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Key Points:

- The strength of the geomagnetic field is a proxy for the flux of ultraviolet radiation (UVR)
- The disappearances of the Neanderthals and many large mammals during the Late Quaternary occurred during minima in geomagnetic field strength
- Human phylogeny from mitochondrial DNA and Y chromosomes can also be linked to minima in field strength, hence UVR flux

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The Role of Geomagnetic Field Intensity in Late Quaternary Evolution of Humans and Large Mammals

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Abstract It has long been speculated that biological evolution was influenced by ultraviolet radiation (UVR) reaching the Earth's surface, despite imprecise knowledge of the timing of both UVR flux and evolutionary events. The past strength of Earth's dipole field provides a proxy for UVR flux because of its role in maintaining stratospheric ozone. The timing of Quaternary evolutionary events has become better constrained by fossil finds, improved radiometric dating, use of dung fungi as proxies for herbivore populations, and improved ages for nodes in human phylogeny from human mitochondrial DNA and Y chromosomes. The demise of Neanderthals at ~41 ka can now be closely tied to the intensity minimum associated with the Laschamp magnetic excursion, and the survival of anatomically modern humans can be attributed to differences in the aryl hydrocarbon receptor that has a key role in the evolutionary response to UVR flux. Fossil occurrences and dung-fungal proxies in Australia indicate that episodes of Late Quaternary extinction of mammalian megafauna occurred close to the Laschamp and Blake magnetic excursions. Fossil and dung fungal evidence for the age of the Late Quaternary extinction in North America (and Europe) coincide with a prominent decline in geomagnetic field intensity at ~13 ka. Over the last ~200 kyr, phylogeny based on mitochondrial DNA and Y chromosomes in modern humans yields nodes and bifurcations in evolution corresponding to geomagnetic intensity minima, which supports the proposition that UVR reaching Earth's surface influenced mammalian evolution with the loci of extinction controlled by the geometry of stratospheric ozone depletion.

Plain Language Summary The strength of Earth's magnetic field in the past, recorded by rocks and sediments, provides a proxy for past flux of ultraviolet radiation (UVR) to Earth's surface due to the role of the field in modulating stratigraphic ozone. About 40,000 years ago, mammalian fossils in Australia and Eurasia record an important die-off of large mammals that included Neanderthals in Europe. In the Americas and Europe, a large mammalian die-off appears to have occurred ~13,000 years ago. Both die-offs can be linked to minima in Earth's magnetic field strength implying that UVR flux variations to Earth's surface influenced mammalian evolution. For the last ~200,000 years, estimates of the timing of branching episodes in the human evolutionary tree, from modern and fossil DNA and Y chromosomes, can be linked to minima in field strength, which implies a long-term role for UVR in human evolution. New fossil finds, improved fossil dating, knowledge of the past strength of Earth's magnetic field, and refinements in the human evolutionary tree, are sharpening the focus on a possible link between UVR arriving at the Earth's surface, magnetic field strength, and events in mammalian evolution.

1. Introduction

The apparent spacing of mass extinction events at long (~26 Ma) repeat times (e.g., Raup & Sepkoski, 1986), and the supposed role of geomagnetic polarity reversal in extinction (e.g., Raup, 1985), has received intermittent attention over the last 50 years, since early studies of Quaternary radiolarian evolution and polarity reversal in deep-sea sediments (Hays, 1971). The reader is referred to Aubry et al. (2009) for abbreviations denoting geological time in the past (Ma for millions of years ago and ka for thousands of years ago) and equivalent durations (Myr and kyr, respectively). Linkage between polarity reversal (with its concomitant low field intensity) and extinction or speciation has not gained significant traction, perhaps because of uncertainties in the polarity timescale itself, and in the chronology of extinction/speciation outside of the few well-documented mass extinctions. On the other hand, we now know from Quaternary studies that although polarity reversals coincided with relative paleointensity (RPI) minima, intervals

between polarity reversals are also characterized by numerous RPI minima, some of which coincide with magnetic excursions (see Laj & Channell, 2015). The chronology of both the RPI record and the paleontological record remains poorly constrained, even for the Quaternary, such that a linkage between extinction and RPI minima cannot be ruled out.

The geomagnetic field helps to preserve stratospheric ozone, as well as atmospheric composition, density, and oxygen levels that are vital to Earth's biosphere (Wei et al., 2014). The field shields Earth from galactic cosmic rays (GCRs) and solar wind, and from harmful ultraviolet radiation (UVR) that affect the function of living systems (Belisheva et al., 2012; Mendoza & de La Peña, 2010). The demise of the Martian magnetic field, several billion years ago, is widely believed to have been the root cause for the near disappearance of the Martian atmosphere and the resulting dramatic change in the Martian environment from one featuring surface water and aqueous sedimentation to its present relative inactivity and sterility. The explosion of life in the Early Cambrian period at ~530 Ma has been associated with growth of Earth's inner core, the supposed strengthening of the dipole geomagnetic field, and the resulting thickening of Earth's atmosphere (Doglioni et al., 2016), although there is little evidence for strengthening of the geomagnetic field at this time (e.g., Biggin et al., 2015). On the other hand, the Late Ediacaran and Early Cambrian periods (~550 and ~530 Ma, respectively) may have been times of unusually high polarity reversal frequency (Bazhenov et al., 2016; Pavlov & Gallet, 2001), although precise estimates of reversal frequency are elusive due to poorly constrained age control in stratigraphic sections where the reversals were recorded. Meert et al. (2016) proposed that high reversal frequency (up to ~20 reversals/Myr) at this time would have been associated with low geomagnetic field intensity that therefore lowered shielding from UVR, which created an evolutionary advantage for burrowing and shelled organisms. These proposals for the role of the geomagnetic field in evolution are controversial partly because of poor knowledge of the state of the geomagnetic field 500-550 million years ago. Oxygenation of the oceans and atmosphere after the Gaskiers glaciation at ~580 Ma (Canfield et al., 2007) may have been the principal driver of the Early Cambrian explosion of life, both through oxygen levels at Earth's surface and increased UVR shielding through enhanced stratospheric ozone concentrations.

Several strategies in modern organisms reflect the evolutionary impact of UVR. Behavioral adaptations to UVR include vertical water-column migration in aquatic organisms, the presence of UVR-screening pigmentation decreasing with water depth, and complete disappearance of pigments for deep-water and cave-dwelling animals (e.g., Hessen, 2008). The red coloration of alpine plankton and the "red sweat" of the hippopotamus (Saikawa et al., 2014) are examples of evolutionary adaptation to high UVR at altitude and at low latitudes, respectively. UVR causes two classes of DNA lesions: cyclobutane pyrimidine dimers and 6-4 photoproducts. Both lesions distort DNA structure, introducing bends or kinks and thereby impeding transcription and replication (e.g., Branze & Foiani, 2008; Clancy, 2008). Relatively flexible areas of the DNA double helix are most susceptible to damage. One "hot spot" for UV-induced damage is found within a commonly mutated oncogene TP53 (Benjamin & Ananthaswamy, 2006), which in normal function has an important role in tumor suppression. At low concentrations, reactive oxygen species (ROS) play vital roles during mutagenic activity in response to pathogen attack. Higher concentrations of ROS produced by UVR give rise to oxidative stress where ROS attack DNA bases and the deoxyribosyl backbone of DNA (see MacDavid & Aebisher, 2014). Production of antioxidant enzymes neutralizes ROS, and ROS modulation is controlled by the aryl hydrocarbon receptor (AhR) that plays a key role in mammalian evolution.

The consequences of ionizing radiation associated with GCRs and solar particle events (solar wind) for human health have received attention in recent years in an effort to evaluate the health effects of future space travel outside Earth's protective magnetosphere (e.g., Delp et al., 2016). Earth's atmosphere is opaque to all but the highest energy GCRs, and, together with the geomagnetic field, serves to shield Earth's surface from GCRs. The intensity of UVR arriving at Earth's surface decreases with increasing latitude and is attenuated by stratospheric ozone (O₃) that acts as a sink for UVR. The geomagnetic field plays an important role in preserving the atmosphere, including stratospheric ozone that would otherwise be stripped away by solar wind and GCRs (Wei et al., 2014). UVR triggers dissociation of oxygen molecules (O₂) into oxygen radicals that combine to form stratospheric ozone that in turn absorbs UVR as it splits into oxygen atoms. Certain ozone-depleting agents (such as nitrogen oxides) are produced naturally by energetic particle precipitation from solar wind, particularly during solar proton events, and therefore, times of low geomagnetic field

strength lead to higher ozone depletion (Randall et al., 2005, 2007). Atmospheric modeling implies substantial increases in hydrogen and nitrogen oxide concentrations due to enhanced ionization by GCRs during the Laschamp excursion, with significant decrease in stratospheric ozone particularly at high latitudes (Suter et al., 2014). Modeling of ozone depletion during polarity reversals, based on a geomagnetic field intensity ~10% of the present value, leads to enhanced UVR flux at the Earth's surface, particularly at higher latitudes, that is 3-5 times that resulting from the anthropogenic ozone hole (Glassmeier & Vogt, 2010; Winkler et al., 2008). Prior to anthropogenic emission of ozone-depleting chlorofluorocarbons and halons, energetic particle precipitation at times of low geomagnetic field strength played an important role in ozone depletion. A well-defined nitrate peak, together with a broader ¹⁰Be peak, are associated with low field strength at the time of the Laschamp magnetic excursion (~41 ka) in the EPICA-Dome C Antarctic ice core (Traversi et al., 2016), which indicates that geomagnetic shielding played a role in the production of both cosmogenic isotopes (such as ¹⁰Be) and ozone-depleting nitrogen compounds. It is noteworthy that bacterial UVR proxies in sediments from Lake Reid (Antarctica) imply more than three times higher UVR flux during part of the last glacial than during the Holocene (Hodgson et al., 2005). UVR exposure affects the early stage of life in modern marine plankton, and plankton-benthos coupling in coastal waters (e.g., Hernandez Moresino et al., 2011). Furthermore, UVR plays a role in photosynthesis (e.g., Hollosy, 2002) and can cause changes in vegetation and habitat modification.

The aim of this paper is to review the record of geomagnetic field intensity over the last ~300 kyr, and compare this record with the fossil record of extinction in terrestrial mammals (including Neanderthals), and with nodes in hominin phylogeny determined from mitochondrial DNA (mtDNA) and Y chromosomes.

2. The Geomagnetic Field

Knowledge of the Holocene geomagnetic field (i.e., over the last ~12 kyr) has been based on models built from archaeomagnetic, lava, and sediment data (Constable et al., 2016; Korte et al., 2011; Pavón-Carrasco et al., 2014). Beyond the Holocene, geomagnetic field strength during the Quaternary, over the last ~2.5 Myr, has been acquired primarily from RPI data from marine sediments recovered by deep-sea drilling (e.g., Channell et al., 2009, 2018; Laj et al., 2000, 2004; Valet et al., 2005; Xuan et al., 2016; Ziegler et al., 2011). Absolute paleointensity data from lavas are hampered by unknown time gaps between lava flows and inadequate age control in young (Quaternary) lavas. RPI data from sediments have been acquired by normalizing the natural remanent magnetization (NRM) intensity by a laboratory-acquired magnetization designed to activate the same population of magnetic grains that carry the NRM, thereby compensating for variations in concentration of NRM-carrying grains throughout the sedimentary section. The laboratory-applied normalizers are typically anhysteretic remanent magnetization (ARM) and/or isothermal remanent magnetization (IRM). ARM is acquired in a decreasing alternating field (AF, with a peak AF of ~100 mT) with a weak direct current bias field (typically 50 μT), and IRM is acquired in a strong direct current field (up to ~1 T). The appropriate normalizer is usually chosen such that its coercivity (response to AF demagnetization) closely matches that of the NRM. A typical RPI proxy comprises the slope of NRM intensity versus ARM (and/or IRM) intensity determined over a particular peak demagnetization field range (such as 20-80 mT). The definition of the slope, often determined at 1-cm spacing down-core, can be gauged by determining its linear correlation coefficient (e.g., Xuan & Channell, 2009).

There is substantial agreement among Late Quaternary stacks of global sedimentary RPI data (Channell et al., 2009; Valet et al., 2005; Xuan et al., 2016; Ziegler et al., 2011) that are independently supported by 10 Be/ 9 Be data from marine sediments (e.g., Simon et al., 2016) due to the role of the geomagnetic field in modulating cosmogenic isotope (10 Be) production. Agreement is, however, poor among stacks of sedimentary RPI data and 10 Be/ 9 Be data in the 0-40 ka interval (see Figure 1 of Channell et al., 2018) that is attributed to subtle and often unrecognized drilling disturbance in poorly consolidated uppermost sediments recovered from the ocean floor. A recently published RPI stack for 10-40 ka based on high-deposition-rate sediments from the Iberian Margin and elsewhere (Figure 1) is supported by paleointensity estimates from revised calculation of 10 Be flux in Greenland ice cores (Channell et al., 2018) using the GICC05 Greenland ice-core age model (Svensson et al., 2008). Models and stacks covering the same 0- to 40-ka interval that use RPI data from lower sedimentation rate sequences (e.g., Panovska et al., 2018) cannot resolve the detail that is revealed by higher sedimentation rate sequences and by 10 Be flux in ice cores.

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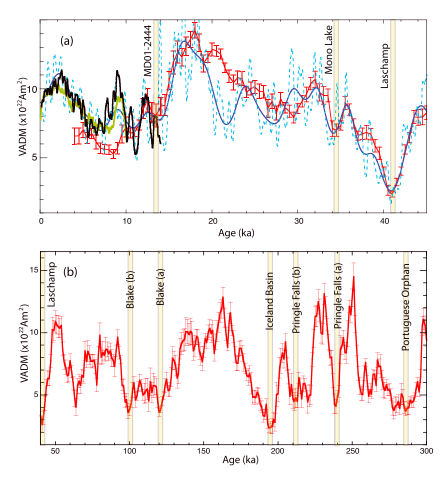


Figure 1. Relative paleointensity (RPI) calibrated to virtual axial dipole moment (VADM) for the interval 0–45 ka (a) and 40–300 ka (b). (a) Overall RPI stack of 23 sedimentary records (red) with standard error (2σ) compared with the ¹⁰Be-based VADM from Greenland ice cores (blue) filtered with cutoffs of 1/3,000 years (dark blue) and 1/500 years (dashed light blue; Channell et al., 2018). Also shown are Holocene VADM models of Constable et al. (2016; green) and Pavón-Carrasco et al. (2014; black). (b) PISO RPI stack for 40-300 ka with standard error (Channell et al., 2009). The yellow shading indicates RPI minima in the stack that have been associated with directional magnetic excursions. See text for references that document the labeled excursions.

When adequately recorded, "magnetic excursions" are manifested in both lavas and sediments as antipodal magnetization directions defining short-lived polarity reversal, and they occupy minima in RPI records (Laj & Channell, 2015). At least seven magnetic excursions have been documented in lavas and sediments of the last 300 kyr (Figure 1) with ~13 in the Brunhes Chron (last 775 kyr). Directional magnetic excursions have been named after the location where they were initially recorded such as Mono Lake (34 ka), Laschamp (41 ka), Blake (~115 ka), Iceland Basin (190 ka), and Pringle Falls (211 and/or 238 ka). Although the first record of a magnetic excursion (the Laschamp excursion) was published over 50 years ago (Bonhommet & Babkine, 1967), magnetic excursions remain controversial mainly because the aberrant magnetization directions that define them have submillennial to millennial duration and are therefore fortuitously recorded in geological archives. Although the Laschamp excursion has now been recorded in scores of globally distributed sedimentary sequences (e.g., Channell et al., 2017; Collins et al., 2012; Evans et al., 2007; Laj et al., 2000, 2006; Lund et al., 2005; Mazaud et al., 2002), the Laschamp records represent a small fraction of the total number of paleomagnetically studied sedimentary sequences that cover the Laschamp interval. The submillennial duration of the excursion means that recordings of the excursion are generally restricted to sequences with mean sedimentation rates >10-15 cm/kyr that are unusual in deep-sea sediments (see Roberts & Winklhofer, 2004). Lava records of the Laschamp are restricted to the Massif Central in France (e.g., Laj et al., 2014), the Auckland volcanic field (Cassata et al., 2008), and New Zealand's Mt. Ruapehu (Ingham et al., 2017), that just happened to have erupted with high frequency during this brief period.

The age of the Laschamp excursion, and the low geomagnetic field strength associated with it (Figure 1), is based on correlation of deep-sea cores that record the Laschamp excursion to ice-core chronologies (e.g., Channell et al., 2017; Laj et al., 2000; Nowaczyk et al., 2012), by 40 Ar/ 39 Ar age determinations in lavas from the Massif Central (e.g., Laj et al., 2014), by U/Th ages in a speleothem from Missouri (Lascu et al., 2016), and by the age of the related cosmogenic-isotope flux peak in ice cores (Baumgartner et al., 1998; Svensson et al., 2006, 2008; Traversi et al., 2016; Wagner et al., 2000; Yiou et al., 1997). Current estimates of the duration and midpoint age of the Laschamp directional excursion are <1 kyr and 41 ka, respectively. A less abrupt geomagnetic field intensity decrease (over several kyr) defines a paleointensity minimum that brackets the directional excursion.

Directional records of the Iceland Basin excursion at ~190 ka have been recovered from numerous deep-sea sediment cores from the Northern Hemisphere (see review in Channell, 2014), in cores from Lake Baikal (Oda et al., 2002), and two sites from the South Atlantic ocean (Channell et al., 2017; Stoner et al., 2003). The age of the excursion has been established by correlation to marine oxygen isotope records, and to ice core records. There are no records of the Iceland Basin excursion from well-dated lava sequences.

Apart from the Laschamp and Iceland Basin excursions, magnetic excursions during the last 300 kyr are controversial because of the paucity of records and/or poor age control in available records. On the other hand, RPI records and hence recordings of RPI minima associated with the excursions are numerous, presumably because RPI minima associated with excursions are manifest over longer timescales than the associated directional excursions.

The Mono Lake excursion at ~34 ka has been recorded in both deep-sea sediments and lava flows (Cassata et al., 2008; Channell, 2006; Kissel et al., 2011; Laj et al., 2014; Negrini et al., 2014) as well as at Mono Lake in California (e.g., Benson et al., 2003), although the age at the Mono Lake type-location is controversial (e.g., Cassata et al., 2010; Kent et al., 2002). The age of a RPI minimum associated with the Mono Lake excursion can be estimated from a cosmogenic-isotope flux maximum in Greenland ice cores (Muscheler et al., 2005; Wagner et al., 2000).

Although the older of the two Blake excursions (100 and 120 ka) was first recorded almost 60 years ago (Smith & Foster, 1969), there have only been a handful of observations of either Blake excursion since (see Laj & Channell, 2015). The older Blake excursion has, however, been observed in diverse media: marine sediments (e.g., Tric et al., 1991), Chinese loess (e.g., Zhu et al., 1994), and a speleothem from Spain (Osete et al., 2012). The younger "Blake" excursion, also known as the Skalamaelifell excursion, has been observed in Icelandic and Italian lavas (Jicha et al., 2011; Singer et al., 2014). The age of the Pringle Falls excursion (at 211 ka and/or 238 ka) is not yet settled due to a lack of records in well-dated sequences (Herrero-Bervera et al., 1994; Laj & Channell, 2015; Singer, 2014; Singer et al., 2008).

An apparent RPI minimum at ~13 ka appears in some individual RPI records, as a notch in RPI stacks (Figure 1), and in virtual axial dipole moment (VADM) proxies from 10 Be flux in Greenland ice cores (Channell et al., 2018). In one core from the Iberian Margin (MD01-2444), a pronounced RPI minimum at ~13 ka is associated with a directional magnetic excursion (Channell et al., 2013). The apparent RPI minimum at ~13 ka in core MD01-2444 coincides with the old-end of the 0-14 ka Holocene paleointensity model of Pavón-Carrasco et al. (2014). The RPI stack of Channell et al. (2018) contains a prominent field intensity peak (VADM ~13 × 10^{22} Am²) around 18-17 ka followed by a field collapse between 16 and 14 ka culminating in a minimum VADM value (~7 × 10^{22} Am²) that has been smoothed by stacking (Figure 1). The VADM decrease at 17-13 ka corresponds to a rate of decrease (12.1 nT/year) that is comparable to the rate (12.7 nT/year) observed in the 10 Be-derived field at the same time (Channell et al., 2018) and to the rates observed during the Laschamp/Mono Lake excursions (Laj & Kissel, 2015).

To provide a reference template for geomagnetic field intensity over the last ~300 kyr, we combine the RPI stack for 14-45 ka (Channell et al., 2018) with the Holocene model for 0-14 ka (Pavón-Carrasco et al., 2014) and with the PISO paleointensity stack (Channell et al., 2009) beyond 45 ka. This paleointensity template was calibrated to VADM by aligning it with Holocene archaeomagnetic models (Constable et al., 2016; Korte et al., 2011; Pavón-Carrasco et al., 2014) and assuming a value of ~1.5 \times 10²² Am² for the VADM minimum at the Laschamp excursion (Laj et al., 2014). This VADM template provides a proxy for geomagnetic intensity variations that is compared with events in mammalian evolution, in order to test potential linkages between geomagnetic field intensity variations and Late Quaternary mammalian evolution.

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3. Late Quaternary Extinctions

Causes of extinction of mammalian megafauna (adult weight >45 kg) during the Late Quaternary, the so-called Late Quaternary Extinction (LQE), have been the subject of prolonged debate (e.g., Koch & Barnosky, 2006; Martin, 1967; Stuart, 2015). Prior to ~13 ka, the mammal assemblage of the Americas included large-bodied animals such as mammoths, horses, camels, saber-tooth cats, and the short-faced bear. Extinction was total for mammals larger than 1,000 kg, >50% for size classes between 32 and 1,000 kg, and ~20% for those between 10 and 32 kg (Koch & Barnosky, 2006). Within a short time window, >150 species were lost in the Americas, including all mammals over ~600 kg. An analogous catastrophic size-controlled LQE affected 14 of 16 Australian mammalian genera; however, extinction ages are at least ~30 kyr older than in North America. Although fewer species were affected by the LQE in Africa and Eurasia, a similar size-biased extinction has been observed with end-Pleistocene (~13 ka) ages being prominent. Current explanations for the LQE in North America and Australia involve a combination of two hypotheses: climate change, and "overkill" by human hunting, modulated by the knock-on effect of herbivore extinction on the environment and on the survivability of other groups (e.g., Owen-Smith, 1987). Although "overkill" was originally used to explain North American extinctions (Martin, 1967), a forerunner of the hypothesis was popular in 19th century Europe, before it was eventually abandoned as archaeological evidence for human migration showed little evidence for the impact of human hunting on the LQE. Grayson and Meltzer (2003) argued that island settings (e.g., New Zealand or the West Indies), where human hunting and habitat degradation can be unequivocally associated with extinction, should not be the model for continental extinctions. Extraterrestrial impact as a contributing cause for the LOE in North America, and for the onset of the Younger Dryas cold period (Firestone et al., 2007), has not been supported by subsequent analyses (Holliday et al., 2014; Pinter et al., 2011).

In North America, the brief (\sim 200 year) duration of Clovis-tool finds at \sim 13 ka (e.g., Waters & Stafford, 2007) is usually associated with rapid dispersal of modern humans across North America and is closely contemporaneous with the LQE peak. On the other hand, the lack of mammalian kill sites in the Clovis record argues against a direct linkage between Clovis technology and "overkill." Humans were present in North America at least several kiloyear prior to the Clovis horizon (Gilbert et al., 2008; Waters et al., 2011), and perhaps prior to \sim 24 ka (Bourgeon et al., 2017).

According to Faith and Surovell (2009), the LQE in North America was abrupt and requires a mechanism capable of wiping out ~35 genera across the continent in a "geological instant" in the 13.8- to 11.4-ka interval (Figure 2), with the spread in last appearances being largely explained by an incomplete fossil record and the resulting Signor-Lipps effect (Signor & Lipps, 1982). Abrupt versus staggered megafaunal extinction at the LQE is central to determination of cause. In their "continental simulation," Faith and Surovell (2009) determined the empirical probability (3.4%) of observing a terminal Pleistocene (10-12 ka) age from 1955 stratigraphic occurrences (from 31 genera) of which 66 taxa (from 16 genera) yield terminal Pleistocene ages, assuming that all occurrences are equally likely to receive a terminal Pleistocene age. The simulation randomly assigned pre-12-ka or post-12-ka ages to all 1955 observations based on this probability (3.4%). The total number of genera that received a terminal Pleistocene age was tallied for each of 10,000 simulations to determine the probability of observing 16 or fewer terminal Pleistocene genera. The authors concluded that the observed pattern is consistent with synchronous (i.e., 10-12 ka) extinction for all 31 genera.

Bradshaw et al. (2012) proposed a Gaussian-resampled, inverse-weighted McInerny (GRIWM) approach, which weights observations inversely according to their temporal distance from the last observation of a confirmed species occurrence, and samples radiometric ages from the underlying probability distribution. In Figure 2, we show GRIWM estimates of continent-wide European extinctions from the fossil record aided by DNA analyses (Cooper et al., 2015), excluding regional disappearances. An extinction age estimate in North America for *Arctodus simus* (the short-faced bear) at 10.8 ka (Schubert, 2010) and the onset of population decline of *Bison priscus* (the steppe bison) at ~37 ka (Shapiro et al., 2004) are included in Figure 2. Note that the horse and woolly mammoth (*Mammuthus primigenius*) persisted in interior Alaska until ~10.5 ka (Haile et al., 2009), and the woolly mammoth survived on St. Paul Island (Alaska) until ~5.6 ka (Graham et al., 2016). Zazula et al. (2014) pointed out that the American mastodon (*Mammut americanum*) occupied eastern Beringia (Alaska/Yukon) during the last interglacial before its range contracted southward at the onset of glacial conditions at ~75 ka. The range of the species appears to have expanded northward again as interglacial conditions returned at the end of the Pleistocene, before extinction of the species at ~11.5

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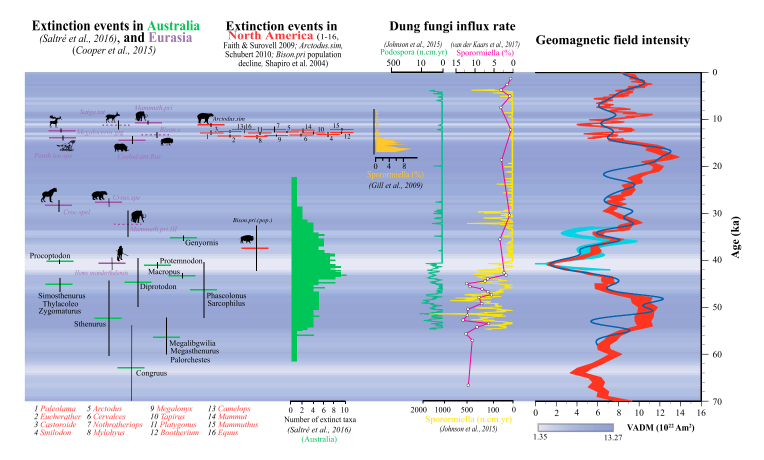


Figure 2. Geomagnetic field intensity (VADM) for the last 70 kyr compared with timing of continent-wide extinction events for genera from Australia (Saltré et al., 2016), Eurasia (Cooper et al., 2015), and North America (Faith & Surovell, 2009), including extinction of *Arctodus simus* (Schubert, 2010) and population decline for *Bison priscus* (Shapiro et al., 2004). The solid and dashed bars in the Eurasia record refer to fossil and DNA derived events, respectively. Last appearance datums for extinct North American genera (Faith & Surovell, 2009) were recalibrated using Calib 7.1 (Stuiver et al., 2018) with 2σ errors for entries that pass selection criteria established by Meltzer and Mead (1985). Dung fungi (*Sporormiella* and *Podospora*) in sedimentary archives, proxies for herbivore population, are for North America (Gill et al., 2009) and Australia (Johnson et al., 2015; van der Kaars et al., 2017). Geomagnetic field intensity during the last 70 kyr (red) combines VADM for the last 14 kyr (Pavón-Carrasco et al., 2014), VADM for 14-45 ka (Channell et al., 2018), and the PISO VADM record (Channell et al., 2009) for 45-70 ka. The light blue band represents the VADM estimate from 36 Cl-flux in Greenland ice-cores (Muscheler et al., 2005). The dark blue line corresponds to the ice-core 10 Be-derived VADM record (Channell et al., 2018). The background shades of blue are scaled to the maximum (darker blue color) and minimum (white) VADM intensity in the 1.35-13.27 × 10^{22} Am 2 range.

ka (10,000 ¹⁴C years before present [BP]). Zazula et al. (2014) posed the question: why was this species stopped in its tracks when favorable conditions beckoned in Beringia?

An important proxy for herbivore population is the abundance in sedimentary sequences of coprophilous (dung) fungal spores, such as *Sporormiella*. The proxy was first proposed over 30 years ago (Davis, 1987), requires careful interpretation and laboratory techniques (e.g., van Asperen et al., 2016), but provides a measure of herbivore population independent of the bone-fossil record. Lake sediments in New York and Indiana imply a decline in *Sporormiella* spores beginning at 14.8 ka (Figure 2) that falls below the 2% threshold by 13.7 ka (Gill et al., 2009). This result has been closely replicated at Silver Lake (Ohio) where *Sporormiella* decline was dated at 13.9 ka (Gill et al., 2012). Importantly, the *Sporormiella* decline at these sites predates Younger Dryas cooling, and concurrent changes in the pollen record, and immediately predates a marked charcoal deposition increase, implying that herbivore decline and resulting landscape changes provide an explanation for subsequent (natural) landscape burning. The onset of the demise of North American herbivores at ~14.5 ka (Gill et al., 2009) lies within the Bølling-Allerød warm period with the Younger Dryas cold period beginning ~2 kyr later (e.g., Deplazes et al., 2013).

The South American LQE was even more profound than that in North America (Barnosky & Lindsey, 2010; Koch & Barnosky, 2006), with the loss of 50 megafaunal genera (~83%). Robust dates are scarce for the South American LQE, although it appears that many taxa were lost near the Pleistocene-Holocene boundary

(Barnosky & Lindsey, 2010). *Sporormiella* decline in lake sediments from SE Brazil implies herbivore population collapse at ~12 ka (Raczka et al., 2018).

In Northern Eurasia, nine genera (35%) were lost during the LQE. Available age data are consistent with a two-phase extinction in the 45- to 35-ka and 15- to 10-ka intervals (Koch & Barnosky, 2006). Up to 50% of worldwide megafaunal extinctions at 15-10 ka apparently occurred in Northern Eurasia (Cooper et al., 2015), but the extinction pattern is more complex than in North America with megafaunal range contractions culminating in extinction for some species but not others (Stuart, 2015). In Figure 2, we plot continent-wide Eurasian megafaunal extinction events from Cooper et al. (2015). Several well-studied species disappeared continent-wide at ~26-31 ka (Figure 2); hence, their last appearances significantly postdate the Laschamp excursion (at ~41 ka). Post-Laschamp extinction for *Crocuta crocuta* (spotted hyaena) and *Crocuta spelaea* (cave hyaena) at ~26 ka were, however, preceded by severe range contraction from Asia into Europe (Stuart & Lister, 2014). Fossils of *Ursus spelaeus* (cave bear) also indicate E to W range contraction before extinction at ~26 ka with abrupt population decline, based on DNA analyses, after 50 ka (Baca et al., 2016; Stiller et al., 2010; Stiller et al., 2014).

In Africa, at least 24 species and ~10 genera of mammals became extinct in the 13- to 6-ka interval, representing 25% of Pleistocene African megafauna (Faith, 2014). Species-level extinction was, again, most intense for larger-bodied megafauna (Koch & Barnosky, 2006). The African LQE was considered to have been less severe than elsewhere, accounting for the relatively rich diversity of modern African megafauna. On the other hand, the number of extinct African species dated to the last 100 kyr exceeds the number in Europe and matches the number in Australia, and is only surpassed by the LQE in the Americas (Faith, 2014). In east Africa alone, the number of securely dated latest Pleistocene mammal extinctions has risen from two to seven in the last decade, with most being of grazers associated with open habitats (Faith, 2014).

Estimation of extinction ages for Australian megafauna (and for some Eurasian genera) is complicated by the majority of last appearances being at or beyond the practical range of radiocarbon dating (i.e., >40 ka). The LQE in Australia was apparently catastrophic for large mammalian megafauna, with the complete loss of all animals heavier than ~100 kg. Fourteen of 16 genera of Pleistocene mammalian megafauna disappeared, together with all megafaunal reptiles (6 genera), in the vicinity of, or prior to, ~40 ka (Figure 2). Ten Australian genera disappeared in the 44- to 35-ka interval based on a variety of frequentist statistical methods (including GRIWM) to determine extinction ages for 16 megafaunal genera (Figure 2; Saltré et al., 2016). The mass extinction of megafauna at this time, including the largest-known (~3,000 kg) marsupial (Diprotodon), has been linked with climate variability and aridity (e.g., Wroe et al., 2013), although this linkage has been disputed (e.g., Saltré et al., 2016), often in favor of human predation or "overkill" (e.g., Brook & Johnson, 2006; Johnson et al., 2016; Miller et al., 2016; van der Kaars et al., 2017). It is noteworthy that the extinction age for the ~200-kg flightless bird Genyornis newtoni at ~35 ka (Figure 2; Saltré et al., 2016) is younger than the ~43-ka estimate given by Miller et al. (2016) based on dated eggshell fragments. Even if final extinction was delayed until ~35 ka, the population of Genyornis newtoni crashed close to the time of the Laschamp excursion (~41 ka) based on the egg-shell data (Miller et al., 2016), although some egg-shells attributed to Genyornis newtoni may be from other species (Grellet-Tinner et al., 2016).

At Lynch's Crater (NE Queensland), an abrupt decline in dung fungi including *Sporormiella* (Figures 2 and 3) implies abrupt demise of large Australian herbivores at 40-44 ka (Johnson et al., 2015). An abrupt increase in charcoal lags *Sporormiella* decline by ~100 years, and evidence for grasses and sclerophyll vegetation lags *Sporormiella* decline by ~300-400 years (Johnson et al., 2015; Rule et al., 2012). The charcoal-rich levels can be explained by natural lightning-induced biomass burning as a result of fuel buildup triggered by herbivore extinction (Johnson et al., 2016; Rule et al., 2012). Off the southern coast of Western Australia, marine core MD03-2614G records a sharp decline in *Sporormiella* in the 45- to 43-ka interval, relative to values recorded back to 140 ka (Figures 2 and 3; van der Kaars et al., 2017).

A role for humans in the extinction of large animals in Australia remains popular (e.g., Brook & Johnson, 2006; Johnson et al., 2016; Miller et al., 2016; Turney et al., 2008; van der Kaars et al., 2017), although the arrival of humans in Australia (Sahul) may have predated the LQE at ~40 ka by ~25 kyr (Clarkson et al., 2017) but the arrival date is not unequivocal (O'Connell et al., 2018). There is no evidence for a spike in the human population in Australia at the time of the most prominent extinction event at ~40 ka, when the entire Australian human population may not have exceeded a few tens of thousands (Williams, 2013).

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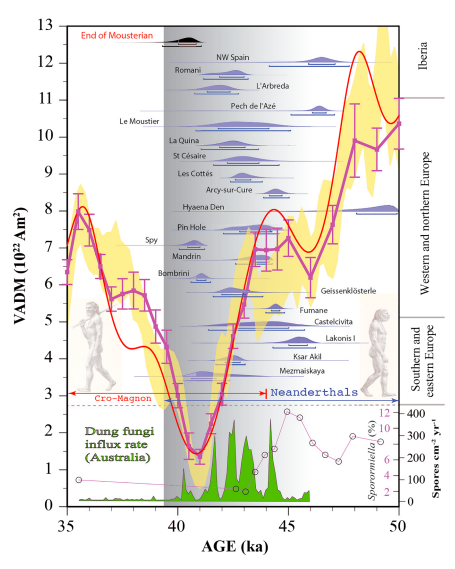


Figure 3. Geomagnetic field intensity (VADM) for the 35- to 50-ka interval compared to probability density functions for the timing of Neanderthal disappearance at different sites (blue) and for the end of the Mousterian tool industry (black; Higham et al., 2014), with dung-fungal proxies for large herbivore extinctions from NE Queensland (green; Johnson et al., 2015) and southern Western Australia (open circles joined by purple line; van der Kaars et al., 2017). The yellow band represents the GLOPIS paleointensity record converted to VADM (Laj et al., 2004, 2014), the purple curve is the sedimentary VADM record with 2σ errors, and the red line is the VADM reconstructed from 10 Be flux in Greenland ice-cores (Channell et al., 2018). The gray shading indicates the age range associated with Neanderthal demise, and the temporal overlap between the two human groups (Neanderthals and Cro Magnon). The end of the Mousterian flint-tool industry has been dated at 39.260-41.030 ka (Higham et al., 2014), and 39.528-41.013 ka (Cooper et al., 2015), corresponding closely with low field intensity associated with the Laschamp geomagnetic excursion (39.7-41.9 ka; Laj et al., 2014).

Tasmania had a land bridge to the Australian mainland during the last glacial, becoming an island in the early Holocene. The extinction of megafauna in Tasmania at ~40 ka does not correspond to climate or environmental change and has been associated with the late arrival of humans in the region (Turney et al., 2008). More recent results place the Tasmanian extinction of *Protemnodon anak* and other megafauna at ~41 ka, predating human arrival on the island at ~39 ka and hence precluding human involvement in the extinctions (Cosgrove et al., 2010; Lima-Ribeiro & Diniz-Filho, 2014). Extant smaller (more accessible) prey, particularly the common wombat (*Vombatus ursinus*) and the red-necked wallaby (*Macropus rufogriseus*), characterize the early archaeological kill-sites on the island (Cosgrove et al., 2010).

Apart from Australian extinctions concentrated close to the Laschamp excursion at ~40 ka, fossils from the King's Creek Catchment (SE Queensland) indicate additional concentrations of megafaunal last

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appearances at ~83, ~107, and ~122 ka (Price et al., 2011; Wroe et al., 2013). The older two dates (107 and 122 ka) correspond to magnetic field intensity minima associated with the Blake excursions (Figure 1b).

4. Neanderthal Extinction

The extinction of Neanderthal represents one of the great puzzles of human evolution. Neanderthal and anatomically modern humans (AMHs) cohabited Western Europe for ~2-5 kyr, prior to ~39 ka, supporting the contention that competition may have contributed to the demise of Neanderthal (Higham et al., 2014). Brief cold and dry conditions in Europe associated with Heinrich Stadial (HS) 4 were proposed as an additional likely stressor on Neanderthal (Sepulchre et al., 2007). Analyses of Campanian Ignimbrite (CI) cryptotephra from archaeological sites in Greece and elsewhere in Eastern Europe and Libya indicated that the CI eruption occurred early in a dry period associated with HS4, postdated the end of the Middle Paleolithic and the Mousterian tool industry, and hence postdated the demise of Neanderthal (Lowe et al., 2012). In Black Sea sediment cores, CI tephra overlie, and therefore postdate, the Laschamp excursion (Nowaczyk et al., 2012).

The extinction of Neanderthal and the demise of the Mousterian tool industry (Figure 3) can be placed at 41,030–39,260 calibrated years before present (41-39 ka) with 95.4% probability (Higham et al., 2014). Cooper et al. (2015) estimated the extinction of Neanderthal at 41,227 calibrated years BP with a standard deviation of 219 years, and 39,528-41,013 calibrated years BP using the GRIWM method (Figure 2). Mousterian ages outside this range have been recorded at several locations in southern Iberia including Gorham's Cave in Gibraltar (Finlayson et al., 2006; Tzedakis et al., 2007), but these ages should now be disregarded according to Higham et al. (2014). Recent findings cast doubt on the existence of Neanderthal after ~39 ka and lead to a closer correspondence of the demise of Neanderthals with the Laschamp magnetic excursion and the associated brief interval of very low geomagnetic field intensity centered at ~41 ka (Laj et al., 2014). It is important to note that the IntCal13 radiocarbon calibration (Reimer et al., 2013) may be offset to older ages by ~1 kyr in the vicinity of the Laschamp excursion, relative to ice-core chronologies (Muscheler et al., 2014).

Valet and Valladas (2010) proposed that low magnetic field strength in the Laschamp/Mono Lake excursion interval (40-33 ka) was an important factor in Neanderthal demise. Why AMHs were not similarly affected has remained an open question considering that the two populations shared habitats for 2,600-5,400 years (Higham et al., 2014) or >5 kyr (Lowe et al., 2012). There is no evidence for differences in skin pigmentation between European AMHs and Neanderthals, and at least a fraction of Neanderthals apparently had the same pale skin and/or red hair observed in some modern humans (Lalueza-Fox et al., 2007). Natural skin pigmentation in humans mitigates the harmful effects of UVR, but its advantage is offset by the importance of sunlight for vitamin D3 synthesis. The skin protection factor of "red ochre" (hematitic iron oxides) is traditionally utilized by some African tribes (Rifkin et al., 2015) and has been used since at least the last interglacial (~120 ka) based on ochre coatings on strung beads and residues on storage shells from Africa (Hodgskiss & Wadley, 2017), SE Spain (Hoffmann et al., 2018), and Levantine Mousterian sites (Bar-Yosef Mayer et al., 2009). The mystery of AMH survival at the time of Neanderthal demise may have been resolved by the discovery of differences in amino acid substitution in an intracellular chemosensor (the AhR) for AMHs and for Neanderthals and other primates (Hubbard et al., 2016).

5. The Role of the AhR

Defense mechanisms against UVR include the production of quenching agents and antioxidant enzymes that neutralize ROS produced by UVR. The ROS modulation is controlled by the AhR intracellular chemosensor that plays a key role in the evolutionary response to UVR. Experimental results indicate an adaptive response of mitochondria to varying ROS levels under a phenomenon called mitohormesis (Becker et al., 2016).

The primary role of AhR is to regulate the transcription of genes mediating responses to the biochemical and toxic effects of dioxins, polyaromatic hydrocarbons, and related compounds (Abel & Haarmann-Stemmann, 2010). AhR is expressed in all skin cells and can be generated by UVR through an endogenous ligand formed in situ from an amino acid called tryptophan (Esser et al., 2009). UVR and the more harmful UVB (wavelength 290-320 nm) are absorbed by free tryptophan in the cytosol of epidermal cells, and AhR plays a key role in translocating UVR stress response to the nucleus (Fritsche et al., 2007; Tigges et al., 2014; Wei et al., 1999). Exposure to UVR, particularly UVB, generates highly mutagenic DNA photoproducts. The process initiates apoptosis and involves damage to nuclear DNA accompanied by mitochondrial dysfunction

(Frauenstein et al., 2013). There is a general consensus that the AhR of modern humans is implicated in DNA repair (Dittmann et al., 2016; Schreck et al., 2009), tumor suppression (Fan et al., 2010; Yu et al., 2017), epidermal barrier function (Noakes, 2015), skin tanning response, and melanocyte homeostasis (Jux et al., 2011; Luecke et al., 2010). Phylogenetic analysis suggested that the ability of vertebrate AhR to sense xenobiotics was acquired at a late stage of evolution, implying that the driving force for evolutionary conservation of AhR lies not only in its role in xenobiotic metabolism but also in normal cell development (Hahn et al., 2017; Hao & Whitelaw, 2013).

Hubbard et al. (2016) showed that the AhR variant in modern humans contains Val381 residue in the ligand-binding domain, while the AhR of Neanderthals, and a Denisovan individual, as well as nonhuman primates and other vertebrates (rodents) encode the ancestral Ala381 variant. The Val381 variant is fixed in the genome of all modern humans as well as in the genome of the oldest (45 ka) AMH individual sequenced to date (Fu et al., 2014). Hubbard et al. (2016) suggested that the unique modification of AhR in AMHs led to significant competitive advantage over their Neanderthal neighbors, due to decreased sensitivity in AMHs to toxins associated with fire-smoke, the effects of which may have been exacerbated by troglodytic lifestyles.

Our focus here is on AhR involvement in the regulation of the skin responses to UVR, especially to harmful UVB radiation, and its modulation of the immune system (Agostinis et al., 2007; Esser et al., 2013; Rannug & Fritsche, 2006). UVB induces two signaling routes in mammalian cells: first, UVB is absorbed by nuclear DNA that results in generation of DNA photoproducts, and second, UVB activates cell-surface receptors (Merk et al., 2004). AhR plays an important role in skin integrity and immunity. AhR activation leads to transcriptional gene activation and is involved in the cutaneous stress response to UVR (Agostinis et al., 2007; Dittmann et al., 2016; Navid et al., 2013; Schwarz, 2005) and alterations of gene expression (Dugo et al., 2012). Activation of AhR by UVB leads to signaling both to the nucleus and to cell membranes (Fritsche et al., 2007). The findings show that UVB irradiation affects cell surface receptors with subsequent activation of mitogen-activated protein kinases that in turn affect DNA in the nucleus.

6. The Early Fossil Record of Homo sapiens

From analyses of the fossil and stone-tool record over the last 250 kyr (Figure 4), Lahr (2016) proposed five transitions in the evolutionary history of *Homo sapiens*: (1) the origins of the species at 240–200 ka, (2) the first major expansion at 130–100 ka, (3) a period of dispersals at 70–50 ka, (4) a period of local/regional structuring of diversity at 45–25 ka, and (5) an early Holocene phase of significant extinction of hunter-gatherers and expansion of farmers (the Holocene Filter).

Until recently, Member 1 of the (Omo) Kibish Formation of Ethiopia yielded the earliest known AMH cranial remains (Day, 1969). A volcanic tuff about 2 m below the level of the fossil finds has a 40 Ar/ 39 Ar weighted mean age of 196 \pm 2 ka (McDougall et al., 2005). An age of 196 ka for Member 1, combined with sedimentological evidence for rapid deposition, is consistent with Member 1 having been deposited synchronously with Mediterranean sapropel S7 (Brown et al., 2012; McDougall et al., 2008). Jebel Irhoud, Morocco, has been an important archaeological site since the 1960s when human fossils were found alongside Mousterian stone tools and were once dated at ~40 ka (Ennouchi, 1962). Recent fossil discoveries at this location support the presence of fossils with characteristics of AMHs (Hublin et al., 2017), and new luminescence age dating (Richter et al., 2017) indicates that the fossils are considerably older, at ~300 ka, than the Ethiopian AMH finds at Omo Kibish (Figure 4). The finds in Morocco are now among the earliest known hominin fossils with AMH characteristics, in common with characteristics of the Florisbad fossil from South Africa discovered by T. F. Dreyer in 1932 and dated at ~260 ka (Grün et al., 1996). With such a sparse hominin fossil record, it is not possible to predict when AMHs first appeared, although the period from 300 ka to 200 ka in Africa appears to have been a critical time in development of AMH, and may constitute the evolutionary cradle of our species.

7. Time to Most Recent Common Ancestor (TMRCA) From mtDNA and Y Chromosomes

Thirty years ago, Cann et al. (1987) demonstrated that ethnically diverse surveys of modern mitochondrial DNA (mtDNA) are a major source of human evolutionary history. The advantages of mtDNA as an evolutionary tool include faster mutation rates in mtDNA than in nuclear genes, and mtDNA is inherited maternally and does not recombine. Cann et al. (1987) demonstrated the African origin of the human mitochondrial

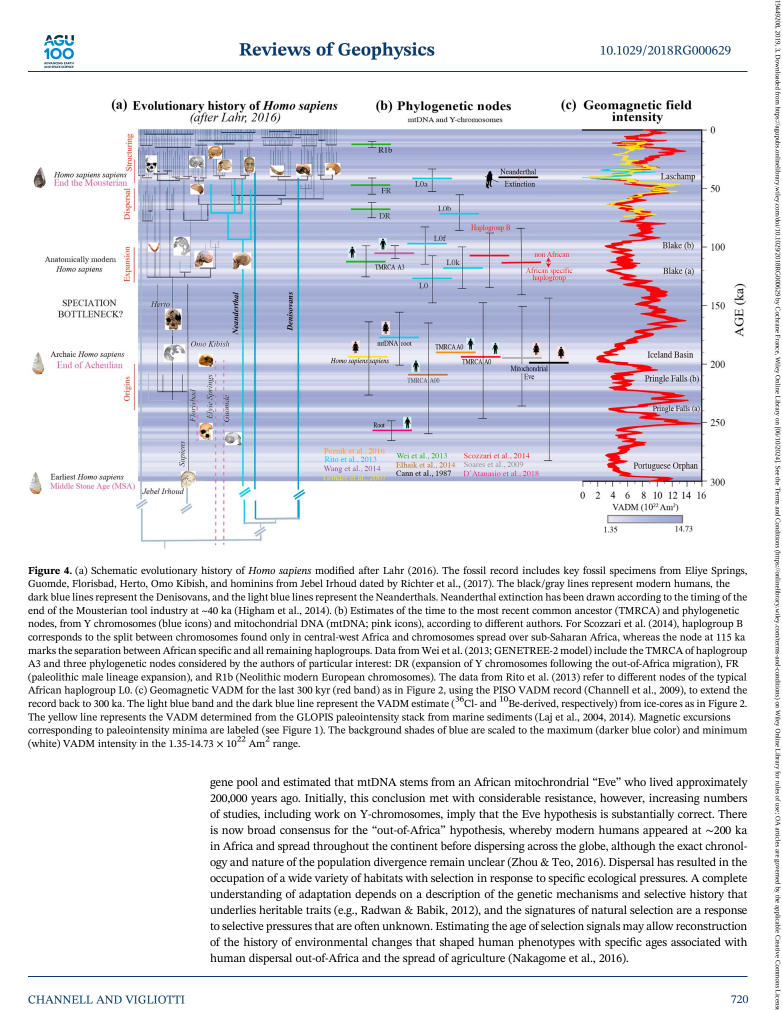


Figure 4. (a) Schematic evolutionary history of Homo sapiens modified after Lahr (2016). The fossil record includes key fossil specimens from Eliye Springs, Guomde, Florisbad, Herto, Omo Kibish, and hominins from Jebel Irhoud dated by Richter et al., (2017). The black/gray lines represent modern humans, the dark blue lines represent the Denisovans, and the light blue lines represent the Neanderthals. Neanderthal extinction has been drawn according to the timing of the end of the Mousterian tool industry at ~40 ka (Higham et al., 2014). (b) Estimates of the time to the most recent common ancestor (TMRCA) and phylogenetic nodes, from Y chromosomes (blue icons) and mitochondrial DNA (mtDNA; pink icons), according to different authors. For Scozzari et al. (2014), haplogroup B corresponds to the split between chromosomes found only in central-west Africa and chromosomes spread over sub-Saharan Africa, whereas the node at 115 ka marks the separation between African specific and all remaining haplogroups. Data from Wei et al. (2013; GENETREE-2 model) include the TMRCA of haplogroup A3 and three phylogenetic nodes considered by the authors of particular interest: DR (expansion of Y chromosomes following the out-of-Africa migration), FR (paleolithic male lineage expansion), and R1b (Neolithic modern European chromosomes). The data from Rito et al. (2013) refer to different nodes of the typical African haplogroup L0. (c) Geomagnetic VADM for the last 300 kyr (red band) as in Figure 2, using the PISO VADM record (Channell et al., 2009), to extend the record back to 300 ka. The light blue band and the dark blue line represent the VADM estimate (36 Cl- and 10 Be-derived, respectively) from ice-cores as in Figure 2. The yellow line represents the VADM determined from the GLOPIS paleointensity stack from marine sediments (Laj et al., 2004, 2014). Magnetic excursions corresponding to paleointensity minima are labeled (see Figure 1). The background shades of blue are scaled to the maximum (darker blue color) and minimum (white) VADM intensity in the 1.35-14.73 \times 10²² Am² range.

gene pool and estimated that mtDNA stems from an African mitochrondrial "Eve" who lived approximately 200,000 years ago. Initially, this conclusion met with considerable resistance, however, increasing numbers of studies, including work on Y-chromosomes, imply that the Eve hypothesis is substantially correct. There is now broad consensus for the "out-of-Africa" hypothesis, whereby modern humans appeared at ~200 ka in Africa and spread throughout the continent before dispersing across the globe, although the exact chronology and nature of the population divergence remain unclear (Zhou & Teo, 2016). Dispersal has resulted in the occupation of a wide variety of habitats with selection in response to specific ecological pressures. A complete understanding of adaptation depends on a description of the genetic mechanisms and selective history that underlies heritable traits (e.g., Radwan & Babik, 2012), and the signatures of natural selection are a response to selective pressures that are often unknown. Estimating the age of selection signals may allow reconstruction of the history of environmental changes that shaped human phenotypes with specific ages associated with human dispersal out-of-Africa and the spread of agriculture (Nakagome et al., 2016).

Both mtDNA and Y-chromosomes have been used to reconstruct human history. The former (mtDNA) is often the better option for analyzing ancient DNA because it is easier to obtain, is present in higher number in human cells, and does not undergo recombination. However, mtDNA reflects only the maternal history of a population and the history of a single individual may not accurately reflect the history of a population. For this reason, mtDNA studies should be complemented by data on the male-specific Y chromosome (Pakendorf & Stoneking, 2005). Rates of human evolution have been estimated by applying a number of substitution models to mtDNA or Y-chromosome sequence data (Behar et al., 2008; Kivisild, 2015; Soares et al., 2013; Wang et al., 2014). In some examples, calibration has relied on the assumption that the genetic separation between humans and chimpanzees occurred at ~6 Ma and that the evolutionary process has been clocklike since that time, while more complex models involve different substitution rates for coding and control regions of the genome (see Endicott et al., 2009). Molecular clocks can be tested by comparison with archaeological data pertaining to human migration.

Phylogeny based on the first complete mtDNA sequence data available in year 2000 (from 52 modern individuals selected from around the world) yielded a time to the most recent common ancestor (TMRCA) of ~190 ka (seeOppenheimer, 2003, 2009; Soares et al., 2009), apparently consistent with the conclusions of Cann et al. (1987) and with dated fossil finds of early AMH in Ethiopia (McDougall et al., 2005). More recent studies of human evolution from mtDNA (Oppenheimer, 2012), and of paternal evolution from Y chromosomes (Poznik et al., 2016), yielded a broadly consistent picture of human evolution over the last ~200 kyr although the timing of branches in the evolutionary tree is poorly constrained and depends on estimated rates of mutation and population size (Figure 4). As pointed out by Wang et al. (2014), Y-chromosomal substitution rates obtained using different calibration modes vary considerably and produce disparate reconstructions of human history. An additional determinant of substitution rate is the efficacy of purifying selection, which in turn depends not only on the particular constraints of each chromosome but also on the long-term effective population size for each chromosome type (Elhaik et al., 2014). Furthermore, mutation rates may have changed during hominoid evolution (Scally, 2016).

The TMRCA is not a unique number, but rather a probability distribution based on two fundamental assumptions: the number of mutations and the mutation rate. Estimates of TMRCA depend strongly on the substitution rate, and different results can be obtained by using different rates. For example, Mendez et al. (2013) estimated a very early date of 338 ka for the TMRCA of the Y-chromosome tree (A00) from a population of African-Americans. The authors explained this early age by either long-standing population structure among modern human populations or archaic introgression from unknown species into the ancestors of modern humans in western Central Africa. However, other researchers (e.g., Wang et al., 2014) have pointed out that this ancient TMRCA can be partially attributed to the low substitution rate used by the authors. By using either a higher mutation rate or more extensive sequencing data, the estimate of TMRCA becomes much younger at about 208 ka (Elhaik et al., 2014) or 257 ka (D'Atanasio et al., 2018).

Poznik et al. (2013) reported entire Y-chromosome and mitochondrial genome sequences using a within-human calibration point to estimate the substitution rate. The results indicate small TMRCA differences from Y chromosomes (120–156 ka) and mtDNA (99-148 ka) that disagree with the conventional suggestion that the common ancestor of male lineages lived considerably more recently than that of female lineages. Analyzing the entire Y-chromosome dataset of the 1,000 Genomes Project (1,000 Genomes Project et al., 2012; 1,000 Genomes Project, 2015) using the pedigree-based substitution rate, Wang et al. (2014) estimated the TMRCA at 105 ka that is consistent with the estimate (105 ka) of Cruciani et al. (2011) and the estimate (101-115 ka) of Wei et al. (2013).

Phylogeny based on mtDNA sequence data has provided an estimate of TMRCA (190 ka) with branching episodes at ~120, ~70, ~40, and ~15 ka, which coincide with AMH dispersal patterns (Oppenheimer, 2012). Wei et al. (2013) used five substitution models to assess phylogenetic nodes from the TMRCA of 29 Y chromosomes that yielded branching ages for one of the models (GENETREE-2) of 112 ± 12 ka, 68 ± 7 ka, 49 ± 6 ka, and 13 ± 2 ka. Applying the BEAST method to 68 worldwide Y chromosomes, Scozzari et al. (2014) dated the first two splits in their tree at ~196 and ~167 ka, followed by TMRCAs at ~110 ka, 85-77 ka, 51-33 ka, and 8-22 ka (Figure 4).

It remains challenging to reconstruct population structure prior to ~60 ka using existing data, but the modern-human African mtDNA pool contains phylogenetic patterns that can be used to estimate the ages

of several haplogroups. Analyzing L0 HVS-I sequences, Rito et al. (2013) classified five branches (L0a, L0b, L0d, L0f, and L0k) of the mtDNA tree. The age estimates indicate that the mtDNA tree split to form L0 at \sim 180 ka, and later diversity follows a geographical distribution from southern Africa northward. Around \sim 128 ka, two distinct AMH groups coexisted in Africa with a first split around 119 ka (L0k) followed by major clades at 98.7 ka (L0f), 70.9 ka (L0b), and 42.4 ka (L0a). It is noteworthy that the splitting of the widespread and common L0a, the diversification of L0k, and population increases of haplogroups L0 and L3 can be dated to \sim 40 ka (Figure 4).

8. Discussion

Past episodes of low geomagnetic dipole field strength affect UVR levels arriving at the Earth's surface because reduction in magnetospheric shielding results in lower stratospheric ozone levels, and hence lowered UVR shielding (Wei et al., 2014). Solar storms enhance nitrogen oxide production in the stratosphere that, in early 2004, led to ozone reduction of more than 60% at high northern latitudes (Randall et al., 2005). Similarly, atmospheric N_2O concentrations are enhanced during times of low geomagnetic field strength when shielding from solar storms and GCRs is diminished. Earth's surface naturally emits N_2O from the oceans and from soils, with emissions having increased due to anthropogenic practices. N_2O emissions are enhanced during interglacial and interstadial (warm) climate states, and increased by ~50% at the last glacial termination (Schilt et al., 2013, 2014). Enhanced atmospheric N_2O concentrations during the Bølling-Allerød warm period, which coincided with an apparent geomagnetic field strength minimum at ~13 ka, would have elevated UVR reaching the Earth's surface at this time. It is noteworthy that the drastic magnetic field intensity decrease after the peak at 18 ka (Figure 1) coincided with paleoclimatic changes especially in the southern hemisphere (Boex et al., 2013; Moreno et al., 2015; Martínez-García et al., 2014). Evidence from Antarctic ice cores indicated sudden enhanced tropospheric UVR that has been related to a Mount Tahahe eruption dated to 17.7 ka (McConnell et al., 2017).

A selection process resulted in Neanderthal disappearance at ~41 ka that apparently did not affect AMHs. Previous hypotheses for Neanderthal disappearance, and expansion of AMHs, include differences in subsistence strategies, language skills, and technical, economic, and social systems, and the ability to adapt to changing environments (e.g., Kochiyama et al., 2018). Response to environmental change such as UVR flux at Earth's surface would have involved the AhR, a chemosensor that regulates immunity and differs in AMHs versus Neanderthals and other primates (Hubbard et al., 2016). Ages for the end of the Mousterian tool industry and Neanderthal demise (Higham et al., 2014) are now tightly constrained to the Laschamp magnetic excursion (at 41 ka) implying a role for high UVR levels during the Laschamp field intensity minimum (Figure 3).

Prominent low geomagnetic field intensity episodes at 285 ka, 190 ka, 100-120 ka, 64 ka, 41 ka, and 13 ka (Figure 1) appear to correspond to important times in evolution of hominins and other large mammals. At the LQE, megafauna in Australia, Europe, and the Americas were thought to have become extinct over a protracted time in the Late Quaternary; however, improved age estimates and discovery of new fossil sites (Barnosky & Lindsey, 2010; Cooper et al., 2015; Faith, 2014; Faith & Surovell, 2009; Koch & Barnosky, 2006; Price et al., 2011; Roberts et al., 2001; Saltré et al., 2016; Stuart, 2015; Wroe et al., 2013) have led to extinction peaks becoming progressively constrained to ~13 ka in the Americas, to ~40 ka (with earlier episodes at 84, 107, and 122 ka) in Australia, and a complex combination of Late Pleistocene ages (including ~13 and ~40 ka) in Eurasia and Africa.

One of the outstanding and intriguing aspects of the LQE is the strong correlation between extinction and body mass. As the vast majority of cell mutations are deleterious, large long-lived organisms are at an evolutionary disadvantage. In addition, small mammals often have opportunities to avoid UVR through burrowing. In modern mammal populations, there is, however, no apparent correlation between body mass and cancer occurrence, known as Peto's Paradox (Peto et al., 1975). The elephant genome includes 20 copies of an oncogene (TP53) that is a crucial tumor suppressor gene involved in apoptosis in response to DNA damage, whereas other mammals usually have small numbers of this gene (Abegglen et al., 2015; Sulak et al., 2016). This discovery may explain why elephants are one of few large mammals to pass the LQE barrier. On the other hand, the remains of two extinct mammoth species included more than a dozen copies of TP53 in their genomes, and the American mastodon, that disappeared at ~ 50 ka, had 3-8 copies in its genome

which implies an evolutionary selective trend in the increased number of copies of TP53 in the genomes of these related megafauna (Sulak et al., 2016). TP53 is considered the guardian of the genome due to its role in mitigating DNA damage, and is itself a target of UV-induced mutations (Aylon & Oren, 2011; de Pedro et al., 2018).

Huang et al. (2017) analyzed the body weight of species in two orders of large ungulate herbivores (Artiodactyla and Perissodactyla) from the Neogene fossil record in Europe and North America. They found a significant and progressive increase in body weight from early Miocene to late Pliocene for both orders in North America and for Artiodactyla in Europe. This work was followed by the analysis of Smith et al. (2018) who documented a global increase in body-size of megafauna during the Cenozoic, with abrupt downsizing at the LQE in Late Pleistocene. We speculate that large mammals may have reached a natural body-size limit by Late Pleistocene, due to increased likelihood of cell mutation at times of high UVR flux, as life spans and body weights increased during the Cenozoic. The envisaged role of UVB in reducing megafaunal populations at the LQE does not involve an instantaneous "blitzkrieg," but rather an accumulation of UVB-triggered mutations over multiple (~30) generations or ~0.5-1 kyr, the approximate duration of RPI minima associated with magnetic excursions.

There is no clear correlation of LQE events, either at ~13 ka (in North America and Eurasia) or ~40 ka (in Australia and Eurasia), with the first appearances of humans, which apparently preceded the LQE by at least 10 kyr in Australia, and at least several kyr in North America. The human population of Australia at ~40 ka was likely no more than a few tens of thousands, with no evidence of an increase in population at this time (Williams, 2013). According to Webb (2013), the "overkill" hypothesis "is more sensational, granted, but the arguments are unrealistic and the evidence for it, at least in Australia, is non-existent." The case for "overkill" in North America at ~13 ka involves the close coincidence of the LQE with the brief (few century duration) Clovis-tool horizon. On the other hand, based on Sporormiella decline in lake sediments from the eastern United States, the demise at North American herbivores began at ~14.5 ka (Gill et al., 2009, 2012), prior to the first appearance of Clovis tools. Clovis tools have only rarely been found in association with megafaunal remains. Evidence for pre-Clovis human occupation in the Americas includes locations in southern Chile (Dillehay et al., 2015), and the Florida panhandle where human occupation at ~14.5 ka predated the Sporormiella decline by ~2 kyr (Halligan et al., 2016), consistent with Sporormiella disappearance at ~12.7 ka at another Florida panhandle location (Perrotti, 2018). Paucity of evidence for human occupation prior to ~13 ka can be attributed to habitation being concentrated in coastal regions that were largely submerged during the subsequent (last) deglaciation, when sea level rise necessitated migration of humans and other terrestrial mammals into the continental interior. We speculate that large mammals, particularly those that did not burrow, were particularly susceptible to DNA damage associated with low geomagnetic field strength and the resulting increase in UVB reaching the Earth's surface. The coincidence of low magnetic field strength with LQE events, particularly at ~40 and ~13 ka, but also at 107 ka and 122 ka in Australia, implies that UVB flux was a contributing cause of the LQE in North America, Europe, and Australia.

The ~190-ka paleointensity minimum at the Iceland Basin excursion corresponds to TMRCAs determined from the mtDNA and Y chromosomes of modern humans (Gonder et al., 2007; Oppenheimer, 2009, 2012; Poznik et al., 2016; Soares et al., 2009; Wei et al., 2013). Other paleointensity minima at 100-120 ka, 64 ka, 41 ka, and 13 ka (Figure 1) correspond to branches in phylogeny estimated from mtDNA and Y-chromosome analyses (Figure 4). For example, Wei et al. (2013) used five models to assess phylogenetic nodes from the TMRCA of 29 Y chromosomes that yielded branching ages for one model (GENETREE-2) of 112 \pm 12 ka, 68 \pm 7 ka, 49 \pm 6 ka, and 13 \pm 2 ka, corresponding closely to minima in RPI records (Figures 1 and 4). Although the transition from Middle Paleolithic to Upper Paleolithic at ~40 ka did not correspond to significant changes in AMH anatomy in the fossil record, it is an important time for human population structure (Figure 4; Lahr, 2016) with mtDNA implying rapid population growth at a time of major advance in tool design and the first appearance of artwork (Stoneking, 1994).

Phylogeography, the study of human dispersal, demonstrates that when migration occurs from one region to another, new mutations unique to that region accumulate (Soares et al., 2016). Local adaption to different habitats, including changes in exposure to mutagenic solar radiation, partially controlled by the magnetic field, are potential sources of phenotypical divergence (Jablonski & Chaplin, 2000, 2010). During

migrations, humans have adapted to differences in climate, altitude, and resource availability. Migration to new environments alters selection pressures on the human genome, and genetic studies have identified certain loci that were likely targets of selection for adaption to UVR (Ding et al., 2014). Highly pigmented skin protects against skin cancer but reduces synthesis of vitamin D3, so differences in the amount of UVR place differing selection pressures on pigmentation genes (Creanza & Feldman, 2016).

9. Conclusions and Outlook

Although coincidence does not prove causality, the timing of geomagnetic field strength minima (hence enhanced UVR flux at Earth's surface) appears to correspond to events in mammalian evolution. Improvements in knowledge of past geomagnetic field strength, new mammalian fossil finds, advances in radiocarbon dating and DNA analyses of fossils, use of dung-fungal proxies for herbivore populations, and advances in the use of mtDNA and Y chromosomes to map human phylogeny have all contributed to this possible linkage. Minima in geomagnetic dipole field strength at ~13 and ~41 ka (Laschamp; Figure 1) led to stratospheric ozone depletion and UVB levels at the Earth's surface that may have contributed to extinction of large mammals at these times, although extinction dates are associated with large errors due to the inadequacy of the fossil record (Figure 2). The Neanderthals were apparently victims of the ~41 ka (Laschamp) magnetic field minimum, and differences in the AhR of modern humans and Neanderthals may explain why we passed the Laschamp evolutionary barrier. Episodes of low field strength further back in time notably at ~64 ka, ~100-120 ka (Blake; Figure 1), and ~190 ka (Iceland Basin; Figure 1) may have contributed to phylogenetic nodes in hominin evolution revealed by fossil finds and by studies of mtDNA and Y-chromosomes (Figure 4). According to this hypothesis, the geomagnetic field influenced evolution of large long-lived mammals through exposure to UVR at times of low field strength, with foci of the extinction (e.g., Australia and Europe at ~41 ka, and North America and Europe at ~13 ka) depending on the specific geometry of stratospheric ozone depletion during episodes of low field strength. Ozone holes are preferentially located at high latitudes because of the role of stratospheric temperatures and polar stratospheric clouds in ozone depletion. UVR arriving at the Earth's surface may have had an influence on evolution due its strong mutagenic effects, its potential for promoting oxidative damage on membranes and proteins, and the role of AhR in translocating UVB stress response to the nucleus. Lower levels of UVR reaching Earth's surface at low latitudes, due to the role of polar-stratospheric clouds and stratospheric temperature in ozone depletion, may partly explain the relative diversity of modern African megafauna.

As the chronologies of both the RPI record and the paleontological record become better resolved, we speculate that unconvincing efforts to establish a linkage between polarity reversal and extinction over the last ~50 years will, in future, be superseded by a linkage between extinction and paleointensity minima particularly for land-dwelling mammals at higher latitudes where the magnetic field strength is linked to loss of stratospheric ozone and enhanced UVR reaching Earth's surface. The importance of Australian extinction events at the time of Laschamp excursion (41 ka) and the Blake excursions (100 and 120 ka) may imply that stratospheric ozone depletion was largely in the Southern Hemisphere at these times, and in the Northern Hemisphere when the North American extinctions are manifested at ~13 ka.

The magnetic dipole moment has decreased by $\sim 10\%$ since 1833 (Gauss' first direct field intensity measurement) or $\sim 5\%$ per century. Although the present dipole field strength may not be appreciably lower than the average during the Brunhes Chron, the current field intensity decrease, combined with asymmetry in the field detected in satellite data, has led to speculations that the geomagnetic field may reach intensity levels appropriate for a magnetic excursion or polarity reversal in the next 1,000-2,000 years (Glassmeier et al., 2009; Hulot et al., 2002; Laj & Kissel, 2015).

Twenty years ago, no more than five Quaternary magnetic excursions had been recorded (see Opdyke & Channell, 1996), whereas today ~20 Quaternary magnetic excursions have credible documentation (Laj & Channell, 2015). In the next few decades, we speculate that integration of excursion records with RPI and cosmogenic isotope records, documentation of mammalian extinction, understanding the role of geomagnetic field in UVR flux, and the role of Ahr in modulating the deleterious effects of UVR in mammals, as well as improvements in human phylogeny from mtDNA and Y chromosomes, will clarify the role of the geomagnetic field in mammalian evolution.

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Glossary

6-4 photoproducts

(6-4 PPs)

6-4 pyrimidine-pyrimidone photoproducts (6-4 PPs) and cyclobutane pyrimidine dimers (CPDs, see below) are the common ultraviolet radiation (UVR) products formed in the human skin during exposure to sunlight. Although the 6-4 PPs are not as prevalent as CPDs (in lesion formation), they are significantly more mutagenic.

Anhysteretic remanent magnetization

(ARM)

A laboratory-induced remanent magnetization imparted by a strong alternating magnetic field in the presence of a weak direct current bias field. The bias field usually has an intensity comparable to the geomagnetic field. ARM provides an indication of magnetic mineral concentrations, and the efficiency of ARM acquisition varies with magnetic grain size.

Apoptosis

A form of programmed cell death that occurs in multicellular organisms. It is a highly regulated and controlled process that confers advantages during an organism's lifecycle. Because apoptosis cannot be halted once initiated, it is a highly regulated process.

Artiodactyla

Order of even-toed ungulates, hoofed animals that bear their weight on two toes including hippopotamuses, camels, antelopes, sheep, giraffes, and deer.

Aryl hydrocarbon receptor (AhR)

A protein that regulates gene expression, cell physiology, and organ homeostasis with a key role in skin integrity and immunity. Its functions are related to cell proliferation, adhesion, and migration as well as to cell differentiation. It has a strong function in mediating the response to xenobiotic and toxins particularly Dioxins, and to UVR.

Brunhes Chron

The most recent time interval of the geomagnetic polarity timescale, characterized by a normal polarity of the geomagnetic field that has existed since ~773 ka.

Campanian Ignimbrite (CI)

The CI eruption was located near Napoli (Italy) and was probably the largest volcanic eruption in the Mediterranean region during the last 200 kyr. CI ash was deposited on the Russian Plain, and throughout the Eastern Mediterranean and northern Africa. The event coincided with the onset of a cold climatic phase known as Heinrich Stadial 4 (HS4) approximately 40,000 years ago.

Chlorofluorocarbons

(CFCs)

An organic compound (commonly known as Freon) that contains carbon, chlorine, and fluorine, produced as a volatile derivative of methane and ethane. Many CFCs have been commonly used as refrigerants, propellants (in aerosol applications), and solvents. Although the concentration of CFCs in the atmosphere is very small, measured in parts per trillion, they do contribute significantly to the enhancement of the greenhouse effect. CFCs contribute significantly to ozone depletion in the upper atmosphere. The manufacture of such compounds has been phased out under the Montreal Protocol, and their use is being replaced by other products such as hydrofluorocarbons.

Clade

In taxonomy, a clade is defined as a group of organisms consisting of a single common ancestor and all descendants of that ancestor.

Clovis horizon

The short-lived Clovis horizon comprises distinctive stone tools marking early human occupation of North America at ~13 ka. The Clovis culture is a prehistoric Paleo-Indian culture, named after characteristic stone tools found in close association with Late Pleistocene fauna at Blackwater near Clovis, New Mexico, in the 1920s and 1930s. Most of the indigenous cultures of the Americas are considered to descend from the Clovis people.

Coercivity

CHANNELL AND VIGLIOTTI

The coercivity, also called magnetic coercivity, is a measure of the ability of a remanent magnetization to withstand an external demagnetizing magnetic field. For ferromagnetic materials (*sensu lato*), the coercivity (or coercive force) is the intensity of the applied magnetic field required to reduce the remanent magnetization of that material to zero after the magnetization of the sample has been driven to saturation in the opposite direction. Thus, coercivity measures the resistance of a ferromagnetic material (*sensu lato*) to demagnetization by an external field.

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Cosmogenic isotopes

Radioactive isotopes created in the upper atmosphere when galactic cosmic rays collide with atmospheric molecules at high speed. The production rate of these isotopes depends on the intensity of the cosmic radiation, which is related to the strength of the Earth's magnetic field and solar activity. Therefore, records of cosmogenic isotope production are useful as a proxy for the strength of Earth's magnetic field and for understanding the relation between the Earth magnetic field, and variations in solar activity.

Cyclobutane pyrimidine dimers (CPDs) One of two major photoproducts generated by UV irradiation. CPDs are molecular lesions formed in the human skin during exposure to sunlight due to UV radiation and photochemical reactions.

Cytosol

It is the fluid inside living cells. Proteins, organelles, and other structures of cells live in this water-based fluid.

Deoxyribosyl

A univalent radical derived from deoxyribose, a monosaccharide (simple sugar). It is derived from the sugar as indicated by its name: deoxy sugar. Deoxyribose is the five-carbon sugar molecule involved to form the phosphate backbone of DNA molecules. DNA, or deoxyribonucleic acid, is a polymer formed of several nucleic acids. Each nucleic acid is composed by a molecule of deoxyribose bound to a phosphate group and either a purine or a pyrimidine. As part of DNA, 2-deoxyribose derivatives have a significant role in biology. The DNA molecule, which is the main repository of genetic information in life, consists of a long chain of deoxyribose-containing units called nucleotides, linked via phosphate groups.

DNA photoproducts

Exposure to the ultraviolet component of sunlight causes DNA damage, which subsequently leads to mutations, cellular transformation, and cell death through the creation of different photoproducts. There are two important DNA photoproducts, namely CPDs and pyrimidine-pyrimidone photoproducts (6-4 PPs). A ligand is a protein that attaches (binds) to another protein called a receptor; the latter have specific sites into which the ligands fit like keys into locks. Ligands that are produced in the body are called endogenous.

Endogenous ligand

Energetic particle precipitation (EPP)

Precipitation of a group of highly energetic electrons, protons, neutrons, and ions that are accelerated into the atmosphere through various heliophysical and geomagnetic processes. They enter the atmosphere mainly in the vicinity of the geomagnetic poles. When energetic particles enter the atmosphere they ionize and dissociate atmospheric constituents, resulting in the formation of reactive nitrogen oxides. EPPs have been shown to contribute up to 10% of the stratospheric NO_x budget and up to 40% of the polar stratospheric NO_x budget. Once in the stratosphere, NO_x produced by EPPs (EPP-NO_x) interferes with catalytic cycles involving ozone (O_3). Theoretically, changes in O_3 can also lead to changes in temperature and winds, which means that EPPs have the potential to impact climate.

Gaussian-resampled, inverse-weighted McInerny et al. (GRIWM)

Dating method for estimating the probability of extinction by using an approach that weighs observations inversely according to their temporal distance from the last observation of a species' confirmed occurrence. For dates with associated radiometric errors it is able to sample individual dates from an underlying fossilization probability distribution.

Geomagnetic excursions

Excursions represent a significant, but short-lived, change of direction of the Earth's magnetic field and are apparently manifested globally. Intervals of constant polarity are punctuated by geomagnetic excursions where magnetic directions depart from the usual geocentric axial dipole, and when adequately recorded, achieve the opposite polarity direction for a short time. Excursions have been recorded in both volcanic and sedimentary records. These directional aberrations typically have durations of a few thousand years, or less than 1 kyr in some cases, and are characterized by a decrease in strength of the main axial dipole.

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Halons

A group of hydrocarbon compounds in which some hydrogen atoms are replaced by bromine and fluorine atoms, and sometimes also chlorine. Halons are nonconductors of electricity and may be utilized in fighting fires in ignitable liquids and most solid flammable materials. Halons in the atmosphere are responsible for ozone depletion and contribute to greenhouse warming.

Haplogroup

Group of individuals sharing a common ancestor identified by DNA sequences defined by shared mutations and which tend to show regional specificity. In human genetics, the most commonly studied haplogroups are Y chromosome (Y-DNA) haplogroups and mitochondrial DNA (mtDNA) haplogroups, each of which can be used to describe genetic populations.

Haplotype

A group of genes in the chromosome of an organism that are inherited together, because of genetic linkage from a single parent. The word "haplotype" is derived from the word "haploid" that describes cells with only one set of chromosomes and from the word "genotype," which refers to the genetic makeup of an organism.

Heinrich Stadial

(HS) 4

A Heinrich event is a natural phenomenon. Several occured during a period known as Marine Isotope Stage 3 (about 59-27 ka). Large icebergs containing rock mass eroded by the glaciers broke off from glaciers and traversed the North Atlantic, as they melted this material was deposited on the sea floor as ice rafted debris. One of the stronger Heinrich stadials is HS4 at ~40 ka.

Isothermal remanent magnetization (IRM)

IRM is an artificial (usually laboratory acquired) remanent magnetization acquired by applying a strong direct field to a ferromagnetic (*sensu lato*) material. It is a useful parameter used for detection of magnetic minerals with high coercivity such as hematite or goethite.

Late Quaternary extinction (LQE)

Widespread extinction of large terrestrial mammals (megafauna) during the Late Quaternary, observed in the Americas, Europe, Africa, Asia, and Australia. As the ages of extinction have become better constrained, the timing of these extinctions has become focused in the vicinity of 40 and 13 ka.

Ligand binding

Intermolecular interactions occurring between or among proteins, nucleic acids, or small molecules. The interaction of ligands with their binding sites is often characterized in terms of a binding affinity. In general, high-affinity ligand binding occurs from greater intermolecular force between the ligand and its receptor, whereas low-affinity ligand binding involves less intermolecular force between the ligand and its receptor.

Luminescence age dating

Refers to a group of dating methods based on the determination of how long ago mineral grains were last exposed to sunlight. The techniques include optically stimulated luminescence, infrared stimulated luminescence, and thermoluminescence dating. It is useful to geologists and archaeologists who want to know the age of a particular event. Sediments and soils contain traces of radioactive isotopes such as potassium, uranium, thorium, and rubidium. The ionizing radiation emitted during the decay of these elements is absorbed and trapped by quartz and potassium feldspar grains occurring in the sediment. Stimulating these mineral grains using either light (blue or green for optically stimulated luminescence; infrared for infrared stimulated luminescence) or heat (for thermoluminescence dating) produces a luminescence signal emitted as stored unstable electron energy, the intensity of which varies depending on the

Mediterranean sapropel S7

(for thermoluminescence dating) produces a luminescence signal emitted as stored unstable electron energy, the intensity of which varies depending on the amount of radiation absorbed during burial, and specific properties of the mineral. The sediments of the Mediterranean (central-eastern) are characterized by the presence of organic-rich (>2% organic carbon) layers called sapropels. They were deposited during periods of reduced oxygenation related to increasing riverine input occurring in correspondence with peaks in solar insolation. Each peak has been numbered starting from the youngest (S1; 8-10 ka) and the sapropel S7 occurred around 190 ka.

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Melanocyte homeos-

tasis

Mitochondria

The maintenance of systems within a cell as conditions change is called homeostatic regulation, and melanocyte (a mature melanin-forming skin cell) homeostasis is a paradigm for understanding the formation of melanoma. Mitochondria are part of a eukaryote cell found in the cytoplasm. They oxidise glucose to provide energy for the cell through the creation of a molecule, called adenosine triphosphate that cells use as an energy source.

Mitochondrial DNA (mtDNA)

Mitochondria are tiny organelles that live in the cytoplasm of cells, called mitochondria. Each cell contains thousands of mitochondria with its own small circle of DNA, a reminder of their distant bacterial ancestry. MtDNA is not located in the cell nucleus, and in most species, including humans, it is inherited solely from the mother and is not subject to genetic recombination during meiosis. Therefore, it remains unchanged from generation to generation. Human mtDNA was the first significant part of the human genome to be sequenced. Since animal mtDNA evolves faster than nuclear genetic markers, it represents a pillar of phylogenetics and evolutionary biology. It also makes possible an examination of the affinity of populations, and so has become important in anthropology and biogeography.

Mitochondrial Eve

Analyses of the mitochondrial DNA of living humans have shown that we descend from a common female ancestor that has been dated to about 200 ka. This woman has been named "Eve" or "Mitochondrial Eve" after a journalist's confusing reference to the unrelated biblical story of the first woman being created by God from Adam's rib. ("The man called his wife Eve ["life"], because she was the mother of all who live." Genesis 3:20)

Mitohormesis

Mitohormesis, known also as mitochondrial hormesis, is a particular form of hormesis that is a nonlinear response to potentially harmful substances. Mitochondrial function often results in excessive production of reactive oxygen species (ROS) that are responsible for many chronic diseases. However, moderate levels of mitochondrial ROS can protect against chronic disease by stimulating mitochondrial capacity and endogenous antioxidant defense. This phenomenon is called mitohormesis.

Mousterian tool industry

The predominant industry of the Middle Paleolithic is termed Mousterian, named for its type location at Le Moustier, in Dordogne, France. Mousterian industry is the tool culture traditionally associated with Neanderthal man in Europe, western Asia, and northern Africa. Mousterian tools disappeared abruptly from Europe with the passing of Neanderthal man.

Natural remanent magnetization

(NRM)

The magnetic remanence carried by rocks or sediments before the laboratory demagnetization or magnetization treatments. The NRM of a rock or sediment is usually represented by multiple magnetization components acquired through its geologic history. Thermal or alternating field laboratory treatments are usually necessary to identify the primary remanence acquired at the time of rock formation or sediment deposition.

Pedigree-based substitution rate

A method to estimate mutation rate by comparing the mtDNA sequences of a sample of parent/offspring pairs or analyzing mtDNA sequences of individuals from a deep-rooted (well established) genealogy. The number of new mutations in the sample is counted and divided by the total number of parent-to-child DNA transmission events to calculate a mutation rate.

Perissodactyla

Order of odd-toed ungulates, hoofed animals that bear most of their weight on a single toe including horses, zebras, rhinoceroses, and tapirs.

Peto's paradox

Peto's paradox, named after Richard Peto, is the observation that at species level, the incidence of cancer does not appear to correlate with the number of cells in an organism (e.g., body weight). Although cell mutations are usually deleterious, the incidence of cancer is not apparently related to the body-size of individual mammals.

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Phenotypical

The observable physical or biochemical characteristics of an organism determined by genetic and environmental influences.

Phylogeny

The evolutionary history and relationships among people or groups of organisms (e.g., species or populations). These relationships can be discovered through phylogenetic methods that evaluate observed heritable traits, such as DNA sequences or morphology, using an evolutionary model for these traits. The results of these analyses represent the phylogeny (also referred to as the phylogenetic tree).

Polar-stratospheric clouds

Polar-stratospheric clouds are typically formed in the stratosphere (15- to 25-km altitude) at high latitudes especially in winter because they require very low temperatures. They are implicated in the formation of ozone holes because they convert benign forms of chlorine into reactive, ozone-destroying forms, and they remove gaseous nitric acid that would otherwise moderate the destructive impact of chlorine.

Quaternary

The most recent of the three periods of the Cenozoic Era in the geologic time scale. It spans the last 2.588 Myr and is divided into two epochs: the Pleistocene (2.588 million years ago to 11.7 thousand years ago) and the Holocene (11.7 thousand years ago to present).

Reactive oxygen species (ROS)

Oxygen-derived molecules that act as powerful oxidants and may form via a large number of physiologic and nonphysiologic processes as a result of natural consequences of aerobic metabolism. Although these molecules play a role in the oxygen-dependent defense mechanism against bacteria, they may also be highly damaging, as they can attack biologic macromolecules, including lipids, proteins, and DNA, and lead to significant tissue damage. In normal conditions, ROS can act as immune system modulation and can activate various signal transduction pathways. ROS levels can increase dramatically during times of environmental stress (e.g., UVR or heat exposure).

Relative paleointensity (RPI)

The intensity of the Earth's axial dipole field is globally coherent and can be reconstructed in the past by studying suitable rock and sediment samples or archaeological materials. The RPI proxy in sediments usually comprises the NRM intensity normalized by the intensity of a laboratory-acquired magnetization designed to activate the same population of magnetic grains that carry the NRM, thus compensating for variations in concentration of NRM-carrying grains through the sedimentary section. The laboratory-applied normalizers are typically ARM and/or IRM.

Signor-Lipps effect

The first and last occurrence of a taxon will never be accurately recorded in the fossil record because of the inherent incompleteness of the record.

Sclerophyll vegetation

Type of vegetation characterized by hard, leathery, evergreen foliage well adapted to prevent moisture loss. Broad-leaved sclerophyll vegetation, including species such as holly (Ilex), is known as Mediterranean vegetation because it develops in regions with a Mediterranean climate—hot, dry summers and mild, wet winters. Pines are examples of narrow-leaved sclerophyll vegetation.

Solar proton events (SPE)

A SPE or "proton storm" occurs when particles (mostly protons) emitted by the Sun are accelerated either close to the Sun during a flare or in interplanetary space by coronal mass ejection. Solar protons normally have insufficient energy to penetrate the Earth's protective magnetosphere. However, during unusually strong flares, protons can be produced with sufficient energies to penetrate the Earth's magnetosphere and ionosphere around the poles. They can pass through the Earth's magnetic field and cause ionization in the ionosphere. The effect is analogous to an auroral event, except that protons instead of electrons are involved. Energetic protons represent a significant radiation hazard to spacecraft and astronauts.

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The time to the most recent common ancestor (TMRCA) TP53 (p53) gene Time to the most recent individual from which an entire population of the group is directly descended. The term is used in reference to the ancestry of groups of genes (haplotypes) or species.

Tumor protein 53 is a protein-coding gene that acts as a tumor suppressor for many tumor types and induces growth arrest or apoptosis depending on physiological circumstances and cell type. The TP53 gene is responsible for instructions to create a protein called tumor protein p53. This protein acts as a tumor suppressor, which means that it regulates cell division by keeping cells from growing and dividing (proliferating) too fast or in an uncontrolled way. The p53 protein is found in the nucleus of cells throughout the body, where it attaches (binds) directly to DNA. If the DNA in a cell is damaged by agents such as toxic chemicals, radiation, or UVR, this protein plays a critical role as to whether the DNA can be repaired or the damaged cell will self-destruct (undergo apoptosis). If it is possible to repair the DNA, p53 activates other genes to fix the damage. On the other hand, this protein can prevent the cell from dividing and signals it to undergo apoptosis. By stopping cells with mutated or damaged DNA from dividing, p53 helps prevent tumor development.

Tryptophan

Tryptophan is a α -amino acid that is used in the biosynthesis of proteins. Many animals (including humans) cannot synthesize tryptophan: they obtain it through their diet. Tryptophan is among the less common amino acids found in proteins, but it has critical structural and functional roles.

Virtual axial dipole moment (VADM)

Intensity of an imaginary axial (along the Earth's rotation axis) centric (located in the center of the Earth) dipole that would produce the estimated magnetic field reconstructed from paleomagnetic data. The reconstruction is virtual because the VADM is an approximation of a geocentric axial dipole moment.

Xenobiotics

A xenobiotic is a chemical substance within an organism that is foreign to the biological system and not naturally produced or expected to be present within the organism. It can also cover substances that are present in much higher concentrations than usual. The term is often used in the context of pollutants and their effect on biota.

Y chromosome

One of the two sex chromosomes in humans (the other is the X chromosome) and most mammals. The Y chromosome occurs in males, who have one X and one Y chromosome, while females have two X chromosomes. The Y chromosome contains genes that provide instructions for making proteins. Because only males have the Y chromosome, the genes in this chromosome are involved in male sex determination and development. The Y chromosome represents almost 2% of the total DNA in male cells.

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Erratum

In the originally published version of this article, discoloration appeared in Figure 2 and Figure 4 was not large enough. Figures 2 and 4 have been corrected, and this version may be considered the authoritative version of record.