TRANSLATIONAL SCIENCE

Active immunisation targeting nerve growth factor attenuates chronic pain behaviour in murine osteoarthritis

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ABSTRACT

Objectives Nerve growth factor (NGF) has emerged as a key driver of pain in osteoarthritis (OA) and antibodies to NGF are potent analgesics in human disease. Here, we validate a novel vaccine strategy to generate anti-NGF antibodies for reversal of pain behaviour in a surgical model of OA.

Methods Virus-like particles were derived from the cucumber mosaic virus (CMV) and coupled to expressed recombinant NGF to create the vaccine. 10-week-old male mice underwent partial meniscectomy to induce OA or sham-surgery. Spontaneous pain behaviour was measured by Linton incapacitation and OA severity was quantified using OARSI histological scoring. Mice (experimental and a sentinel cohort) were inoculated with CuMV\textsuperscript{tGF} (Vax) or CuMV\textsuperscript{tGF} (Mock) either before surgery or once pain was established. Efficacy of anti-NGF from the plasma of sentinel vaccinated mice was measured in vitro using a neurite outgrowth assay in PC12 cells.

Results Anti-NGF titres were readily detectable in the vaccinated but not mock vaccinated mice. Regular boosting with fresh vaccine was required to maintain anti-NGF titres as measured in the sentinel cohort. Both prophylactic and therapeutic vaccination demonstrated a reversal of pain behaviour by incapacitance testing, and a meta-analysis of the two studies showing analgesia at peak anti-NGF titres was highly statistically significant. Serum anti-NGF was able to inhibit neurite outgrowth equivalent to around 150 ng/mL of recombinant monoclonal antibody.

Conclusions This study demonstrates therapeutic efficacy of a novel NGF vaccine strategy that reversibly induces neutralising anti-NGF antibodies and suppresses pain behaviour in surgically induced murine OA.

What is already known about this subject?
► Nerve growth factor (NGF) is a validated target for pain in human and mouse OA.
► Neutralising antibodies to NGF show therapeutic efficacy in Phase III clinical studies.

What does this study add?
► Here, we demonstrate efficacy of an NGF vaccine that reversibly induces neutralising anti-NGF antibodies and suppresses pain behaviour in murine OA.

How might this impact on clinical practice or future developments?
► Vaccination potentially offers a tunable, cheaper and easier to use alternative to biological therapy in patients.

OA is the most prevalent joint disease costing approximately 1%–2.5% of the gross domestic product of developed countries. Greater than 75% of patients experience pain on a daily basis. Current standard therapies for pain relief, such as non-steroidal anti-inflammatory drugs (NSAIDs) and opioids are limited by their modest efficacy and long-term safety. In the last decade, nerve growth factor (NGF), a key pain sensitisier, has emerged as a promising target for OA pain. In humans, neutralising antibodies to NGF significantly suppress pain associated with late-stage OA. However, biological therapy in OA is likely to be limited by cost and by treatment failure due to anti-drug antibodies. Active immunisation targeting NGF represents an attractive alternative to deliver effective analgesia, while potentially providing a more economically sustainable substitute for patients. The latter is particularly the case as biosimilars replace proprietary products.

Chronic pain in late OA can be modelled using surgical models of joint destabilisation in mice. Spontaneous pain behaviour is assessed by differential weight distribution of the hind limbs using incapacitance testing. Following joint destabilisation, mice display two phases of pain behaviour: one immediately following surgery (postoperative pain) and a second late phase that starts between weeks 7 and 11 after surgery and which is progressive (online supplementary figure 1a) with worsening joint destruction (online supplementary figure 1b, c). Both phases of pain behaviour correspond to an increase in NGF expression in the joint (online supplementary figure 1d, e) and can be neutralised by delivery of NGF’s soluble receptor (TrkAd5).

Immunisation against self-proteins can be achieved by displaying the antigen of interest on virus-like particles (VLPs). Due to their optimal size and geometry, VLPs can effectively transit to
draining lymph nodes to drive antigen-dependent immunogenicity. Antigens are arranged as a repetitive array on the particles’ surfaces via genetic fusion or chemical conjugation to generate good polyclonal antibody responses without breaking T cell tolerance. This means that the antibody response will only occur when the antigen is presented on the VLP.

Here, we describe a novel plant virus derived VLP based on the cucumber mosaic virus, which incorporates a tetanus toxoid epitope for T cell help (herein referred to as CuMVtt).

To test the therapeutic efficacy of NGF vaccination, mice were immunised with either CuMVttNGF (Vax) or CuMVttctrl (Mock) 2 weeks prior to joint destabilisation (figure 1B). Non-operated sentinel control mice also underwent vaccination to enable regular blood sampling over the experimental course. Immunisation led to seroconversion by week 3, followed by a decline in antibody titres. Additional boosts were necessary to maintain antibody levels (figure 1C). No difference in pain behaviour was detected in NGF immunised animals 24 hours postoperatively (postop), but CuMVttNGF vaccinated animals recovered from pain behaviour faster than mock-vaccinated animals (within 48 hours) (figure 1D). As expected mice were pain free for several weeks, but pain behaviour started to develop from 8 weeks postsurgery. Following a boost at 10 weeks postop, and in keeping with a concomitant rise in the serum levels of anti-NGF antibody, a reversal of pain behaviour was observed. This was maintained for 3 weeks until anti-NGF titres fell and pain behaviour resumed. At termination of the experiment, joints were harvested and scored for OA severity. No difference in disease severity between mock and vaccinated groups was observed (figure 1E,F).

A second experiment was carried out to establish whether analgesia could be induced by immunisation after induction of pain behaviour i.e. therapeutic vaccination (figure 2A). When pain behaviour was established (10 weeks postop) mice were randomised into two groups: vaccinated and mock-vaccinated. Vaccine boosts were delivered at weeks 12 and 15 postop to maintain titres. Higher titre anti-NGF levels at the end of the experiment (around OD50 103) appeared to be associated with an analgesic response between weeks 14 and 18 postop (figure 2B,C). A subsequent spontaneous reduction in titres was associated with resumption of pain behaviour. Direct correlation between antibody levels and pain behaviour during the experiment was not possible as titres were only measured in the sentinel group. A meta-analysis comparing the analgesic effects across both studies at the point of highest titre in the sentinel group (week 13 for the prophylactic study and week 17 for the therapeutic study, marked by †) yielded a significant difference (p=8.93e-05) between mock and vaccinated cohorts (figure 2D).

Figure 1  Prophylactic NGF vaccination blocks murine OA pain. (A) VLP is chemically cross-linked (SMPH) to enable conjugation with His-NGF. (B) Prophylactic vaccination protocol. (C) Anti-NGF titres in sentinel cohort (n=10). (D) Painful behaviour following surgical induction of OA (n=40) measured by Linton incapacitance where 100% represents equal weight distributed across R and L limbs. Repeated-measures two-way ANOVA with Bonferroni multiple comparisons test applied, * adjusted p<0.05. SEM shown. Differences between treatment groups during late OA pain phase were not significant after correcting for multiple testing. † identifies time point of highest anti-NGF titre (see figure 2D). (E) Representative histological sections for (F) cartilage degradation (OARSI) scores 18 weeks after PMX surgery in mice treated with mock or NGF vaccine. Statistical significance is shown by two-tailed t-test. Bars represent mean±SEM, n.s.—non-significant. **p<0.01 by t-test. CuMVtt adapted from EMD: 3855. ANOVA, analysis of variance; NGF, nerve growth factor; PMX, partial meniscectomy; VLP, virus-like particle.
Recent studies using refined vaccine platforms have demonstrated requiring the use of codelivery of adjuvant in preclinical models. have been due to poor immunogenicity of the vaccine platform, preclinical success but with limited clinical efficacy, which may after 5 ug/mL. The effect appeared to plateau to that seen with 150 ug/mL monoclonal anti-NGF antibody neurite outgrowth in PC-12 cells (figure
control mice was able to dose-dependently inhibit NGF induced IgG purified from the serum of CuMVtt NGF vaccinated, but not
maintained to follow the fall in antibody titres over the following 10
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Contributors AE-T, ISV and TLV designed the studies. MFB originated the concept of the vaccine. JM-Z and ISV conducted the mouse surgery and ISV conducted the behavioural studies. AZ provided VLP constructs and developed purification strategies. AE-T produced and characterised the CuMVttNGF vaccine. AET performed the vaccinations and immunological assays. IP performed the histological preparation and in conjunction with ISV conducted the histological analysis. LJ conducted and approved of the statistical analysis. JM-A conducted the ELISA experiment.

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Competing interests MFB is founder of SAIBA GmbH and Hypopet AG, that own the VLP-platform IP and is involved in the development of therapeutic VLP-based vaccines for commerical purposes.

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Data sharing statement All data supporting this study are available on reasonable request from the corresponding authors.

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