A Neanderthal Sodium Channel Increases Pain Sensitivity in Present-Day Humans

Highlights

- Some present-day humans carry a Neanderthal variant of the sodium channel Nav1.7
- The Neanderthal variant carries three amino acid differences to the common variant
- The Neanderthal channel is more likely to be in a ready-to-open state
- Carriers of the Neanderthal variant experience more pain

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In Brief

Zeberg et al. show that a Neanderthal variant of the sodium channel Nav1.7, which is crucial for the initiation of the pain signals, occurs in some humans today. The Neanderthal variant differs from the common variant in its electrophysiological properties and is associated with increased sensitivity to pain.
A Neanderthal Sodium Channel Increases Pain Sensitivity in Present-Day Humans

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SUMMARY

The sodium channel Nav1.7 is crucial for impulse generation and conduction in peripheral pain pathways [1]. In Neanderthals, the Nav1.7 protein carried three amino acid substitutions (M932L, V991L, and D1908G) relative to modern humans. We expressed Nav1.7 proteins carrying all combinations of these substitutions and studied their electrophysiological effects. Whereas the single amino acid substitutions do not affect the function of the ion channel, the full Neanderthal variant carrying all three substitutions, as well as the combination of V991L with D1908G, shows reduced inactivation, suggesting that peripheral nerves were more sensitive to painful stimuli in Neanderthals than in modern humans. We show that, due to gene flow from Neanderthals, the three Neanderthal substitutions are found in ~0.4% of present-day Britons, where they are associated with heightened pain sensitivity.

RESULTS AND DISCUSSION

Neanderthals and their Asian relatives, Denisovans, evolved largely separately from the ancestors of present-day humans for about 500,000 years [2]. During that time, each group independently accumulated genetic changes that became frequent or fixed. However, late in their history, Neanderthals and Denisovans mixed with modern humans, which resulted in many genetic variants from Neanderthals and Denisovans being present in humans today [3, 4]. As several Neanderthal genomes of high quality are now available [2, 5, 6], it is possible to identify genetic changes that occurred in many or most Neanderthals, investigate their physiological effects, and assess their consequences when they occur in people today.

Whereas most genetic differences between Neanderthals and modern humans that affect gene products occur singly in genes across the genome, genes that carry several such differences are of particular note. One such case is the gene SCN9A, which encodes the Nav1.7 protein, a voltage-gated sodium channel in which all Neanderthal genomes sequenced to date carry three amino acid substitutions relative to modern humans: M932L; V991L; and D1908G. At these positions, extant monkeys and apes share the modern human residues. Nav1.7 forms complexes with the β3 subunit [11–13], encoded by SCN3B, which carries no amino acid differences between Neanderthals, Denisovans, and modern humans. When the β3 subunit is expressed together with Nav1.7 in the oocytes, the inactivation curve of the Neanderthal ion channel is shifted by ~6.1 mV toward less negative values relative to the modern human channel (Figures 1A and 1B; Table S1; p = 2.9 x 10^-5; n = 26 and n = 31, respectively). This results in an increased availability of sodium channels for activation and increased probability that the sodium channel remains open for a longer time once activated and is expected to lower the threshold for the generation of an action potential (see computational model; Figures S1M and S1N).

To examine which of the amino acid substitutions mediates the shift in the inactivation curve, we synthesized and injected mRNAs encoding each of the three amino acid substitutions...
singly (Figure S2). No single amino acid substitution had any effect on the inactivation of the ion channel (Figure 1E; Table S1). We next investigated the effect of the three possible combinations of two amino acid substitutions. Two of the combinations, M932L+D1908G and M932L+V991L, had no effect on the inactivation. However, V991L+D1908G caused a shift in the inactivation curve, similar to that of the three substitutions (Figure 1F; Table S1; $p = 2.4 \times 10^{-5}$; $n = 24$). Thus, an epistatic interaction of the two substitutions V991L and D1908G, which are both located in the intracellular part of the protein, is necessary to elicit the electrophysiological effects. In contrast, M932L, which is located extracellularly, had no detectable effects. We note that the Denisovan genome carries the D1908G substitution in a homozygous form but lacks the other two substitutions. This suggests that the D1908G substitution occurred in the common ancestor of Neanderthals and Denisovans but that it may have had a functional effect only in Neanderthals, where also the V991L substitution occurred.

To test whether the effects of the Neanderthal substitutions are also seen in mammalian cells, we transfected human embryonic kidney (HEK) cells with vectors expressing the mRNAs encoding the modern human and Neanderthal variants of Nav1.7 and the $\beta$3 subunit. As in the oocytes, we observe a depolarizing shift in inactivation for the Neanderthal variant of Nav1.7 (Figure 1C; Table S1; $p = 9.1 \times 10^{-3}$; $n = 8$ and $n = 9$, respectively). In addition, we observe a depolarizing shift of the activation curve (Figure S1C; Table S1; $p = 1.1 \times 10^{-3}$; $n = 8$ and $n = 9$, respectively). When investigated in a simple model of a human peripheral nerve, the depolarizing shift in the inactivation curve has an excitatory effect for suprathreshold stimuli, even in the presence of a depolarizing shift in activation (Figure S1N). Thus, we conclude that the shifts seen in both oocytes and HEK cells are likely to have an excitatory action.

Figure 1. Effects of Neanderthal-Derived Amino Acid Substitutions on Nav1.7 Inactivation
(A) Current traces from *Xenopus laevis* oocytes expressing modern human and Neanderthal (M932L+V991L+D1908G) Nav1.7 proteins together with the auxiliary subunit Nav$\beta$3. Following prepulses to three different potentials (shown as examples), currents were elicited by stepping to 0 mV ($t = 0$). Less negative prepulse potentials resulted in progressively smaller currents due to channel inactivation. This inactivation is more pronounced for the modern human than the Neanderthal protein.

(B) Fraction of current as a function of prepulse potential for the modern human and Neanderthal channel proteins. Current amplitudes, measured at $t = 10$ ms, were normalized to the largest amplitude obtained in each cell. The voltage required for half-maximal inactivation is shifted by $+6.1$ mV for the Neanderthal protein.

(C) Fraction of current as a function of prepulse potential for the modern human and Neanderthal channel proteins when expressed in human embryonic kidney (HEK) cells. Current amplitudes, measured as peak inward current, were normalized to the largest amplitude obtained in each cell. The voltage required for half-maximal inactivation is shifted by $+11.7$ mV for the Neanderthal protein.

(D) Current traces from HEK cells expressing modern human and Neanderthal (M932L+V991L+D1908G) Nav1.7 proteins together with the auxiliary subunit Nav$\beta$3

(E) Fraction of current as a function of prepulse potential in *Xenopus* oocytes for the three single amino acid substitutions individually.

(F) Fraction of current in *Xenopus* oocytes as a function of prepulse potential for Nav1.7 proteins with two substitutions. The voltage required for half-maximal inactivation is shifted by $+6.2$ mV for the V991L+D1908G protein. Compared to mammalian cells, the steady-state inactivation curves are shifted to the right due to the recording temperature (10°C ± 2°C), the co-expression of the $\beta$3 subunit, and the use of *Xenopus* oocytes, as described [14–16]. Error bars represent SEM. See also Figures S1 and S2 and Table S1.
We next investigated whether any of the three SCN9A missense mutations (M932L, V991L, and D1908G) exist in present-day humans by examining 2,535 genomes available in the phase III 1000 Genomes (1000G) dataset (Figure 2A). We did not find any of the three variants in the African (N = 507) and European (N = 505) 1000G populations, but M932L and V991L occur in almost perfect linkage disequilibrium ($R^2 = 0.99$) at frequencies of 0.9%–7.8% in Asia and 0.5%–23.8% in the Americas. In addition, D1908G occurs at a frequency of 0%–17.1% in Asia and 0.5%–52.9% in the Americas (Table S2) and tends to be in linkage disequilibrium with M932L+V991L ($R^2 = 0.26$).

These variants could have entered the present-day human population from a common ancestor shared with Neanderthals and Denisovans or by introgression from archaic hominins when these met modern humans some 40,000–60,000 years ago. In the latter case, the size of the archaic-like DNA segments on which the variants sit are expected to be substantially larger than if they were inherited from a common ancestor of the three groups, as meiotic recombination will have had less time to reduce their size. To estimate the size of these segments, we identified alleles in this genomic region that are absent in the 1000G Yoruba individuals but occur in homozygous form in the Neanderthal and/or Denisovan genomes when these met modern humans some 40,000–60,000 years ago. We next typed the two DNA segments and their recombination rates, we concluded that these variants entered the modern human population by introgression from Neanderthals or Denisovans, rather than being inherited from a common ancestor of the groups ($p = 0.059$ for V991L and M932L; $p = 7.8 \times 10^{-16}$ for D1908G). We estimated the phylogenetic relationships of the two DNA segments overlapping M932L+V991L and D1908G to the corresponding present-day human and Neanderthal DNA segments. For both segments, one group of present-day human DNA sequences is most closely related to the Neanderthal segments (Figures 2B and 2C), and in both instances, these DNA sequences are found in Asia and the Americas.

The amino acid substitutions M932L and V991L have previously been associated with increased pain sensitivity and small-fiber neuropathy in experimental and clinical studies in humans [1, 17]. To investigate whether the three introgressed Neanderthal variants affect pain sensitivity in the general human population, we selected 362,944 unrelated individuals of British ancestry in the UK Biobank (UKBB) [18] who had answered questions relating to their experience of pain (Table S3). Although no individual was homozygous for the Neanderthal variants, 1,327 individuals (0.4%) carried all three amino acid substitutions in a heterozygous form. Based on their answers to 19 pain-related questions, these carriers of the three Neanderthal substitutions had experienced one or more forms of pain more often than non-carriers (Figure 3; $p = 0.0078$; adjusted for age and sex). They also experienced more pain than carriers of only one or two of the Neanderthal substitutions, although these did not differ significantly from individuals carrying none of the Neanderthal substitutions (Figure 3). The amino acid substitutions are associated with an increased probability to report pain ($p = 0.82$ and $p = 0.72$, respectively), suggesting that their effects are smaller than the Neanderthal–derived variants.

Notably, sex did not affect the extent to which individuals in the UKBB report pain ($p = 0.84$; 198,047 females; 164,897 males),...
whereas age showed an approximately linear and positive correlation with reported pain between the ages of 40 and 70 (Figure S3; p = 4.6 × 10−282). Individuals who carried the three amino acid substitutions inherited from Neanderthals in heterozygous form are 7% more likely to report at least one form of pain compared to people without these substitutions. This corresponds to an increase of approximately 8.5 years in terms of pain reported. At the present time, we can only speculate about the consequences of having these substitutions in homozygous form, as was the case in the three Neanderthals genomes have been sequenced to high coverage.

We note that Nav1.7 is expressed also in other cell types, for example, olfactory neurons, raising the possibility that the substitutions studied here might have additional effects beyond modulation of pain. However, given the electrophysiological effects of the substitutions, nociception, i.e., the input to the central nervous system from peripheral nerves in response to harmful or potentially harmful stimuli, is likely to have been higher in Neanderthals than in modern humans. The translation of such input into the conscious perception of pain is modulated both at the level of the spinal cord and the brain. Thus, it is not possible to conclude that Neanderthals necessarily experienced more pain than modern humans do. Yet the input from Neanderthal peripheral nerve endings would have allowed that Neanderthals to be more sensitive to stimuli, as suggested by the observations in present-day people heterozygous for the Neanderthal Nav1.7 variant.

STAR+METHODS

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Figure 3. Pain Sensitivity among UK Biobank Individuals (n = 362,944)

Odds ratios (ORs) for reporting pain among individuals carrying all three amino acid substitutions (M932L+V991L+D1908G) and for each amino acid substitution singly or in a combination with one other substitution (but not all three) relative to the rest of the UKBB individuals. No individual carried V991L+D1908G. The low frequencies of the three individual variants of M932L: 0.21%, V991L: 0.23%, and D1908G: 0.29% are consistent with observations in the 1000G (0 of 100 in England and Scotland; Table S2). Horizontal lines indicate 95% confidence intervals. See also Figure S3 and Table S3.

|                  | n   | OR (95%CI)     |
|------------------|-----|---------------|
| M932L            | 1327| 1.16 (1.04-1.29) |
| M932L+V991L      | 149 | 0.91 (0.69-1.20) |
| M932L+D1908G     | 22  |               |
| Σ = 201          |     |               |
| V991L+M932L      | 153 | 0.93 (0.74-1.16) |
| Σ = 302          |     |               |
| D1908G+V991L     | 780 | 0.99 (0.86-1.14) |
| Σ = 802          |     |               |
| 0.6              | less likely to report pain | more likely to report pain |
| 0.8              |     |               |
| 1                |     |               |
| 1.2              |     |               |
| 1.4              |     |               |

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AUTHOR CONTRIBUTIONS

H.Z., M.D., J.K., and S.P. designed the study; H.Z. performed the electrophysiological experiments with assistance from K.S., H.P.C.R., and W.H. M.D., K.T., J.K., and S.P. did phylogenetic analyses; M.D. and H.Z. studied the phenotypic effects in UKBB; V.W. performed the in vitro translation; and H.Z., M.D., and S.P. wrote the manuscript with input from all authors.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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STAR METHODS

KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Chemicals, Peptides, and Recombinant Proteins | | |
| Normal Frog Ringer | EcoCyte Bioscience | Cat#: 1688 |
| Modified Barths Solution | EcoCyte Bioscience | Cat#: 1697 |
| Penicillin-Streptomycin | Sigma-Aldrich | Cat#: P4333 |
| Sodium pyruvate | Sigma-Aldrich | Cat#: P2256 |
| Dulbecco’s modified Eagle medium | Sigma-Aldrich | Cat#: D5796 |
| TrypLE Express Enzyme | Thermo-Fisher | Cat#: 12605010 |
| Fetal Bovine Serum | Sigma-Aldrich | Cat#: F7524 |
| Critical Commercial Assays | | |
| mMESSAGE mMACHINE T7 ULTRA Transcription Kit | Thermo-Fisher | Cat#: AM1345 |
| Deposited Data | | |
| Archaic genomes | Max-Planck Institute EVA | http://cdna.eva.mpg.de/neandertal |
| 1000 genome project | The International Genome Sample Resource | http://www.internationalgenome.org/data |
| UK BioBank | UK Biobank Limited | https://www.ukbiobank.ac.uk/ |
| GTEx gene expression | The Genotype-Tissue Expression (GTEX) project | https://gtexportal.org/home/datasets |
| Experimental Models: Organisms/Strains | | |
| Xenopus oocytes | EcoCyte Bioscience | First grade |
| Human embryonic kidney cells | University of Cambridge | HEK293T |
| Recombinant DNA | | |
| SCN9A (Nav1.7) cDNA | GenScript | Cat#: OHu22884 |
| SCN3B (Nav3) cDNA | GenScript | Cat#: OHu16302 |
| pXOOM vector | Søren Olesen [19] | N/A |
| Software and Algorithms | | |
| MATLAB | MathWorks | R2018a |
| R | R Foundation | ver. 3.5.3 |

RESOURCE AVAILABILITY

Lead Contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Hugo Zeberg (hugo.zeberg@ki.se).

Materials Availability
The plasmids encoding the Nav1.7 variants are available from the Lead Contact upon request.

Data and Code Availability
The recording software, utilizing the MATLAB Data Acquisition Toolbox, and electrophysiological data, including steady-state inactivation and activation as a function of voltage, are available at https://gin.g-node.org/hugozeberg/SCN9A. Code used for this study and raw data are available upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

*Xenopus laevis* (EcoCyte Bioscience, Castrop-Rauxel, Germany) oocytes were kept at 12°C in Modified Barth Solution, supplemented with 2.5 mM sodium pyruvate and 25 U/mL penicillin and 25 μg/mL streptomycin. HEK293T cells were cultured in Dulbecco’s Modified Eagle Medium (DMEM), supplemented with 10% fetal calf serum and 100 U/mL penicillin and 0.1 mg/mL streptomycin, at
37°C in a 5% carbon dioxide humidified incubator. For the human phenotypic analysis, 198,047 adult females and 164,897 adult males from United Kingdom were included in the study (the UK Biobank).

**METHOD DETAILS**

**Molecular biology**
SCN9A (Nav1.7) and SCN3B (Navβ3) cDNAs were synthesized by GenScript (Piscataway, NJ). The SCN9A construct used is the most abundant splice isoform in dorsal root ganglia (DRG), and corresponds to the RefSeq sequence NM_002977.3, which encodes a W at position 1150, as in the majority of modern humans, as well as in all three high-coverage Neanderthal genomes. In contrast, the human reference carries an R at this position seen in 13% of present-day humans (rs6746030). pXOOM [19] was used as the plasmid backbone and all constructs were verified by sequencing. Plasmids were linearized with the restriction enzyme Xhol and transcribed in vitro using the T7 mMessage mMACHINE ULTRA kit (Ambion, Austin, TX). cRNA concentration and purity were determined using a spectrophotometer.

**Oocyte injection**
Stage V-VI oocytes injected using a Nanoinject II Auto-Nanoliter Injector (Drummond Scientific, Broomall, PA) with 2 ng SCN9A cRNA and (in experiments with the Navβ3 subunit) 20 ng SCN3B cRNA in a volume of 50 nL RNase-free water. SCN3B was injected in excess relative to SCN9A, in order to assure that Nav1.7 channels were complexed with Navβ3 subunits. After injection, oocytes were incubated at 12°C for 4-6 days, before electrophysiological recordings.

**Transfection of HEK293T cells**
HEK293T cells were transfected with 2 μg of each of the two cDNA vectors using 10 μg branched polyethylenimine (~25kDa) in 2 mL of DMEM in 9.5 cm² wells for 4 hours. Cells were harvested using TrypLE Express Enzyme, seeded in 3.5 cm dishes, and used for voltage clamp experiments 24-48 hours post transfection.

**Voltage clamp recordings in Xenopus oocytes**
Experiments were performed at 10 ± 2°C using a two-electrode voltage-clamp setup and either a CA-1 amplifier (Dagan, Minneapolis, MN), or an OC-725C amplifier (Warner Instruments, Hamden, CT). Data were acquired at 100 kHz via a NI-6221 DAQ using custom written software in MATLAB (Mathworks, Natick, MA). External solution consisted of (in mM); 90 NaCl, 2 KCl, 1 MgCl₂, 2 CaCl₂, and 5 HEPES, adjusted to pH 7.2. Micropipettes were pulled from borosilicate glass capillaries (Harvard Apparatus, Holliston, MA) and had a resistance of 0.5-1 MOhm when filled with 3M KCl. The bath potential was actively controlled to minimize series resistance in the bath and across the bath-grounding electrode. Oocytes had an input resistance of 1.6 ± 0.9 MOhm (n = 216).

The voltage clamp protocols are shown in Figure S1A. The first 300 ms pulse from the baseline of ~90 mV was used to measure channel activation and the time constant for inactivation. Steady-state inactivation was measured by a second 0 mV pulse immediately following the first pulse, given the temperature coefficient of the fast inactivation (Q10 ~3 [20]), a 300-ms prepulse is equivalent to 100-ms prepulse at room temperature. Membrane potential was clamped at ~90 mV for 5 s between consecutive trials. Current traces (Figure S2) were filtered after recordings using a rolling average of 100 data points. In Figure 1A, capacitive transients (time constant of ~1 ms) have been eliminated by fitting and subtracting a single exponential. For data analysis, we measured the current amplitude at t = 10 ms (into the 0-mV step), a time point at which virtually all capacitive transients had declined (~0.004% of the initial stimulation, or ~2 nA, remained). For inactivation measurements, the current following a prepulse to +10 mV was taken as baseline (I_base, compensating for any non-inactivating currents, including leak currents) and current (I) was normalized to the maximal observed current in that cell (I_max) according to (I-I_base)/(I_max-I_base). The channels were considered to have fully inactivated following the +10 mV prepulse, based on previous work in Xenopus oocytes [14].

Activation as a function of voltage was measured as peak inward current relative to the current observed at the end of the 300 ms into the first pulse (i.e., immediately before the second pulse to 0 mV). Currents were converted to conductance using g = I/V-E_{rev,Na}. The reversal potential E_{rev,Na} was calculated using the Nernst equation, assuming an intracellular sodium concentration of 22.5 mM [21], yielding a reversal potential of +33.8 mV.

Data were fitted to a first (inactivation) or second-order Boltzmann equation (activation; as suggested by the Frankenhaeuser-Huxley model [22]). The curves were fitted over the dynamic range of the functions (~40 to 0 mV for activation and ~70 to ~10 mV for inactivation). For the activation function, a fixed slope of 5.9/mV provided the overall best fit (see Figures S1D–S1G) and for the steady-state inactivation function, 15.1/mV (Figures S1I–S1L). The numerical values of the fitted midpoint potentials are given in Table S1.

The time-constants for inactivation were measured by fitting a single exponential between 10 and 20 ms into the 300-ms pulse, in traces showing a current decay > 10 nA from 10 to 20 ms. Outliers were detected using the built in outlier detection in MATLAB. Whereas the data points are experimental measurements, the bell-shaped curves in Figures S1O–S1Q are derived using a kinetic model [23] with values from the steady-state inactivation curves.
Voltage clamp experiments in HEK293T cells

HEK293T cells were analyzed at room temperature (~23 °C) using either an Axon Multiclamp 700A or a 700B patch-clamp amplifier, Molecular Devices, San Jose, CA. The pXOOM vector encodes an enhanced green fluorescent protein, which was excited to identify transfected cells. The external solution consisted of (in mM): 140 NaCl, 2.5 KCl, 1 MgCl₂, 2 CaCl₂, 10 glucose, and 10 HEPES, adjusted to pH 7.4 with NaOH. Patch pipette were pulled from borosilicate glass capillaries (Harvard Apparatus, Holliston, MA), and the tip of the electrode was fire-polished. For low noise voltage-clamp recordings (Figure S2) with potassium currents blocked, pipettes were filled with a solution, containing (mM): Cs methane sulfonate 125, CsCl 5, KCl 10, EGTA 1, 10 HEPES, 4 Mg-ATP, 0.3 GTP-Na₂, 10 creatine phosphate-Na, pH adjusted to 7.3 with CsOH. Electrodes had a resistance of 4-11 MΩ, and the membrane potential signal was corrected for nulling of the liquid junction potential before seal formation. Data was filtered using a low-pass filter at 6 kHz (Bessel, 8-pole) and sampled at 20 kHz via a NI-X-series DAQ using custom software written in MATLAB (Mathworks, Natick, MA). Recording pipettes were positioned with LM-Mini stepper-motor-controlled micromanipulators (Luigs and Neumann, Ratingen, Germany). Cells had an input resistance of 1.5 ± 1.6 GΩ (min = 112 Mohm, max = 4.5GΩm, n = 17). A P/n protocol consisting of 4 negative steps in every sweep, each 5 mV and 500 ms long, was used to compensate for passive membrane properties. Capacitance was also compensated for by the built-in capacitance correction circuit of the amplifier. In order to adjust for the effect of nonlinearity in the background current, the P/n correction was scaled to between 30% and 100% to minimize the amplitude of the residual capacitance transient. The leak conductance was estimated using the current response between −120 to −70 mV, before any activation of the Na current.

Activation and inactivation were investigated with the voltage clamp protocol shown in Figure S1A. Amplitude was measured as the peak inward current. Given that inactivation is hyperpolarized in mammalian cells, the voltage protocol spanned the range −120 mV to +10 mV. Activation of conductance as a function of voltage was computed using the sodium equilibrium potential of +65mV. Data were fitted over the entire range to a first (inactivation) or second-order Boltzmann equation (Figures S1C and S1H). The slopes from the oocyte experiments provided a good fit for HEK cells and were kept as fixed constants. The numerical values of the fitted midpoint potentials are given in Table S1.

Computational modeling

The effect of a 10-mV shift of activation and fast inactivation was studied using a model of a peripheral nerve, based on voltage clamp data from the human node of Ranvier [24] (ModelDB; accession: 261436). The model was implemented in MATLAB.

Amino-acid substitutions in Neanderthal ion channels

Fifteen voltage gated-like ion channels (CACNA1D, CACNA1H, CACNA1S, CATSPER3, CNGB1, HCN4, KCNH3, KCNK2, KCNV2, PKD2L, SCN9A, TRPC6, TRPM1, TRPM4, and TRPV4) carry amino-acids substitutions in the three high-coverage Neanderthal genomes available. Expression data of these 15 channels in single-cell RNA sequencing data from rodent DRG [7] showed high expression of SCN9A only.

Archaic introgression

Potentially introgressed segments in and around SCN9A were identified by studying “archaic single-nucleotide polymorphisms” ("aSNPs") defined as those that are homozygous ancestral in all 1,000G (phase III) Yoruba individuals [25], homozygous derived in at least one of the Altai Neanderthal, the Vindija Neanderthal, and the Denisovan genomes [2, 6, 26], and for which the derived allele is present in at least one non-African 1,000G individual. In addition, aSNPs were required to overlap previously identified regions of Neanderthal introgression in present-day humans, i.e., having a Neanderthal posterior probability greater than 0.9 and a length of at least 0.02 cM [3].

The likelihood of incomplete lineage sorting was tested using a published method [27], an age of the Altai Neanderthal based on a mutation rate of 1.0x10⁻⁹ per base pair per year [28], and the local recombination rate estimate [29].

The relationships of present-day and Neanderthal DNA segments were estimated using genotype information from 1,000G (phase III) individuals, excluding sites that were missing in at least one individual or were heterozygous in one of the three archaic individuals. This resulted in 727 variable positions in the regions overlapping V991L and M932L, and 2,868 variable sites in the region overlapping D1908G. Haplotypes that had less than five differences per 10,000 base pairs to each other were clustered, resulting in six clusters per region. Consensus sequences were generated by randomly selecting a variant at each position. Based on 100 and 363 positions that were variable among these consensus sequences, maximum-likelihood phylogenies were constructed using the Hasegawa-Kishino-Yano-85 model [30] with a proportion of invariant sites and a discrete-gamma distribution to account for variable substitution rates among sites.

Phenotypic consequences of Nav1.7 variants

Excluding individuals that were related, showed unusually low levels of heterozygosity, or who had substantial non-British ancestry left 362,944 of 487,409 individuals from the UK Biobank [18], which were used for subsequent analysis. All combinations of the three variants were tested for correlations with the 19 pain phenotypes and additionally with a composite binary measure that reflects whether an individual was positive for at least one of the 19 pain-related phenotypes (Table S3). Correlations between genotypes and pain phenotypes were assessed using logistic regression, including sex and age as co-factors in the linear model. Whereas the primary analysis involving the composite pain measure for the Neanderthal Nav1.7 variant rejected the null hypothesis, none
of the other secondary analyses resulted in a p value < 0.05, and hence these p values were not adjusted for multiple comparisons. The potential impact of noncoding aSNPs was identified using the Variant Effect Predictor [31, 32]. Identified noncoding variants were then checked for expression changes in 48 GTEx tissues [33].

**QUANTIFICATION AND STATISTICAL ANALYSIS**

Oocytes were randomly selected for injection of different mRNA transcripts. Each oocyte was used only once (n = 216). HEK cells were selected based on expression of GFP (n = 17). For the electrophysiology recordings, data were collected until the SEM of the midpoint potentials of the steady-state activation and inactivation curves were less than ± 1.5mV. Recordings from oocytes that had a holding current greater than 5 μA at −90mV were excluded. HEK cells were excluded if they had a leak current larger than 500 pA at −120mV, did not display a Na current (an inward current at −20mV with a peak around 1-2 ms), displayed other inward ion current fractions, or showed an atypical current-voltage relationship [34, 35] due to suboptimal recording conditions (sum of square errors > 0.05 of the fit of a standard first-order Boltzmann curve to the inactivation process). The significance of any difference in the midpoint potentials was determined using a two-tailed t test with n-2 degrees of freedom, corrected for multiple comparisons using Bonferroni correction. The normality of midpoint potentials was verified using the Shapiro-Wilk test.

**ADDITIONAL RESOURCES**

The archaic genomes are available at http://cdna.eva.mpg.de/neandertal/. The genomes from the 1000 Genomes Project are available at http://www.internationalgenome.org/data. The UK BioBank is available at https://www.ukbiobank.ac.uk/. GTEx gene expression data is available at https://gtexportal.org/home/datasets.