Topical ambroxol for the treatment of neuropathic pain
An initial clinical observation

Background and problem
Neuropathic pain is common and difficult to treat. However, topical treatments such as transient functional desensitization of TRPV1 receptors as well as C-fibres with capsaicin 8% and blockade of neuronal signal transmission with lidocaine 5% are often inadequate. Other potent inhibitors of cutaneous nociceptors with few side effects would be welcome as topical therapeutic options.

Ambroxol has been authorized for the treatment of respiratory disorders since 1979 and can now be freely purchased over the counter. Since its local anaesthetic properties were recognized at an early stage [23], ambroxol-containing pastilles have also been authorized for the treatment of sore throat [10]. However, the substance has never been used to date as an analgesic, although pain-related behaviours have been suppressed in animal studies, even in chronic pain situations [14, 16, 27, 32]. Compared with local anaesthetics, ambroxol is, interestingly, a very potent blocker of voltage-dependent sodium channels, blocking these channels about 40 times more strongly than lidocaine [47]. Moreover, the sodium channel subtype Na1.8, which is preferentially expressed in nociceptive C-fibre neurons [2, 6, 35, 52], is blocked more potently compared to the other channel subtypes. Since the toxicity of the substances is comparatively very low [51, 53], the use of an ambroxol-containing semisol-
this treatment as their sole medication, and all continued taking primary medication, with occasionally varying dosages, over what was a prolonged treatment period in some cases. The patient with the longest treatment period has been using topical ambroxol for 48 months (patient 1), while the shortest case report (patient 4) covers a period of 9 months (Table 2).

### Allodynia

Dynamic allodynia was present in six of the seven patients and a hyperalgesia in response to pinprick stimuli in four patients. Both forms were present in four cases, while one patient suffered from neither. Dynamic allodynia on its own was observed in two cases, and no patient was sensitive exclusively to pinprick stimuli (Table 1). Irrespective of these configurations, all patients were sensitive to ambroxol.

### Pretreatment with capsaicin 8%

Three of the seven patients had also received capsaicin 8% as topical treatment. In two patients this was helpful, and one continues to use the medication. Both patients whose pain was relieved by capsaicin 8% also experienced this relief with ambroxol. By contrast however, the patient in whom the capsaicin had proved ineffective benefited additionally from the effect of ambroxol.

### Pretreatment with lidocaine 5%

Six of the seven patients had previously used lidocaine plasters, which proved ineffective in four cases and helpful in two patients, one of whom was unable to tolerate them. Both of the patients experiencing pain relief also reported this with ambroxol. By contrast however, all four patients in whom the lidocaine had proved ineffective profited additionally from the effect of ambroxol.

**Case reports**

**Case report 1 (Neuropathic forefoot pain)**

Born in 1942, this man presented in October 2010 (clinically) with neuropathic pain in both feet. He felt as if he was running on coals, while his right foot felt as if it were gripped in a vice. Walking or gardening was almost impossible. The foot pain could not be explained in orthopaedic terms, either on the basis of radiographs or as a result of a spondyloesis of L4/5 in 2007, and no polyneuropathy was detected on neurographic investigation. The problem was thought to be ‘likely of vertebral origin’. Clinical examination showed pronounced dynamic allodynia and hyperalgesia in response to pinprick stimuli in the arch of the right foot, and to a lesser extent in the left foot. Gabapentin had been prescribed up to a dosage of 3 x 600 mg, supplemented by buprenorphine 20 µg/h, and no further dose increase were possible in either case. Topical lidocaine 5% plasters had not proved helpful.

The first test treatment with topical ambroxol was started in June 2011, with a very successful outcome: within 5 min the stabbing pain, with an initial intensity of 8/10 on the NRS scale during the heel-to-toe roll of the right foot, and the touch sensitivity of the forefoot disappeared completely for over 8 h. The intermittent treatment with topical ambroxol has been continued to date successfully (Fig. 1). The pain has also meanwhile been treated successfully on 11 occasions with capsaicin 8% plasters, and the patient now uses ambroxol cream in phases when the pain

---

| Case Nr. | Diagnosis of neuropathic pain | Duration of pain (months) | NRS Mean | NRS Max | Alldynia Dynamic | Pin-prick Lidocaine 5 % | Capsaicin 8 % |
|----------|--------------------------------|---------------------------|---------|---------|-----------------|---------------------|----------------|
| 1        | Neuropathic forefoot pain      | 55                        | 5       | 8       | Yes              | Yes                 | No             |
| 2        | Cold phantom pain              | 68                        | 6       | 8       | Yes              | No                  | No             |
| 3        | Neuropathic pain after total knee replacement | 49 | 6 | 10 | Yes | Yes | No | Yes |
| 4        | Deafferentation pain           | 174                       | 4       | 8       | Yes              | Yes                 | No             |
| 5        | Thoracic postherpetic neuralgia| 78                        | 4       | 6       | Yes              | No                  | Yes            |
| 6        | Mononeuropathy multiplex        | 108                       | 4       | 8       | No               | No                  | No             |
| 7        | Trigeminal postherpetic neuralgia | 6                      | 5       | 8       | Yes              | Yes                 | Yes            |

NRS numerical rating scale, n.t. not tested
Topical ambroxol for the treatment of neuropathic pain. An initial clinical observation

Abstract
Background. Neuropathic pain is difficult to treat, and the available options are often inadequate. The expectorant ambroxol also acts as a strong local anaesthetic and blocks sodium channels about 40 times more potently than lidocaine. It preferentially inhibits the channel subtype Na\textsubscript{1.8}, which is expressed especially in nociceptive C-fibres. In view of the low toxicity of ambroxol, it seemed reasonable to try using it for the treatment of neuropathic pain that failed to respond to other standard options.

Material and methods. The medical records of seven patients with severe neuropathic pain and pain reduction following topical ambroxol treatment are reported retrospectively. As standard therapies had not proved sufficient, a topical ambroxol 20% cream was repeatedly applied by the patients in the area of neuropathic pain.

Results. The reasons for neuropathic pain were postherpetic neuralgia (2×), mononeuropathy multiplex, phantom pain, deafferentation pain, postoperative neuralgia and foot neuropathy of unknown origin. The individual mean pain intensity reported was between 4 and 6/10 (NRS), maximum pain at 6–10/10 (NRS). The pain reduction achieved individually following ambroxol cream was 2–8 points (NRS) within 5–30 min and lasted for 3–8 h. Pain attacks were reduced in all five patients presenting with this problem. Four patients with no improvement after lidocaine 5% and one patient with no response to capsaicin 8% nevertheless experienced a pain reduction with topical ambroxol. No patient reported any side effects or skin changes during a treatment that has since been continued for up to 4 years.

Conclusion. Ambroxol acts as a strong local anaesthetic and preferentially inhibits the nociceptively relevant sodium channel subtype Na\textsubscript{1.8}. For the first time, we report below on a relevant pain relief following topical ambroxol 20% cream in patients with neuropathic pain. In view of the positive side effect profile, the clinical benefit in patients with pain should be investigated further.

Keywords
Ambroxol · Neuropathic pain · Topical therapy · Na\textsubscript{1.8} · Local anaesthetic

Case report 2 (Cold phantom pain)

This patient presented with unusual pains: he complained of extremely painful cold sensations in both previously amputated phantom feet. A below-knee amputation of the left leg had been required for pAOD and diabetes mellitus in 2008, followed by the right leg in 2009. The cold pains, with an intensity of NRS 7–9/10, occurred sporadically several times a week, lasted from a few minutes to many hours and frequently woke the patient at night. He described the cold sensation as shifting between the toes and balls of the feet and said that heat applied to the stump was of limited benefit. The cold phantom pain could be triggered by a cold environment, but also by visually perceived cold (e.g., images of snow). He had not experienced the cold sensation prior to amputation (in terms of preamputation recall), and the stump itself was usually warm. Both stumps showed slightly increased cold perception on the distal lateral sides, on the right with additional dynamic al-
lodynia (without pinprick hyperalgesia). Following many unsuccessful treatments (includes opiates and anticonvulsants), the topical ambroxol 20 % cream managed to produce a relevant effect: the extreme cold sensation in the phantom regressed distinctly after approximately 15 min for several hours, and the phantom limb felt warmer. This effect has persisted unchanged for over 11 months now. No skin changes of any kind or any other side effects occurred on the stump, and the case report is documented on video.

Case report 3 (Neuropathic pain after total knee replacement)

This 58-year-old female patient received a total knee replacement in November 2010, after which considerable pains persisted. Clinical examination revealed an extensive dynamic allodynia and pinprick hyperalgesia medially as a sign of central sensitization. There were no signs of inflammation or movement-related pain. The existing treatment with buprenorphine 10 ug/h had been replaced by tapentadol because of the neuropathic nature of the pain, but this had proved no better in relieving the pain. After capsaicin 8 % plasters, the pain became slightly less frequent and of shorter duration. Since the knee pain was still persisting substantially by April 2014 and lidocaine plasters had also not been very helpful, topical ambroxol 20 % was used during the consultation. After just 15 min the patient reported clear pain relief: the burning and stabbing had subsided distinctly, while a ‘raging feeling’ in the knee disappeared almost completely. After applying the cream repeatedly over the next few months, she observed a reduction in pain for 4–6 h after approximately 30 min in each case, from an average of NRS 8/10 before treatment to 4/10 NRS (Fig. 2), and occasionally even down to NRS 1/10. With the most extreme pain (NRS 10/10), she found that the pain no longer returned to this level after the treatment. She has been applying topical ambroxol 20 % cream regularly for 11 months now, without any side effects or skin reactions. The case report is documented on video.

Case report 4 (Deafferentation pain)

The 38-year-old patient presented in April 2014 with deafferentation pains in the left arm after a plexus lesion (motorcycle accident in 1997), with subsequent mechanical allodynia in hand and forearm. A nerve graft proved unsuccessful, as did ketamine, gabapentin and a lidocaine infusion. Mirror therapy ended with a worsening of the pain for several days, while amitriptyline produced excessive sedation. Although cannabis (in connection with a clinical study) managed to reduce the pain by 60 %, it also resulted in substantial mental impairment. His medication now consisted of pregabalin 2 × 300 mg and duloxetine 60 mg. Slight innervation of the biceps muscle and shoulder elevation were possible, as was mental motor imagery to modulate the phantom pain. The patient experienced three types of pain in the arm: a ‘burning pain’ (started by the application of cold to the hand), a ‘crushing underlying pain’ (with no trigger) and ‘shooting tingling pains’. The pain intensity ranged from NRS 4–8/10, and there was no dynamic allodynia at the elbow. Sensations could be triggered in the phantom arm by cold stimuli to the subclavicular area and, as referred pains, from trigger points from the subclavian and pectoral muscles. Lidocaine plasters merely changed the nature of the pain, while trigger point treatments and tapentadol were not tolerated. A test was therefore initiated with topical ambroxol 20 % cream over the pectoral muscle, which successfully relieved the shooting and tingling pains in most cases from NRS 8/10 to 4/10. Since then the effect has been described as starting after approximately 15 min and persisting for 4–6 h, which proved crucially important particularly in falling asleep. However, the ‘deep underlying pain’ remained unaffected. The patient was especially happy about the spontaneous cessation, without triggers, of repeatedly occurring spasms and cramps, which enabled him to continue everyday tasks at these times. The patient’s descriptions are documented on video.

Case report 5 (Thoracic postherpetic neuralgia)

This 55-year-old patient presented in August 2008 with a postherpetic neuralgia on the right side of the chest (intensity 5/10) that had been present for 2 months. Step I and Step II analgesics only reduced the pain slightly, while lidocaine plasters proved very effective. Clinical examination revealed an almost circular, dynamic
alldynia at T6-8 on the right. He was initially treated successfully with increasing doses of gabapentin, and his night-time sleep improved with amitriptyline drops. Since other sedating substances were not possible and capsaicin 8% remained unsuccessful, the use of topical ambroxol 20% cream was tried in April 2012. Although this was unable to control the pain adequately on its own, it did manage to reduce the pain, starting after 30 min, from NRS 6/10 to 4/10 (Fig. 3) for a period of 4–6 h. The patient applies the cream for pain attacks and for those areas that are not well covered by the lidocaine plasters, which he continues to use. No adverse effects or skin reactions have been reported after what is now more than 3 years of use.

Case report 6 (mononeuropathy multiplex)

This patient presented in November 2013 with neuropathic pain in the arch of the left foot and multiplex neuritis as a result of vasculitis. He described two types of pain: a permanent underlying pain that interfered considerably with this everyday activities, and he also reported attacks of fierce pain, particularly in the evenings while watching television and at night, with an intensity up to NRS 8/10. Meanwhile he has been receiving treatment for depression with duloxetine 60 mg and lithium. Lidocaine plasters, peripheral analgesics and low-dose tilidine had already been tried without success. His night-time sleep was now deepened with amitriptyline drops, while the dosage of the tilidine was cautiously increased. Since the central nervous tolerance threshold had already been reached at the second visit in December 2013, a test was arranged with topical ambroxol. This proved highly effective: the patient reported a reduction in his evening, neuropathic pain from NRS 6/10 to 2/10, starting after 15 min and lasting for over 6 h. He was able to watch television without interruption, while his night-time sleep (after applying the cream immediately beforehand) was usually good. The zolpidem that he had been using for a long time was no longer necessary. When he was woken by pain, further application of the cream produced rapid pain relief. After 4 months of use, the permanent underlying pain during the day disappeared almost completely. A notable finding was the absence before treatment in this patient of any kind of alldynia or hyperalgesia in response to dynamic or pinprick stimuli or to cold or draughts of air. Having applied the cream for a period of 17 months now, the patient continues to use this treatment (Fig. 4) and has not reported any form of skin reactions or side effects to date. The patient’s statements have been documented repeatedly on video.

Case report 7 (Trigeminal postherpetic neuralgia)

After contracting a zoster infection of the V2 branch on the left in June 2014, this 91-year-old female complained of facial pain up to the intensity of NRS 8/10. Affective pain perception was considerable, and her night-time sleep was greatly disturbed. Diclofenac, tramadol and metamizole were not sufficient. Her night-time sleep was initially improved with a few amitriptyline drops, but an increase in the tramadol dosage failed due to constipation and sedation. Lidocaine plasters were effective, but had to be discontinued due to skin reactions. Since the dosage of gabapentin (100 mg) could only be increased to a limited extent and capsaicin 8% did not appear to be indicated in this elderly woman, ambroxol 20% cream was attempted. She has now been using this for 11 months and describes the effect as clearly pain-relieving, and felt that her cheek became ‘calmer’ approximately 15 min after each application. If she awoke at night, she was able to fall asleep again without pain sometime after a further application. This use has also been continued now for 10 months without any adverse effects and is documented on video.

Discussion

In this case series we describe, for the first time, patients whose neuropathic pain was relieved by the topical application of ambroxol. The pain reduction was between 33 and 100% (or between 2 and 8 NRS points; Table 1) and can therefore be considered as clinically relevant [12]. The findings can be explained by the sodium channel blockade produced by the substance.
Pharmacology, sodium channels and analgesic effect

Ambroxol is structurally similar to known local anaesthetics (‘Lofgren structure’) and, like these, binds to a specific binding site of the neuronal sodium channel [27]. It likewise blocks the sodium influx at voltage-dependent sodium channels, which leads to a reduction in the action potential frequency and thus in intraneuronal signal transduction.

However, certain properties suggest that ambroxol is of particular interest for the treatment of neuropathic pain. Firstly, it is very potent: ambroxol inhibits neuronal sodium channels at an approximately 40-fold lower concentration than lidocaine [47], thereby alleviating neuropathic pain after local [29] and systemic administration [46]. Secondly, in animal studies (and in contrast with lidocaine), it blocks nociceptor-specific sodium channels more potently than other neuronal sodium channels [47]. This may explain why four of our patients in whom lidocaine had previously proved ineffective nevertheless profited from ambroxol, which also produced a very powerful effect in animal models of neuropathic pain after systemic administration [14, 16], at least as strong as that of gabapentin and at dosages that can certainly be employed in clinical practice [14, 49].

Voltage-gated sodium channels (Nav, v = voltage gated) in nerve cell membranes play an important role in signal transduction, including in nociceptive neurons. Nine subtypes are described (Nav 1.1–1.9) [8]. The neurally expressed channels Nav 1.1, 1.2, 1.3, 1.6 and 1.7 are inhibited with high affinity by tetrodotoxin (TTX), the toxin of the pufferfish. The subtypes expressed predominantly in nociceptive neurons, that is Nav 1.8 and 1.9, are TTX-resistant [37]. Ambroxol blocks TTX-resistant sodium channels to a much greater degree than TTX-sensitive channels. The TTX-r-subtype Nav 1.8, which is preferentially blocked by ambroxol, is expressed particularly in spinal ganglion cells [2]. In nerve injuries, it appears to play an important role in the sensitization of nociceptors and thus contribute to the development and maintenance of neuropathic pain [3, 21, 26, 36]. It is thought to be particularly important in pain caused by cold [52]. A connection with the observation of ‘warming of the ice-cold phantom leg’ by the application of ambroxol to the stump of patient 2 remains to be seen. Nav 1.8 blockade also proved to be clearly analgesic in animal studies [11, 20]. So the improvements in our patients are credible not just on the basis of its continued use for prolonged periods.

**Na₉ 1.8 expression**

After nerve injuries, the redistribution or expression of sodium channels can occur in primary nociceptive neurons, with a resulting low threshold for triggering pain or for the hyperexcitability of DRG neurons. The subtype Na9, 1.8 is selectively expressed in nociceptive, sensory neurons [2, 6, 35, 52], but to a greater extent in nerve pain models [15, 44] and also in patients with neuropathic pain [24, 50]. This is also thought to be important in both animals and humans for the spontaneous activity in neuromas [5, 24, 36], which might have played a role in our amputated patients (case report 2). This may also explain the reduction in attacks of pain in all our five patients concerned. Increased Na9, 1.8 expression also occurs in diabetic polyneuropathy [30], small-fibre neuropathy [19], radiculopathies [17, 44], trigeminal neuralgia [41], but also in arthritis [39, 43] and bone metastases [33]. Na9, 1.8 blockade (e.g. produced by ambroxol) is also ultimately considered as an option for the treatment of neuropathic pain, because investigations with Na9, 1.8-free mice and Na9, 1.8-blocking substances have shown lower mechanical, thermal and visceral hyperexcitability in the animal model [1, 4, 11, 18, 20, 21, 45, 46].

**Clinical significance, safety and dosage**

Although ambroxol binds to the same local anaesthetic binding sites as lidocaine and amitriptyline, none of the substances used to date for neuropathic pain (includes local anaesthetics, antidepressants and anticonvulsants) have shown any relevant selectivity for the Na9, 1.8 channel comparable with that of ambroxol [27, 48]. The sodium-channel blockers used for treating pain, that is amitriptyline and carbamazepine, also do not block the channel types selectively [7, 42]. The selective blockade by ambroxol of the Na9, 1.8 channel type, which is not represented in the cardiac or central nervous systems, may therefore prove clinically beneficial. At least we did not receive any reports of corresponding adverse effects in any of our patients, including in those who have been using the substance topically for a long time now. This is also not surprising in that, despite the potent sodium channel blockade, the tolerability of ambroxol itself is very good after systemic administration: even intravenous doses as high as 1 g (for promoting prenatal lung maturation and the treatment of atelectasis) are well tolerated [51, 53]. There are even isolated reports of doses of up to 3 g a day being administered for up to 53 days [9, 28, 38] and the oral administration of 1.3 g ambroxol a day for 33 days [22]. After animal studies showed that high-dose ambroxol was clearly analgesic and well tolerated, Gaida et al. [14] presumed that these high dosages were probably necessary to produce analgesia. But with our topical treatments, we were able to show that even low-dose peripheral applications can have a clearly analgesic effect. Furthermore, no skin changes were observed by us, even though corresponding case reports exist [31].

In view of this interesting therapeutic approach, a number of other Na9, 1.8-blocking substances are currently being developed. However, its highly potent blockade, good bioavailability and very few side effects make ambroxol a particularly interesting substance, and one that should prove at least as useful as other sodium channel blockers for the treatment of chronic pain [27]. The safety and tolerability of systemically administered ambroxol—even at dosages on the gram scale—have been demonstrated over decades of clinical experience.

**Ambroxol and allodynia**

Apart from case report 6, all patients suffered from hypersensibility to mechanical stimuli which regressed in each case after topical ambroxol. Is this anti-allodynic, analgesic effect really conceivable?
Alldynia and hyperalgesia are considered to be neuroplastic phenomena of the spinal sensory system [25]. The Na\textsubscript{x}1.8 channel is detected mainly in C- or A-delta fibres and neurons of the posterior horn [2, 6, 35, 52], although it is also expressed in A-beta fibres [1, 4, 26, 34, 40]. But since chronic inflammation shifts the excitability of Na\textsubscript{x}1.8 toward hyperpolarization this contributes to the alldynia, which means that ambroxol blockade would therefore be particularly pain-relieving. In animal studies, ambroxol also suppressed alldynia to differing degrees after systemic administration: heat hyperalgesia by 100%, cold hyperalgesia and mechanical alldynia by approximately 75% [14]. The intrathecal administration also showed an anti-alldynic effect in animal experiments [32]. The observation that ambroxol also managed to reduce mechanical alldynia in rats with experimental induced inflammation by approximately 2/3 [4] suggests that the anti-alldynic, analgesic effect is not necessarily limited just to neuropathic pain. In the patients described in this article, at least, the alldynia in response to mechanical stimuli was alleviated in all cases by topical ambroxol, which can now be explained by the rationale described above.

Onset and duration of effect

The onset of Na\textsubscript{x}1.8 blockade in vitro by ambroxol starts within a few seconds and is concentration-dependent and fully reversible [27]. In paraplegic rats hypersensitivity to static mechanical stimuli was reduced after approximately 30 min for approximately 3 h [16]. Both findings correlate closely with the statements of our patients (Fig. 2), according to which the effect started between 15 and 30 min after application. In case report 1, the effect was frequently even said to appear ‘immediately’ and persisted for well over 6 h in this patient. When applied repeatedly, topical lidocaine 5% often shows an increasing effect over time, both clinically and according to the literature [29]. A comparable result was also reported by our patient 6, who felt that his underlying pain had almost disappeared after 4 months of use. The significance of these observations needs to be explored in greater depth.

Limitations

This case series describes (in an open-label, uncontrolled design) examples of the successful outcome of a topical treatment with ambroxol 20% in patients with neuropathic pain. It is therefore still subject to numerous possible sources of error, from placebo effects to temporary therapeutic successes with no long-term significance, even though some of the treatments are documented here for several years. Although these were pooled, by way of example, for many other patients, they were compiled only retrospectively and, in this article, only for neuropathic pain. Since amount and depth of ambroxol’s skin penetration from the cream are not known, it cannot be stated conclusively whether the analgesic effects are based solely on its local action or on partial systemic effects. However, the latter would probably already have come to light clinically in view of the widespread use of ambroxol for respiratory disorders. An improvement to this initial formulation would also doubtless be possible. Therefore the observations do not yet allow any general conclusions to be drawn concerning the degree of efficacy in humans. Systematic and controlled studies on these questions are needed.

Conclusion for practice

The expectorant ambroxol also acts as a very potent local anaesthetic. It significantly and preferentially blocks the nociceptively relevant sodium channel subtype Na\textsubscript{x} 1.8 to a greater extent than all other local anaesthetics. Increased Na\textsubscript{x}1.8 expression is detected in neuropathic pain and is almost exclusively limited to sensory pain fibres. Its blockade is therefore considered to be a useful, simple concept for pain management with few side effects. In the case reports presented in this article, this concept was successfully implemented, for the first time, in the form of topical ambroxol 20% cream in patients with severe neuropathic pain. This preparation has meanwhile been used for over 4 years without any undesirable effects. Since the development of new substances for the treatment of neuropathic pain is very arduous and time-consuming, we believe that there is an urgent need for further extensive research on the clinical use of well-known traditionally used substances with clear anti-nociceptive effects and positive side effect profiles, for example ambroxol. In view of the considerable distress caused by failed treatments the use of such substances should also be considered in connection with individual therapeutic trials.

Corresponding address

PD Dr. K.-U. Kern
Institut für Schmerzmedizin/Schmerzpraxis Wiesbaden, Sonnenberger Str. 68 65193 Wiesbaden
dr.kern@schmerzpraxis-wiesbaden.de

Contributions. The first author treated and documented the patients described in this paper, while the second author characterized the mechanism of action of ambroxol as a sodium channel blocker and investigated its effect in the preclinical pain models. The authors jointly drafted and critically discussed the manuscript. They would like to thank the pharmacist Martin Hofmann, Aukamm Apotheke Wiesbaden, for preparing the ambroxol cream formulation.

Compliance with ethical guidelines

Conflicts of interest. In the past 3 years Dr. K.-U. Kern has worked as a consultant and/or expert for the following companies: Astellas, Berlin Chemie, Boehringer Ingelheim, betapharm, Grünenthal, Hexal, Lilly, MSD, Mundipharma and Reckitt Benckiser. Dr. T. Weiser is an employee of Boehringer Ingelheim.

Open Access This article is distributed under the terms of the Creative Commons Attribution License which permits any use, distribution, and reproduction in any medium, provided the original author(s) and the source are credited.

References

1. Abrahamsen B, Zhao J, Asante CO et al (2008) The cell and molecular basis of mechanical, cold, and inflammatory pain. Science 321:702–705
2. Akopian AN, Sivilotti L, Wood JN (1996) A tetrodotoxin-resistant voltage-gated sodium channel expressed by sensory neurons. Nature 379:257–262
3. Akopian AN, Souslova V, England S et al (1999) The tetrodotoxin-resistant sodium channel SNS has a specialized function in pain pathways. Nat Neurosci 2:541–548
4. Belkouch M, Dansereau MA, Tetreault P et al (2014) Functional up-regulation of Na+, 1.8 sodium channel in the human neocortex fibers subjected to chronic peripheral inflammation. J Neuroinflammation 11:56

5. Black JA, Nikolajsen L, Kroner K et al (2008) Multiple sodium channel isoforms and mitogen-activated protein kinases are present in painful human neovascularisation. J Neurol 245:604–613

6. Blair NT, Bean BP (2002) Roles of tetrodotoxin-sensitive and -insensitive Na+ currents in the action potentials of nociceptive sensory neurons. J Neurosci 22:10277–10290

7. Brau ME, Dreimann M, Olchewski A et al (2001) Effect of drugs used for neuropathic pain management on tetrodotoxin-resistant Na+ currents in rat sensory neurones. Anesthesiology 94:137–144

8. Catterall WA (2012) Voltage-gated sodium channels. In: The Neuroscience (Weissleder C, ed) 3rd edn, p 452. Springer, New York

9. Chiara O, Padalino P, Guadalupi P, Bigallego I, Nespoli A (1983) The prevention of postoperative pain by Ambroxol: a double-blind test in the effects of ambroxol on pulmonary gas exchange. Gerontology Chirurgicae Commentaria 6:1–6

10. De Mey C, Piel H, Kolisch S et al (2008) Efficacy and safety of ambroxol lozenges in the treatment of acute uncomplicated sore throat. EBM-based clinical documentation. Arzneimittel-Forschung 58:557–568

11. Eckers J, Jayamanne A, Vaughan CW et al (2006) mu-conotoxin MrVIB selectively blocks Na+, 1.8 sensory neuronal specific sodium channels and chronic pain behavior without motor deficits. Proc Natl Acad Sci USA 103:17030–17035

12. Farrar JT, Pritchett YL, Robinson M et al (2010) The clinical importance of changes in the 0–10 numeric rating scale for worst, least, and average pain intensity: analyses of data from clinical trials of duloxetine in pain disorders. J Pain 11:109–118

13. Ferrante FM, Paggio J, Cherukuri S et al (1996) The analgesic response to intravenous lidocaine in the treatment of neuropathic pain. Anesth Analg 82:91–97

14. Gaida W, Klinder K, Andrit K et al (2005) Ambroxol, a Na+, 1.8-prefering Na+ channel blocker, effectively suppresses pain symptoms in animal models of chronic, neuropathic and inflammatory pain. J Pharmacol Pharmacother 49:1220–1227

15. Gold MS, Weinreich D, Kim CS et al (2008) Distribution of Na(V)1.8 sodium channels in painful human neuroma-evidence from immunocytochemistry. Acta Neurochir 140:803–810 (discussion 810)

16. Kuner R (2010) Central mechanisms of pathological pain. Nat Med 16:1258–1266

17. Lai J, Gold MS, CD (2000) Inhibition of neuropathic pain by decreased expression of the tetrodotoxin-resistant sodium channel, Na(+), 1.8. Pain 95:143–152

18. Leffler A, Reckzeh J, Nau C (2010) Block of sensory neuronal Na+ channels by the secoerycine ambroxol is associated with an interaction with local anesthetic binding sites. Eur J Pharmacol 630:19–28

19. Luerti M, Laarzarin A, Corbella E et al (1987) An alternative to steroids for prevention of respiratory distress syndrome (RDS): multicenter controlled study to compare ambroxol and betamethasone. J Perinat Med 15:227–238

20. Meier T, Wassen G, Faust M et al (2003) Efficacy of lidocaïne patch 5% in the treatment of focal peripheral neuropathic pain syndromes: a randomized, double-blind, placebo-controlled study. Pain 106:151–158

21. Merit T, Gunnes Y (2012) Antinoceptive activities of lidocaine and the Na+, 1.8 blocker ab03467 in diabetic rats. J Am Assoc Lab Anim Sci 51:579–585

22. Monzon S, Del Mar Garces M, Leuzan A et al (2009) Ambroxol-induced systemic contact dermatitis confirmed by positive patch test. Allergol Immunopathol 37:167–168

23. Moon JH, Song S, Yoon ST et al (2002) The differential effect of intrathecal Na+, 1.8 blockers on the induction and maintenance of capsaicin- and peripheral ischemia-induced mechanical allodynia and thermal hyperalgesia. Anesth Analg 114:215–223

24. Qu F, Jiang Y, Zheng H et al (2012) Increased expression of tetrodotoxin-resistant sodium channels Na+, 1.8 and Na+, 1.9 within dorsal root ganglia in a rat model of bone cancer pain. Neurosci Lett 512:61–66

25. Renganathan M, Cummins TR, Waxman SG (2001) Contribution of Na(V)1.8 sodium channels to action potential electrogenesis in DRG neurons. J Neurophysiol 86:629–640

26. Roza C, Laird JM, Soulos V et al (2003) The tetrodotoxin-resistant Na+ channel Na+, 1.8 is essential for the expression of spontaneous activity in damaged sensory axons of mice. J Physiol 550:921–926

27. Rush AM, Cummins TR, Waxman SG (2007) Multiple sodium channels and their roles in electrogensis within dorsal root ganglion neurons. J Physiol 579:1–14

28. Schilling GJ (1992) Ambroxol in the treatment of ARDS in polytraumatized patients - a report. Klinische Wochenschrift 70:12–18

29. Schuelert N, McDougall JJ (2012) Involvement of Na+, 1.8 sodium ion channels in the transduction of mechanical pain in a rodent model of osteoarthritis. Arthritis Res Ther 14:85

30. Sheds SD, Ahn HS, Yang Y et al (2012) 1.8 expression is not restricted to nociceptors in mouse peripheral nervous system. Pain 153:2017–2030

31. Siqueira SR, Alves B, Malpàrdita HM et al (2009) Abnormal expression of voltage-gated sodium channels Na+, 1.7, Na+, 1.3 and Na+, 1.8 in trigeminal neurones. Neuroscience 164:573–577

32. Song HJ, Ham SS, Shin YK et al (2000) Amtripyrine modulation of Na+ channels in rat dorsal root ganglion neurons. Eur J Pharmacol 401:297–305

33. Strickland IT, Martindale JC, Woodhams PL et al (2008) Changes in the expression of Na+, 1.7, Na+, 1.8 and Na+, 1.9 in a distinct population of dorsal root ganglia innervating the rat knee joint in a model of chronic inflammatory joint pain. Eur J Pain 12:564–572

34. Thakor DK, Lin A, Matsuka Y et al (2009) Increased peripheral nerve excitability and local 1.8 mRNA up-regulation in painful neuropathy. Mol Pain 5:14

35. Veneroni O, Maj R, Calabresi M et al (2003) Anti-alloiodynie effect of NW-1029, a novel Na+(+) channel blocker, in experimental animal models of inflammatory and neuropathic pain. Pain 102:17–25

36. Villareal CF, Sachs D, Cunha FQ et al (2005) The role of Na(V)1.8 sodium channel in the maintenance of chronic inflammatory hypernociception. Neurosci Lett 386:72–77

37. Weiser T (2006) Comparison of the effects of four Na+ channel analogics on TTX-resistant Na+ currents in rat sensory neurons and recombinant Na+, 1.2 channels. Neurosci Lett 395:179–184

38. Weiser T (2006) Na+, 1.8 channel blockade as an approach to the treatment of neuropathic pain. Drugs Fut 31:597–601

39. Weiser T, Wilson N (2002) Inhibition of tetrodotoxin (TTX)-resistant and TTX-sensitive neuronal Na+(+) channels by the secoerycine ambroxol. Mol Pharmacol 62:433–438

40. Yangou Y, Birch R, Sangameswaran L et al (2000) SNS/PN3 and SNS2/NaN sodium channel-like immunoreactivity in human adult and neonate innervated sensory fibres. FEBS Lett 467:249–252

41. Zhang Q, Wu QG, Huang XM et al (2013) Prevention of respiratory distress syndrome in preterm infants by antenatal ambroxol: a meta-analysis of randomized controlled trials. Am J Perinatol 30:529–536

42. Zimmermann K, Leffler A, Babes A et al (2007) Sensory neuron sodium channel Na+, 1.6 is essential for pain at low temperatures. Nature 447:855–858

43. Zwissler B (2002) Pharmakotherapie bei akutem Lungenversagen. J Anaesth Intensivbehandlung 9:59–67