Differential effects of epidermal growth factor receptor inhibitors in a single patient with neuropathic pain

Marte Grønlie Cameron, Christian Kersten

SUMMARY
Neuropathic pain (NP) represents an unmet medical need, where analgesic responses to different epidermal growth factor receptor inhibitors (EGFR-is) have been described. The human EGFR family of receptors consists of four members (human epidermal growth factor receptor, HER 1–4), signalling via different homodimer and heterodimer combinations. A 52-year-old man was treated with the EGFR-I cetuximab in a trial of severe NP. Pain scores decreased dramatically after blinded cetuximab, but not after placebo. On pain recurrence after the trial, he was prescribed the oral EGFR-Is erlotinib, gefitinib, and lapatinib without relief. However, treatment with the pan-HER-inhibitor afatinib was effective. After 4 years on afatinib, pain control remains excellent with manageable side effects. This is the first reported observation of differential effects of EGFR-is on NP in the same patient and the first report describing NP relief with afatinib. Further understanding of the underlying pathophysiology could lead to development of EGFR-is specifically targeting NP.

BACKGROUND
Neuropathic pain (NP) is a clinical description of a pain state resulting from damage to the somatosensory nervous system.1 The pain is lancinating, electric shock-like and burning in character, associated with tingling and crawling sensations. Its severity, chronicity and the poor benefit to side effect ratio of pharmacotherapy for NP frequently lead to diminished physical and psychological functioning among sufferers.2,3 The prevalence of moderate to severe chronic NP in the western world is estimated to be at least 5%–8% and the global burden is escalating due to the ageing, growing population with rising prevalence of the chronic diseases causing NP, including postherpetic neuralgia, diabetic neuropathy, chemotherapy-induced peripheral neuropathy, lumbar radiculopathy and chronic postsurgical pain.4,5

The human epidermal growth factor receptor (EGFR) (human epidermal growth factor receptor, HER) family of cell surface receptors is made up of four members: ErbB1 (HER1, EGFR), ErbB2 (HER2), ErbB3 (HER3) and ErbB4 (HER4) (see figure 1). Each receptor consists of extracellular, transmembrane and intracellular domains. Binding of ligands to some of the extracellular domains leads to receptor dimerisation and conformational changes in the intracellular tyrosine kinases, resulting in receptor autophosphorylation and activation of downstream signal transduction pathways that regulate cellular functions.6 The concept of receptor dimerisation explains why HER2 can be active despite not having a known activating ligand and why HER3 can bind ligands without activating its own kinase.7

Both orally administered tyrosine kinase inhibitors (TKIs) and intravenously administered monoclonal antibodies have been developed to inhibit the EGFR. Several of these are approved and have been in use in oncology for more than a decade.8 Numerous TKIs, which target the intracellular domains of different transmembrane receptors have been under development for the treatment of malignancies. Some of them, such as erlotinib and gefitinib, are designed to specifically target the intracellular tyrosine kinase domain of the EGFR and are primarily used to treat non-small cell lung cancer.9 10 Since cancers develop mutations in the tyrosine kinase domains of their EGFRs, second and third-generation oral EGFR-Is have been developed to overcome this.

Furthermore, TKIs designed to inhibit specific HER family heterodimers have also been developed. One example is lapatinib, a drug used to treat breast cancer, which binds to the EGFR/HER2 protein kinase domains.11 Afatinib inhibits all four HER kinases and is used in lung cancer.

The monoclonal antibody cetuximab inhibits functioning of the EGFR by targeting extracellular EGFR ligand binding domains.12 It has been licensed since 2004 to treat cancers of the head and neck and colorectal cancer.8

Side effects of EGFR-inhibitors are well described, based on their use in millions of cancer patients for over 15 years. Dry skin, acneiform rash and diarrhoea are most frequent side effects and feasibility of using EGFR-Is in the setting of NP presupposes supportive care and close follow-up.13

A serendipitous finding that cetuximab, independent of its anticancer effect, led to rapid relief of NP in a patient with malignant invasion of pelvic nerves sparked interest in the notion that it may also have therapeutic potential in NP.14 Since publication of the initial case, at least 84 additional clinical cases, including both benign and malignant causes of NP that were relieved by EGFR-Is have been reported.15–18 Analgesic responses to both antibodies (cetuximab and panitumumab) which work extracellularly and intracellular tyrosine kinase antagonists (erlotinib and gefitinib) are described.
Although the mechanism behind this phenomenon is not understood, it has clinically appeared to be a class effect. Chronic pain has recently been associated with mutations in the EGFR. Several papers describe a reduction of nocifensive behaviour in rodents with NP after administration of EGFR inhibitors. One of these comprehensive studies convincingly demonstrates colocalisation of and increased molecular interaction between EGFR and HER2 on pain fibres in NP models as well as a synergistic role of combined EGFR and HER2 inhibition in NP relief. Another research group has suggested that EGFR/HER4 heterodimers might be of importance. Together, clinical and preclinical findings point to the EGFR as a plausible target for treatment of NP.

Based on this, we performed a randomised, placebo-controlled double-blind proof-of-concept trial of cetuximab in patients with severe, treatment-refractory NP due to non-malignant compressed peripheral nerves (n=7) or complex regional pain syndrome (n=7). This so-called NoTOPain trial (NoTOPain: Novel Treatment Option for neuropathic Pain) reported relatively high response rates among treatment-resistant patients and the magnitude of pain relief seen in those patients who benefited was encouraging. Eight of the trial participants chose to continue with oral EGFR-Is after study completion. The current case report describes the clinical course of one of the trial participants during and for 48 months after the NoTOPain trial. This particular patient had convincing pain relief after cetuximab treatment within the placebo-controlled NoTOPain trial. Thereafter, he tried four different oral EGFR-Is, with mixed results.

**CASE PRESENTATION**

A 52-year-old previously healthy man with an 11-year history of recurrent, bilateral carpal tunnel syndrome presented with a 22-month history of severe NP in his right hand. The same condition was present on the left side, although to a lesser degree.

The patient had undergone multiple decompressive surgeries (two on the left side, three on the right) during the 11 years prior to the presentation. After each operation, he experienced symptom relief lasting from 6 to 9 months, followed by gradual recurrence. His last operation on his right wrist, 30 months prior to presentation, resulted in roughly 8 months of pain relief. Further surgical intervention was not recommended on pain recurrence. Various treatments were tried without satisfactory pain relief (see table 1). The pain was eventually deemed chronic and treatment refractory.

The patient described the pain as constantly present, with intensity usually ranging between 5 and 8 on a 0–10 Numeric Rating Scale (NRS). The pain was located primarily distal to the wrists, in the area of median nerve innervation. Although there were both pain and sensory symptoms beyond this, both in the hand and lower forearm, he did not describe it as radiating. The pain was associated with numbness, tingling and prickling sensations. The most striking and clinically debilitating aspect of the patient’s pain was the degree to which it was aggravated by cold, in terms of both cold allodynia and the effect of ambient temperature. He scored 22/38 on the PainDETECT questionnaire, indicating >90% probability that the pain had a neuropathic component.

MRI of the right wrist revealed scar tissue in close proximity to the median nerve. Neurological examination found cold and light touch allodynia as well as decreased sensation distal to the wrist scars in both hands. These findings included but also went beyond the distribution of the median nerves. Findings were similar on both sides, but more severe on the right. Neurography confirmed damage to the right median nerve at the level of the carpal tunnel (the left side was not tested). Neurological examination was otherwise normal.

**Cetuximab**

The patient was referred to us for inclusion in the NoTOPain trial which tested EGFR-inhibition in patients with chronic, treatment-refractory NP. He was randomised to receive a single dose of blinded placebo first, followed by a single dose of...
blinded cetuximab and then one open-label cetuximab infusion (see figure 2).

The patient’s pain scores (self-reported daily, on a 0–10 Numeric Rating Scale) showed a clinically significant decrease after blinded cetuximab, but not after placebo (see figure 3). Only the pain scores for the right hand (the most severely affected) were registered in the NoTOPain trial, although improvement was analogous on the left side.

The patient described the pain relief he experienced after cetuximab as having completely transformed his quality of life. Specifically, he was able to sleep right through the night, without being woken up by pain that he otherwise experienced several times each night when his hands were exposed to cold as they slipped out from under the bed covers. Being well-rested improved his concentration and capacity to work. Ability to tolerate cold meant that he no longer required warm gloves and could work outdoors in all types of weather. He was more flexible and functional in general, able to do spontaneously things he enjoyed with family and friends, no longer having to consider pain and disability. He was able to stop taking all other pain medications and no longer needed to use the transcutaneous electrical nerve stimulation apparatus at bedtime.

Mechanical allodynia was assessed during the NoTOPain trial using standardised nylon monofilaments before and after the patient was treated with blinded cetuximab. He was asked to identify the area on his right hand with maximum pain. The monofilaments, in order of increasing size and target force, were then sequentially applied there, and he was asked to indicate when the filament triggered a painful response (using a non-painful area on the opposite side as a control). Just prior to the start of the cetuximab infusion, the patient scored 7 on the 0–10 NRS for ‘pain right now’. The first filament to provoke a pathological pain response at that time had a target force of 60 g (size 5.88). The test was repeated in the same place 2 hours after the cetuximab infusion, when his pain score was 1 on the same NRS (see figure 4). At that time, none of the monofilaments provoked...
a pathological pain response (the largest filament tested having target force 300g (size 6.65)). The test was repeated again the following day, with a pain score of 2 and again allodynia was not provoked by any of the monofilaments.

**Erlotinib**

The patient experienced near-complete pain-relief after one of the two blinded study infusions which at study closure was revealed to be cetuximab. The open-label cetuximab infusion elicited the same response. He was, therefore, prescribed the oral EGFR-inhibitor erlotinib 150 mg daily, which he started taking once pain recurrence was well-established after his last cetuximab infusion. The patient described partial, transient improvement in NP after the first dose of erlotinib but felt that it never measured up to the dramatic improvement that he had experienced after the two intravenous cetuximab infusions in the trial.

The question of whether lack of efficacy of erlotinib could be a dosing question was raised so after 2 weeks taking 150 mg he increased to 300 mg daily. He took 300 mg daily for a total of 6 days but stopped because he was then convinced that erlotinib in fact had no effect at all, that is, he reported that the pain was back to the levels he had before start of the trial.

**Gefitinib**

After a further month of pain, treatment with the oral EGFR-inhibitor gefitinib was started. He took 250 mg gefitinib daily for 3 weeks without any change in NP severity.

**Afatinib**

Having failed both oral EGFR-Is that had shown benefit in several patients before,15 16 a afatinib was prescribed, based on the hypotheses that a germline EGFR mutation may cause resistance to erlotinib and gefitinib, or that a broader inhibition (afatinib is a pan-HER inhibitor) may reproduce the effect he had experienced after the two cetuximab infusions in the NoTOPain trial. Therefore, 6 months after his last dose of cetuximab in the trial, the patient was prescribed afatinib 40 mg daily. He began noticing improvement in his NP on day 4 of this treatment.

There was a gradual decrease in pain scores over the subsequent 2–3 weeks until a new plateau was reached (see figure 3).

Again, pain relief was accompanied by improved function and enjoyment of life, as indicated by Brief Pain Inventory scores (see table 2).

After approximately 3 months, the patient’s afatinib dose was reduced to 20 mg daily in order to test if that was a sufficient dose for pain control. The patient’s impression is that afatinib is not fully as effective as he remembers intravenous cetuximab but that 20 mg daily of afatinib is virtually as effective as 40 mg, with lesser side effects. During the treatment breaks, his pain typically recurs after 3 days.

Afatinib may inhibit EGFRs with mutations that render them resistant to erlotinib and gefitinib. The patient’s blood was therefore analysed for germline EGFR mutations with allele-specific PCR (Cobas EGFR mutation test v2, Roche), covering 42 different mutations in exons 18, 19, 20 and 21. None of these mutations were found.

**Lapatinib**

A year after starting afatinib, the patient was offered a trial of the HER1/HER2 inhibitor lapatinib 1250 mg daily in an attempt to reduce gastrointestinal and cutaneous side effects. He took this treatment for 7 days without noticing any improvement in his NP and therefore reverted to afatinib.

**Side effects**

The patient has intermittently taken tetracycline 500 mg two times per day to treat acneiform rash while under treatment with the various EGFR-Is. He has had maximum grade 2 dry skin and mucositis. His most bothersome side effect has been grade 2 diarrhoea, for which he has taken loperamide as needed, with partial effect.

**OUTCOME AND FOLLOW-UP**

Currently, 4 years after starting afatinib, he characterises his pain as completely gone as long as he takes 25 mg of afatinib daily. He has found that this dose gives him an effect to side-effect balance that results in the best possible quality of life. Based on
the experience with this case, an additional five patients who were using erlotinib (n=4) or gefitinib (n=1) have tried afatinib and reported better effect, tolerability or both.

DISCUSSION
This is the first time different effects of EGFR-Inhibitors on NP have been described in the same patient. This is also the first report describing NP relief with afatinib.

We have previously reported NP relief by the intravenous EGFR-Inhibitors cetuximab and panitumumab, and the oral TKIs erlotinib and gefitinib. However, in all previously reported cases, pain responses were similar across EGFR-Inhibitors and in accordance with their established pharmacokinetics.

This case report resembles an N=1 study in that the same patient received four different EGFR-Inhibitors in series, with pain recurrence both between the different EGFR-Inhibitors tested and during treatment breaks between doses of cetuximab and afatinib. In contrast to a planned N=1 study, however, this patient’s treatment course was guided not by a protocol, but by clinical decisions based on responses to the various EGFR-Inhibitors he was given along the way.

A strength of this case report is the prospective capture of patient-reported outcomes of pain and function. Although not generalisable, the observations support the notion of EGFR-I as a viable treatment option for NP, particularly since this otherwise treatment-refractory patient has continued to take afatinib for 4 years (to date), in order to maintain a better quality of life.

The fact that a patient whose NP responded to cetuximab but not to erlotinib, gefitinib or lapatinib, responded to afatinib is hypothesis generating.

Biological hypotheses
Better understanding of the biological mechanisms underlying the differential effects described in this case could potentially unlock a key to developing EGFR-Inhibitors specifically targeting NP, with a conceivably larger therapeutic window.

In principle, these biological mechanisms could be located at the level of the (1) EGFR ligands, (2) transmembrane receptors or (3) downstream of the receptors. Physiological or pathological mechanisms of resistance could also occur on all these levels. The fact that afatinib inhibits all HER receptors may indicate a broader effect on more extracellular ligands and/or on more intracellular downstream signalling pathways. However, it is important to note that several mechanisms could occur simultaneously, change over time and, importantly, be context dependent.

EGFR ligands
Seven EGFR ligands have been identified, including epiregulin which is involved in the generation of NP. The way each of these ligands interact with the EGFR determines their differential effects. Non-EGFR HER ligands such as neuregulins are also implicated in the pathophysiology of NP. Thus, one can speculate that different HER ligands are involved in the pathophysiology of the individual NP patients. If so, this would also be mirrored by their respective susceptibilities to inhibition by the different EGFR-Inhibitors.
Transmembrane receptors
If this patient’s NP is driven by signalling via an EGFR heterodimer, it follows that extracellular binding of cetuximab to the EGFR would inhibit heterodimer formation regardless of which of the other HERs it is coupling with. This could explain why erlotinib and gefitinib, both highly selective EGFR-Is with an intracellular mode of action, were not effective, while the pan-HER-inhibitor afatinib is. This points to an additive or even synergistic effect of certain HER-receptor inhibitor combinations, which has been hypothesised and shown in rodent models. As indicated in figure 1, afatinib binds to the EGFR, HER2 and HER4. By doing so, it irreversibly inhibits signalling of the entire HER family of receptors, including HER3, to which it does not bind directly.

Lapatinib, a dual inhibitor of the EGFR and HER2 receptors has been shown to inhibit pain in rodent models. It has, however, also been shown to strengthen the inactive monomeric form of the EGFR, thereby preventing its dimerisation. The fact that the patient did not experience pain relief while taking lapatinib, thus raises the question as to the extent to which the EGFR or relevant heterodimer, was being inhibited by that drug, since recent preclinical findings demonstrate a strong synergistic analgesic effect when both EGF and HER2 receptors are inhibited. Interestingly, the relative half maximal inhibitory concentrations of these drugs to inhibit the EGF and HER2 receptors favours EGFR-inhibition by afatinib compared with lapatinib, which may explain the observed differences in pain responses in our patient.

Intracellular downstream signalling
Net outcome of EGFR-I depends on cellular context. A meta-analysis of 20 individual studies comprising 346 microarrays reported that different EGFR-Is had overlapping but quite distinct transcriptional results in target cells. To our knowledge, no such studies have been performed on damaged peripheral nerve cells. This could be relevant in our patient.

Physiological or pathological mechanisms of resistance
Many different germline and acquired resistance mechanisms against EGFR-Is have been described in the context of cancer therapy. Such mechanisms could potentially also play a role in the differential pain relief observed in our patient.

The patient could in theory be a carrier of:
1. A germline mutation known to be targeted by afatinib (the patient has tested negative for these).
2. A rare germline mutation, where afatinib works.
3. Certain single-nucleotide polymorphisms (SNP) involved in pain syndromes, where afatinib is effective but undocumented.
4. An acquired somatic mutation or SNP in the peripheral sensory nerves that involves mechanisms 1–3.

Other acquired resistance mechanisms against EGFR-I include HER2 upregulation, activation of bypass signalling and non-coding RNA. Our patient could in theory have developed one or more such resistance mechanisms. However, of the more than 85 patients that we know have been treated with EGFR-Is for NP, acquired resistance against EGFR-inhibition has been observed only once, after one and a half years of continuous treatment with EGFR-I (unpublished observation).

Psychological hypothesis
The patient experienced the same magnitude of near-immediate pain relief after both blinded and open-label cetuximab infusions within the NoTOPain trial. His pain was not relieved by matched placebo (see figure 3). The rapid, although transient pain relief that began after a single dose of erlotinib was most likely a placebo response. Erlotinib reaches maximum plasma concentration between 0.5 and 4 hours after ingestion and the terminal half-life is 1.5–3 hours. Accordingly, it is not impossible to experience rapid pain relief as described. However, experience in other patients indicates a time to relevant treatment effect of several days when using oral EGFR-Is. The long-lasting effect of afatinib argues against a placebo effect, particularly when the two preceding TKIs (erlotinib and gefitinib) and later lapatinib, were ineffectual.

Limitations
There are obvious limitations to the validity and generalisability of subjective findings reported by a single patient. However, the value of this case lies in its hypothesis-generating role. Lack of translational elements (other than a germline mutation analysis of the EGFR) hinder further validation of the hypotheses presented, including the relevance of heterodimer signalling. Further studies in animal models should explore these avenues, which could subsequently be verified in human series.

Future perspectives
Not all EGFR-Is are created equal. We have previously proposed a class effect for EGFR-Is in the treatment of NP and by large this still seems valid. However, this particular case may indicate that there are nuances of the EGFR/NP association that may open up avenues for drug development.

Although common, cutaneous side effects of EGFR-Is are usually mild and acceptable for patients treated for severe and otherwise treatment resistant NP. However, skin reactions may hamper long-term use of EGFR-Is in patients with less severe NP. If a relationship exists between different HER family heterodimers and toxicity and/or therapeutic effect of anti-EGFR-directed therapy against NP, this could potentially lead to more targeted therapy, reducing toxicity while maintaining or even increasing the analgesic effect.

Learning points
- Several epidermal growth factor receptor inhibitors (EGFR-Is) have previously been reported to relieve neuropathic pain (NP), indicating a possible class effect.
- NP in this case responded differently to different EGFR-Is.
- This case illustrates for the first time that the EGFR-I afatinib can relieve NP.
- Novel observations in this case are supported by preclinical research and indicate a possibility to develop an EGFR-I with an optimal profile to specifically target NP.

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Competing interests The authors have filed an international patent application according to the Patent Cooperation Treaty, for targeting the EGFR in neurological disorders. CK has, since May 2020 been employed by Roche Norway AS which owns erlotinib.

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