoxicity against host cells. For example, the ponerinex peptide isolated from the Amazonian stinging ant *Neoponera goeldii* have broad-spectrum activity against bacteria and fungi, but they are potently hemolytic [174]. Likewise, cupiennin 1a from venom of the wandering spider *Cupiennius salei* exhibits non-specific cytolytic activity against bacteria, insects, protozoan parasites, and human cells [175]. Mammalian cytotoxicity is typically assessed by screens against red blood cells or mammalian cell lines, but an often overlooked property of cytolytic vAMPs is that they can also cause pain. For example, venom-peptide A-5myrtoxin-Mp1a from the Australian ant *Myrmecia pilosula* has submicromolar activity against bacterial pathogens but it causes mechanical allodynia when injected into mice [176]. Melittin, the primary component of honeybee venom, is potently anti-microbial but it also induces pain through direct and indirect activation of sensory neurons [177]. Hence, future research on vAMPs should place greater emphasis on screening across a broad range of cell lines and phenotypic readouts, particularly for cytolytic peptides.

A small number of vAMPs show some degree of inherent taxonomic selectivity. For example, bicaarinin from the ant *Tetramorium bicarinatum* is highly active against bacterial and fungal pathogens (minimal inhibitory concentration (MIC) of 0.7-25 μM, which is comparable to commercial anti-infectives), with at least 10-fold selectivity over human lymphocytes [178]. It also has similar potency to commercial antibiotics against clinical isolates of the *Helicobacter pylori*, a pathogen that causes stomach ulcers [179]. However, most vAMPs will require further engineering of their selectivity in order for them to be therapeutically useful [166], and several strategies have been employed successfully for this purpose. For example, shorter analogues of the scorpion-venom peptides hadurin and jeveoin were engineered with 100-fold improvement in selectivity for *Escherichia coli* over red blood cells [180]. Modification of the hydrophobicity and net charge of the wasp-venom peptide mastoparan through selective amino acid substitutions [181] and N-terminal acylation [182] increased its affinity for negatively-charged bacterial membranes and decreased its cytotoxicity towards mammalian cells.

Although most vAMPs have not progressed beyond basic in vitro studies, some have proven effective in in vivo infection models. ZY4 is a 17-residue cyclised derivative of cathelicidin-BF from venom of the snake *Bungarus fasciatus* with improved potency, selectivity and stability over the parent toxin [183]. In vitro, ZY4 kills multidrug-resistant isolates of the human pathogens *Pseudomonas aeruginosa* and *Acinetobacter baumannii* at lower concentrations than commercial antibiotics, and it reduces bacterial load and inflammation in the lungs of mice in a *P. aeruginosa* infection model, with no reported side-effects [183]. LyeTxI-b, an N-terminal acetylated derivative of LyeTxI from venom of the wolf spider *Lycosa erythroneura*, has improved potency and selectivity against *E. coli* and also reduces both bacteriaemia and inflammation in a mouse model of septic arthritis when administered intraarticularly [184].

Route of administration is a key issue that must be considered for therapeutic deployment of vAMPs; traditionally, peptides have poor oral bioavailability [185], and hence alternative methods of vAMP administration such as topical application should be considered. The effectiveness of this approach was demonstrated for Kn2-7, a derivative of the peptide BmKn2 from venom of the scorpion *Mesobuthus martensii*, which protected the skin of mice in a *Staphylococcus aureus* in vivo infection model [186]. Topical administration can also circumvent or alleviate off-target effects that are evident when the vAMP is present systemically. Finally, it might be possible to achieve additive or synergistic effects by combining vAMPs with existing antibiotics by utilising the pore-forming activity of the vAMP to enhance antibiotic delivery. For example, Garcia et al. found an additive effect when pathogen-specific antibiotics were combined with vAMPs isolated from the venom of arachnids [187].

5.2. Antiviral toxins

Viruses continue to pose an enormous threat to global health, as exemplified by the current global pandemic caused by SARS-CoV-2 [188]. Despite intensive intervention efforts, an estimated 770,000 people continue to die annually from human immunodeficiency virus acquired immunodeficiency syndrome (HIV-AIDS) [189]. Many serious viral diseases such as severe acute respiratory syndrome (SARS), Middle Eastern respiratory syndrome (MERS), dengue, Zika, and Ebola lack vaccines or efficacious therapeutic treatments, highlighting the need for novel antivirals. Most reported venom toxins with antiviral activity are vAMPs with broad-spectrum activity; examples include melittin, snake venom PLA₂ and L-amino oxidases, and mastoparan derivatives [190,191]. Thus, a key challenge will be engineering these compounds to have selectivity for either the viral particles or the process by which they infect host cells. Mastoparan-7, derived from wasp-venom mastoparan, has broad-spectrum activity against lipid-enveloped viruses (including influenza and flaviviruses) through direct insertion and disruption of the envelope, but it remains highly cytotoxic to host cells in vitro [192]. Peptides with activity on specific viral protein targets
rather than lipids may provide better selectivity. Lactacin-1 from the central Asian spider Lachesana tarabaevi inhibits replication of dengue virus-2 (IC50 ~ 7 µM) by virtue of its ability to inhibit the viral protease NS2B-NS3, but it has low selectivity over mammalian cells [193]. The antibacterial activity of mucroporin, a vAMP from the scorpion Lychas mucronatus, was improved through histidine and arginine substitutions to create the analog mucroporin-M1 with increased net charge. Mucroporin-M1 has improved activity against both Gram-positive and Gram-negative bacteria [194], and these modifications also conferred micromolar antiviral activity in vitro (EC50 1–7 µM) against measles virus and pseudovirus models of the highly pathogenic SARS-CoV and avian influenza H5N1 viruses [195].

A key limitation of current research on venom-derived antiviral toxins is that few studies have progressed to in vivo testing. One example found that pre-treating vesicular stomatitis virus, a serious pathogen of cattle, with mastoparan-7 significantly improved clinical outcomes in a mouse challenge model through presumed inactivation of viral particles [192]. However, this needs to be confirmed in a more clinically relevant challenge model of viral infection followed by toxin administration to prove efficacy. Overall, further research is required to ascertain the potential of venom-derived peptides as viable antiviral drugs.

5.3. Antiparasitic toxins

Parasites remain one of the most serious threats to human and veterinary medicine, with current treatment options limited by drug resistance, side-effects and limited efficacy [196]. This has stimulated interest in venom as a source of novel antiparasitic compounds (for a review see [197]). Most such studies have focused on protozoan parasites, which are responsible for a range of human diseases. The most problematic parasitic disease is malaria, which is caused by Plasmodium parasites; despite intensive intervention efforts, malaria kills an estimated 435,000 people each year and control is threatened by drug resistance [198]. A further 20 million people worldwide are infected by parasitic protozoans of the genera Leishmania and Trypanosoma, with approximately 3 million new infections per year [199].

Most antiparasitic compounds isolated from animal venoms are small cationic vAMPs, which again raises the issue of taxonomic selectivity. The vAMPs mastoparan [200] and melittin [199] inhibit development of Trypanosoma cruzi (the causative agent of Chagas disease), but selectivity over host cells is low. However, bicaerinaceins potently inhibits growth of Leishmania infantum (MIC 1.5 µM), the causative agent of infantile visceral leishmaniasis, with good selectivity over human lymphocytes [178]. Remarkably, crotamine, a peptide from venom of the South American rattlesnake Crotalus durissus terrificus, crosses infected red blood cell membranes to lyse the Plasmodium parasitophorous vacuole, thereby killing the parasite [201]. This may present an opportunity for combination therapy with commercial antimalarials, facilitating drug transport to the parasite. Further examples of venom components active against protozoan parasites include trimorphin, a PL2 from venom of the North American rear-fanged snake Trimerophodon biscutatus lambda that inhibits growth of Leishmania major [202], and crovinin, a cysteine-rich secretory protein (CRISP) from the rattlesnake Crotalus viridis viridis that is active against protozoan parasites at concentrations that are not toxic to host cells [199]. The protozoan Toxoplasma gondii, which causes the key zoonotic disease toxoplasmosis, is susceptible to a venom peptide from the marine cone snail Conus californicus [203], while a C-type lectin and a PL2 from B. pauloensis snake venom impair Toxoplasma adhesion and replication [204,205].

The greatest burden of disability-adjusted life years in the neglected tropical diseases comes from helminth infections, including roundworms (Nematoda) and flatworms (Platyhelminthes) [189]. These worms cause important parasitic diseases in humans, including onchocerciasis, lymphatic filariasis, and schistosomiasis. About 200 million people worldwide suffer from schistosomiasis, and it is considered the second most socioeconomically devastating parasitic disease after malaria [206]. The burden of parasitic disease is also a key concern in animal health and production, with the worldwide cost of controlling livestock parasites amounting to tens of billions of dollars [207]. Despite this clear need for novel anthelmintics, it is an underexplored area of venom research with very few reports of anthelmintic venom toxins. Crotamine [208] and ε-latroinsectotoxins from venom of the black widow spider Latrodectus mactans tepecingutattatus [209] both grow mice with the non-parasitic nematode Caenorhabditis elegans, but anthelmintic activity has not been reported against parasitic species. More recently, the spider-venom peptide H1a was shown to inhibit larval development of the barber’s pole worm Haemonchus contortus, a parasite of sheep production [210]. Gomesein, a peptide found in the venom [211] and hemocyes [212] of spiders, has broad-spectrum activity against bacteria, fungi, parasites and tumor cells [213,214], and a cyclized version was recently shown to have improved stability and activity against Plasmodium in vitro [215].

To our knowledge, no venom-derived antiparasitics have yet been demonstrated to have in vivo efficacy. As for antibacterials, antifungals and antivirals, most venom-derived antiparasitics will likely require engineering to enhance their selectivity over host cells. An alternative strategy to developing venom compounds as anti-parasitic drugs is to instead impair Plasmodium development and transmission via transgenic mosquitoes engineered to express the anti-parasitic compound in the mosquito midgut, as recently demonstrated with scorpion-venom peptides [216].

As most research on venom compounds with antibacterial, antiparasitic, or antifungal activity has been performed in the last decade, we expect that more toxins with such activities will be discovered in the near future. Most work to date has focussed on human pathogens, but there is a clear need for novel veterinary anti-infectives which could be further explored in future. Due to the commonly found cytolytic nature of these peptides, a key challenge across all of these diseases will be engineering analogs with selectivity for the targeted pathogen.

6. Targeting vectors of human disease with venom-derived toxins

Arthropods, and in particular hematophagous (blood-sucking) species, are vectors for a wide-range of organisms that can cause human disease [217]. The most pernicious vectors are undoubtedly mosquitoes, which transmit protozoans (e.g. Plasmodium species that cause malaria), parasitic nematodes (e.g. Brugia species that cause filariasis), parasitic plathyhelmints (e.g. trematodes that cause schistosomiasises), and viruses (e.g. flaviviruses that cause yellow fever, West Nile fever, and dengue) [17,217,218]. Other important hematophagous disease vectors include triatome bugs which transmit trypanosomes that cause Chagas disease, fleas which transmit the bacterium Yersinia pestis that causes bubonic plague, and ticks which transmit a wide variety of viral, bacterial and protozoan pathogens [17,217]. Rather than directly targeting disease-causing organisms, venom toxins could be used instead to target the vectors of these diseases.

Insects are the dominant vectors of human diseases, and chemical insecticides are the primary means of controlling these insect pests [17,218]. Unfortunately, all chemical insecticides target one of only a handful of molecular targets, and consequently their indiscriminate use has led to widespread resistance [17]. Thus, new insecticides with novel modes of action are much sought after. Spiders are professional insect killers, and they are the most speciose venomous animal, and consequently their venoms have been intensively studied as a potential source of novel bioinsecticides [17,217–221]. As a result, many insecticidal toxins have been isolated from spider venoms, including peptides, proteins and polypeptides, but they mostly lack the necessary taxonomic selectivity to be useful as insecticides.

An ideal insecticide should be inactive on humans and other vertebrates, as well as beneficial insects such as pollinators and natural
predators of the targeted insect pest [221]. This narrow target-organism profile makes insecticide development particularly difficult. However, spider-venom peptides have been isolated with desirable taxonomic selectivity. For example, by carefully counter-screening insect toxins from the venom of Australian funnel-web spiders in vertebrate assays, a number of insect-selective neurotoxins were isolated [218]. One of these peptide toxins (GS-o-x-hexa toxin-Hv1a), which has complex polynodal pharmacology, has been developed commercially as a bioinsecticide by Vestaron Corporation; this peptide is active against a broad range of insect pests but is inactive against bees and vertebrates [221]. In order to target anopheline mosquitoes that transmit malaria, a gene encoding GS-o-x-hexa toxin-Hv1a was engineered into the entomopathogenic fungus Metarhizium pingshaense [222], and this wep-onised entomopathogen proved highly successful in controlling malaria vectors in a field trial in Burkina Faso [223]. An advantage of this approach is that the range of susceptible species is determined by the entomopathogen rather than the venom peptide, thereby allowing deployment of insecticidal venom toxins that would otherwise lack the desired taxonomic selectivity [221,224].

Some insecticidal venom peptides have exceptional taxonomic selectivity. For example, β-diguetoxin-Dc1a (Dc1a), a 56-residue knottin from venom of the desert bush spider Digenia cantitas [225], and μ-theraphotoxin-Ae1a, a 39-residue knottin from venom of African tar-tanula Augascephalus exsandi [226], have potent insecticidal effects on German cockroaches (Blattella germanica), without significantly affecting American cockroaches (Periplaneta americana). Both knottins target insect NaV channels, and mutational analyses revealed that the susceptibility of insects to these toxins is determined by small sequence differences in their proposed binding site on the extracellular surface of the domain II voltage sensor (VSDII) [226,227]. A high-resolution structure of Dc1a bound to a cockroach Nav channel was recently determined using cryo-arrays microscopy [228], and it not only confirmed that VSDII is the binding site for Dc1a but revealed that the toxin’s inactivity against vertebrates [225] can be explained by sequence variations in the VSDII region of each of the human NaV channels. The Dc1a-NaV channel structure allows predictions to be made about which insects will be susceptible to Dc1a based on their NaV channel VSDII sequences, and it should allow use of structure-guided protein engineering to develop toxin analogs with improved potency and/or altered taxonomic selectivity.

7. Venom-derived drugs

In addition to highlighting how venom compounds can be used to elucidate disease mechanisms and uncover new drug targets, we want to provide examples of venom compounds that have progressed all the way from basic research to marketed drugs. There are currently six venom-derived drugs approved by the FDA, plus a snake-venom-derived serine protease (batroxobin; marketed as Defibrase-PSS) that is approved for clinical use outside of the USA [9,10]. Two further venom-derived drugs — dalazatide for treatment of autoimmune diseases [24] and tozuleristide for intraoperative imaging of brain tumours [25] — are currently undergoing clinical trials. The examples below demonstrate the potential for developing venom compounds into drugs [229] but, like any class of therapeutics, there are many challenges and the success rate is low [61].

The first venom-derived drug was captozide, an angiotensin-converting enzyme (ACE) inhibitor that is used to treat hypertension. The observation that patients envenomated by the lethal Brazilian viper Bothrops jararaca suffered from hypotension and severe intestinal contractions led to the discovery that the venom inhibits ACE [230–232]. Inhibition of ACE lowers blood pressure by preventing conversion of angiotensin I to the vasoconstrictive hormone angiotensin II, and by decreasing degradation of the vasodilatory peptide bradykinin. Researchers at Squibb isolated six ACE inhibitory peptides from B. jar-araca venom, one of which (teprotide, 9 residues) was both potent and stable in vivo [230]. Detailed SAR studies of teprotide ultimately led to a smaller, orally active peptidomimetic known as captopril (D-2-methyl-3-mercaptopyropanoyl-L-proline) [230,232]. Captopril was approved by the FDA for treatment of hypertension in 1981, and by the mid 1990s it had become a blockbuster drug, reaching peak annual sales of more than USD $1.6 billion [233].

Another example of a reptile peptide that was developed into a human drug is exendin-4, a 39-residue peptide isolated from venom of the Gila monster Heloderma suspectum. In striking contrast to captozide, the marketed drug (exenatide) is identical to the peptide found in the lizard’s venom. Exenatide mirrors the effects of human hormone glu-cagon-like peptide-1 (GLP-1), an incretin that stimulates glucose-de-pendent insulin secretion, suppresses glucagon secretion, slows gastric emptying, and promotes satiety [234], but it is much more stable in vivo than human GLP-1 with a plasma half-life of 2.4 h [9]. Byetta®, a formulation of exenatide that requires twice daily subcutaneous injections, was approved by the FDA in 2005 for control of Type 2 diabetes, and it achieved worldwide sales of USD $800 million in fiscal year 2009 [9]. In order to improve patient acceptance and compliance, a new formulation was developed in which the peptide is embedded in biodegradable polymeric microspheres that extends the duration of exenatide release to such an extent that therapeutic levels can be maintained with weekly injections [235]. This new formulation (Bydureon®) was approved by the FDA in 2012, and annual sales have remained steady at USD $500–600 million despite the introduction of competing incretin mimetics.

Another drug that is an exact copy of a venom peptide is ziconotide, a synthetic version of ω-conotoxin MVIIIA, a 25-residue knottin peptide found in venom of the marine cone snail Conus magus. Ziconotide is used to selectively inhibit Ca2,2.2 channels in the spinal cord, thereby preventing the propagation of pain signals from the spinal cord to processing centres in the brain [236]. It was approved by the FDA in 2004 for treating patients with chronic pain. Ziconotide is administered by intrathecal infusion, which limits its use to patients with intractable chronic pain that have failed on frontline analgesic drugs such as morphine. While ziconotide set a landmark as the first venom-derived drug with a neuronal target, its limited market acceptance due to its mode of administration and neuropsychiatric side-effects [237] has caused venom researchers to focus attention on analgesic targets in peripheral nociceptors rather than central neurons, as outlined in Section 4.2.

8. Conclusions and future directions

The integration of advanced venom proteomics with next-genera-tion sequencing of venom-gland transcriptomes has made it possible to rapidly determine, at only moderate cost, the complete venom proteome of even miniature venomous animals [46]. As a result, venom peptides and proteins, often with unique three-dimensional folds, are being discovered at an unprecedented rate. The coupling of these omics workflows with high-throughput screening technologies [56,238] is allowing the pharmacology of these compounds to be explored with unprecedented efficiency, often leading to the discovery of toxins with novel pharmacology [8,74,77,239,240]. All of these advances have combined to create tremendous interest in venoms as a potential source of drugs and bioinsecticides [12,229].

While there have been some notable successes — there are seven venom-derived drugs in clinical use and several in Phase 2/3 clinical trials — the triumphs are scarcer than many predicted. No novel venom-derived drug has been approved by the FDA since 2005 [9,10], and only one venom-derived compound has been commercialised as a bioinsecticide [221]. The reasons for this are manifold. Drug development is inherently difficult and the overall success rate is low [61]; venom-derived drug candidates are no different than any other class of therapeutic in this respect, although the overall approval rate for venom-derived drugs entering clinical trials is higher than other classes.
of therapeutics. Most venom researchers are not trained in the nuances of drug development, and this has led to over-hyped claims about the therapeutic potential of venom compounds, often in the absence of in vivo data. We encourage collaboration between venom researchers and those with experience in drug development so that, at a minimum, in vivo efficacy and toxicity can be assessed in validated disease models, using clinically viable routes of administration, before making claims about therapeutic potential. Renewed interest from pharmaceutical companies in venoms as source of drug leads and pharmacological tools [241–244] will help to drive these collaborations.

Notwithstanding these caveats, there is little doubt that the field of venoms-based drug discovery has rapidly matured over the past decade and is now producing drug leads at an unprecedented rate. Advances in cryoelectron microscopy have facilitated determination of the three-dimensional structure of venom compounds bound to large membrane receptors [228,243,245–248], and this should facilitate the rational design of drug candidates with improved potency and selectivity for their cognate molecular target. We predict that these advances will lead to significant impact in some disease areas. For example, ion channels are now the third most common human drug target after kinases and G-protein-coupled receptors [249]; venoms are unquestionably the best natural source of potent ion channel modulators and venom-derived ion channel modulators have already attracted significant interest from pharmaceutical companies [24,241–244].

CRediT authorship contribution statement

Volker Herzig: Conceptualization, Supervision, Writing - review & editing, Funding acquisition.
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Mathilde R. Israel: Writing - review & editing, Funding acquisition.
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Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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