Supplementary Material For:

Comparison and optimization of ancient DNA extraction

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Supplementary Materials and Methods

1 Initial comparison

1a) Initial comparison of different methods using ~40mg sample material

1a.1 Silica method (1)

40 mg of powder were digested in 1ml buffer composed of 0.45M EDTA pH 8.0, 0.5% N-lauroylsarcosine, 1% PVP, 50mM DTT, 2.5mM PTB and 0.25 mg/ml proteinase K overnight at 37°C in the dark. After centrifugation, the supernatant was added to 4ml binding buffer (L2: 5M GuSCN, 25mM NaCl, 50mM Tris (pH 8.0)) plus 15μl silica suspension (prepared according to (2), using silica from Sigma [ordering number: S5631]) and incubated under agitation for one hour at room temperature. The silica pellet was collected via centrifugation, the supernatant discarded and the pellet washed once with binding buffer and once with NewWash solution (see 3.8 for preparation, Qbiogene, USA). Subsequently, the pellet was dried at 56°C for 7min. DNA was eluted with 50μl 1xTE at 56°C for 10min, and after centrifugation, the aqueous solution was transferred to a new tube.

1a.2 Centricons

40 mg of powder were incubated as described above. After centrifugation, the supernatant was added to 1ml phenol (pH 8.0), gently mixed and centrifuged for 3min. This step was repeated once. The aqueous phase was added to 1ml Ready Red (Chloroform / Isoamylalcohol, Qbiogene), mixed and centrifuged again for 3min. Thereafter, the aqueous phase was added to Centricon YM 30 (MWCO 30,000; Millipore, USA) filter devices and centrifuged at 13,200 rpm until ~ 50μl remained. This solution was washed twice with 100μl 1xTE by centrifugation. The remaining ~50μl extract were transferred using inverse centrifugation of the filter into a new tube.

1a.3 Phenol chloroform (3)

40 mg of powder were decalcified in 2ml 0.5M EDTA (pH 8.0) at room temperature overnight. After centrifugation, the supernatant was discarded and the remaining pellet was digested by incubation at 37°C overnight in 1ml extraction buffer composed of 10mM Tris (pH 8.0), 10mM NaCl, 0.1% SDS, 50mM DTT, 0.25mg/ml proteinase K and 2.5mM PTB (concentrations of DTT, proteinase K and PTB were taken from (1), as the concentrations are omitted in (3)). After centrifugation, the supernatant was added to 1ml phenol (pH 8.0), gently mixed and centrifuged for 3min. This step was then repeated. The aqueous phase was added to 1ml Ready Red, gently mixed and centrifuged for 3min. Then the aqueous phase was added to Centricon YM 30 filter devices, and the solution was concentrated via centrifugation at 5,000 rpm. The solution was washed twice with 200μl 1xTE. The remaining ~50μl extract were transferred by inverse centrifugation of the filter using a new tube.

1a.4 User-developed protocol: Isolation of genomic DNA from compact bone using the DNeasy Tissue Kit (Qiagen, Germany)

40 mg of powder were decalcified in 2ml 0.5M EDTA (pH 8.0) at room temperature overnight. After centrifugation, the supernatant was discarded and the pellet washed with 1ml ddH₂O, centrifuged and again the supernatant discarded. 360μl buffer ATL and 40μl proteinase K were added to the pellet. This mixture was incubated under agitation for one hour at 56°C until lysis of the pellet was complete. After vortexing, 400μl buffer AL were added, and the mixture was incubated at 70°C for 10min. 400μl ethanol absolute and 3ng of DNA size marker serving as a carrier were added, and the samples were gently mixed. The mixture was pipetted onto DNeasy spin columns and centrifuged at 8,000 rpm for 1min. The flow-through was discarded, and the remaining mixture was pipetted to the columns and centrifuged again. Following this step, the column was washed once with 500μl buffer AWl, once with 500 μl AW2, and then centrifuged for 3min at 14,000rpm until it was dry. 50μl of buffer AE were added, incubated for 1min at room temperature and centrifuged for 1min at 8,000rpm. This flow-through represents the extract.

1a.5 "DNA-Präparation aus Knochen" with All-tissue DNA-Kit (GEN-IAL GmbH,

Germany)

40 mg of bone powder were decalcified overnight in 2ml 0.5M EDTA (pH 8.0) at room temperature. After centrifugation, the supernatant was discarded and the pellet was washed with 1ml ddH₂O, centrifuged and the supernatant again discarded. 1ml Lyse1 buffer, 100µl Lyse 2, 3mg DTT and 15µl proteinase K were added to the pellet and

incubated for about one hour at 56°C under agitation until the pellet was completely dissolved. 750µl Lyse 3 buffer were added, vortexed, incubated at -20°C for 5min and centrifuged at 13,000rpm for 20min. The supernatant (~1.8ml) was transferred to a fresh tube and 0.8 volume of isopropanol, plus 10ng of a DNA size marker as a carrier were added. To precipitate the DNA, the mixture was centrifuged for 15min at 13,000rpm. The supernatant was removed and 300µl ice cold 70% ethanol were added and centrifuged at 13,000rpm for 5min. The liquid was discarded and the DNA pellet dried at 56°C for 5min. The DNA was dissolved in 50µl 1xTE.

1a.6 Bone Extraction Protocol to be used with the DNA IQ System (Promega, USA).
40 mg of powder were incubated in 1ml proteinase K digestion buffer composed of
10mM Tris (pH 8.0), 10mM NaCl, 50mM EDTA (pH 8.0), 0.5% SDS and 1mg/ml
proteinase K for one hour at 56°C under agitation. 2ml Lysis buffer (including 10mM
DTT) and 15μl of Resin (magnetic particles) were added to the supernatant. After
incubation at room temperature for 10min with frequent mixing of the solution, the Resin
was collected using the Magnetic Stand. The supernatant was removed and again 100μl
Lysis buffer were added, vortexed, the Resin separated, the solution removed and 100μl
Wash buffer added. After vortexing, separation and removal of the liquid, this washing
step was repeated twice. The Resin was then air dried for 15min in the Magnetic Stand.
50μl 1xTE were added, vortexed and DNA was eluted for 5min at 65°C. The separated
supernatant represents the extract.

1b) Initial comparison of "silica" with "DNA IQ" using 200mg sample material

1b.1 Silica method (1)

The method is the same as described above (1a.1), except that the amount of powder was increased to 200mg, the volume of extraction buffer increased to 5ml, the incubation conditions were modified from 37°C overnight to 56°C for one hour (the same as for the DNA IQ Kit) and the volumes of the binding buffer and the silica suspension were increased to 20ml and 50µl, respectively. All other steps including concentrations, times and temperatures remained identical.

1b.2 Bone Extraction Protocol to be used with the DNA IQ System (Promega, USA). The method is the same as described above (1a.6), except that the amount of powder was increased to 200mg and the volumes of proteinase K digestion buffer and of the Lysis buffer were increased to 1.5ml and 3ml, respectively. All other steps including concentrations, times and temperatures remained identical.

2 Extraction buffer components, incubation temperature and time.

2.1 Surfactants

The "silica" method was used as described under 1b.1, except that the temperature of the initial incubation was again set to 37°C, the amount of silica suspension was 40µl and aside from the 0.5% N-lauroylsarcosine (sarcosyl) used in the original protocol, the performance of other detergents (non-ionic: 1% Tween-20, 1% Triton X-100; anionic: 0.5% SDS; cationic: 1% CTAB, 2M Urea, 2M GuCl) as well as the absence of detergent was tested.

2.2 Incubation temperature and time

Extraction was performed as described under 2.1, without adding detergents to the extraction buffer. Incubation temperature and time of the powder in extraction buffer were varied (56°C for 1hour; 37°C for 1hour; 37°C overnight; room temperature overnight; room temperature for four days, room temperature for four days plus the addition of 0.25mg/ml proteinase K after two days).

2.3 Remaining chemicals in the extraction buffer in all combinations

Extraction was performed as described under 2.2. Incubation was at room temperature overnight and the effects of both the presence and absence of all remaining ingredients in the extraction buffer were tested in all possible combinations. Results were compared for each possible extraction buffer composition in pairwise comparisons of presence versus absence of the four ingredients except EDTA (PTB, PVP, DTT and PK).

2.4 Presence vs. absence of 0.05M Tris (pH 8.0)

Extraction was performed as described under 2.3, reducing the composition of the extraction buffer to 0.45M EDTA, 0.25mg/ml proteinase K and 50mM DTT and testing the presence and absence of 0.05M Tris (pH 8.0) in the extraction buffer.

2.5 EDTA concentration

Extraction was performed as described under 2.4 without adding Tris, varying the EDTA concentration between 0.45M, 0.2M, 0.1M and 0.05M in the extraction buffer.

2.6 Reducing agents

Extraction was performed as described under 2.5 with an EDTA concentration of 0.45M in the buffer. We tested the effect of different reducing agents besides DTT, all in the same concentration of 50mM (Thioglycolic acid; 2-Mercaptobenzothiazole; 2-Naphthalenethiol; 2-Amino-5-mercapto-1,3,4-thiadiazole; 5,5 \subseteq-Dithiobis(2-nitrobenzoic acid); 2-Mercaptoethanol; Dodecanthiol), as well as the absence of any reducing agent.

2.7 Sample preparation technique

Extraction was performed as described under 2.6 without any reducing agent. The difference in this experiment was how the sample was prepared prior to incubation in the extraction buffer. Either the powder was prepared using a freezer mill or part of the compact sample was ground to a more or less coarse-grained "powder" using a mortar and pestle. It should be noted that the different samples could be ground to varying

granularities using the mortar and pestle. For example, sample 891 is extremely compact, and particles with varying diameters of up to several millimeters were used in this experiment, whereas other samples were much easier to grind using mortar and pestle, so that a fine powder was obtained. After completing this experiment, the remaining sample material was extracted a second time using the same procedure as described above, except that it was incubated for 2 hours at 56°C until almost all powder was digested.

3 Binding buffer, silica age, pH during binding, binding time, silica preparation technique, wash solutions

3.1 Age of the silica suspension

Extraction was performed as described under 2.6, again using a fine, homogeneous powder. In this experiment the age of the silica suspension varied from one week to approximately one year.

3.2 pH of the binding buffer (including the extraction buffer plus silica)

Extraction was performed as described under 3.1 using silica suspension not older than three months, and adjusting the pH between 4.0, 6.0 and 8.0 with hydrochloric acid after adding both silica suspension and extraction buffer to the binding buffer.

3.3 Amount of silica suspension

Extraction was performed as described under 3.2 (pH was not adjusted to 4.0, but was ~ 8.0) using different amounts of silica suspension (5μl, 10μl, 20μl, 40μl, 100μl).

3.4 Different binding salts tested on DNA size marker

In this experiment, we used a 100bp DNA size marker (NEB, USA) instead of an extraction buffer containing ancient DNA. The size marker was diluted with 0.5M EDTA (total 400μl) and 8μl silica suspension plus 1.6ml of the respective salt solution (saturated salts: KH₂PO₄, NaCl, NaI, NH₄SO₄, CsCl, KCl, MgSo₄, GuCl, NaAc, NH₄Ac; 5M solution: NaClO₄, GuSCN) were added. The mixtures were incubated for 1 hour at room

temperature under constant agitation. After centrifugation the supernatant was discarded and the silica pellet was washed once with the respective binding salt and once with NewWash solution. After completely removing the liquid, the pellet was dried for 5min at 56°C and DNA was eluted in 15µl 1xTE. 10µl were loaded on a gel together with a size marker in the amount expected at 100% recovery.

3.5 Performance of different binding salts using ancient DNA samples

Extractions were performed as described under 3.2 adjusting the pH to 4.0 and using 100µl silica suspension. Instead of using the usual binding buffer, different salts (saturated KCl, saturated GuCl, 5M GuSCN, 5M NaClO₄, saturated NaCl, saturated NaI) were used as binding buffer, as these salts gave the best results in the size marker test (3.4). After binding to silica and centrifugation, the silica was washed first with the respective binding solution and then with NewWash buffer. When GuSCN was used as a binding salt, two washing variations were performed. In the first variation, the silica was washed once with NewWash and in the second, it was washed twice before it was dried and the DNA eluted.

3.6 Binding time

Extractions were performed as described under 3.5 using 3M NaCl. Washing steps were done once with 3M NaCl and twice with NewWash buffer. This experiment was done using only two samples from 100mg sample material each, because the remaining material from the third sample was needed for the final comparison. The low amounts of bone powder resulted in highly variable outcomes. The binding time was varied in the

first trial (a) between 15 minutes to overnight (0.25h, 1h, 3h, overnight) and in the second trial (b) between 1.5h, 3h and 6h.

3.7 Silica preparation technique

Extractions were performed as described under 3.6 using an incubation time of 3h for binding the DNA to silica. Silica suspensions were prepared using two different techniques. While the "normal" silica was prepared as described in Boom et al. (2), the second preparation technique ("new") was slightly different: 4.8g Silica were suspended in ddH_2O to a final volume of 40ml. After sedimentation for 1 hour, 39ml were transferred into a new tube and left to settle for additional 4 hours. Thereafter 35ml of the supernatant were removed and $48\mu l$ 30% HCl were added to the sediment in order to adjust the pH. After dividing the silica into 400 l aliquots, the solution was stored in the refrigerator.

3.8 Testing the efficacy of different wash solutions

Extractions were performed as described under 3.7 using the new silica preparation technique. Four different possible alternatives for the commercial NewWash solution (1ml NewWash concentrate [Qbiogene], 20ml ddH₂O, 23ml Ethanol absolute) were tested:

Alternative 1: 50% Ethanol, 0.1M NaCl, 1mM EDTA (pH 8.0), 10mM Tris (pH 8.0)

Alternative 2: 50% Ethanol, 0.3M NaCl, 1mM EDTA (pH 8.0), 10mM Tris (pH 8.0)

Alternative 3: 60% Ethanol, 0.1M NaCl, 1mM EDTA (pH 8.0), 10mM Tris (pH 8.0)

Alternative 4: 70% Ethanol, 0.3M NaCl, 10mM EDTA (pH 8.0), 100mM Tris (pH 8.0)

4 Final comparison

4.1 Silica method (1)—method A

Extractions were performed as described under 1a.1 using 500 mg powder and an extraction buffer volume of 10ml. The binding buffer volume was increased to 35ml, the silica suspension to 50µl and the DNA was eluted in 100µl 1xTE.

A 1:10 and 1:50 dilution were prepared immediately after the extraction, and a standard PCR (see 5.2) was performed with these two dilution steps before the extracts were frozen.

4.2 Modified silica method with NaCl as binding salt—method B

500 mg of powder were incubated in 10ml extraction buffer (0.45M EDTA and 0.25mg/ml proteinase K) at room temperature overnight. After centrifugation, the supernatant was added to 35ml 5M NaCl, 90μl silica suspension prepared using the new preparation technique was added and the pH was adjusted to 4.0. This mixture was incubated for three hours at room temperature under rotation. After centrifugation the supernatant was discarded, the silica pellet was washed with 1ml 5M NaCl and then transferred into a 2ml tube. After short centrifugation, the NaCl was removed with a pipette, the pellet resuspended in 1ml wash solution, which is a combination between alternatives 1 and 2 (51.3% Ethanol, 125mM NaCl, 1mM EDTA (pH 8.0), 10mM Tris (pH 8.0)), shortly centrifuged, and the supernatant removed with a pipette. The last washing step was repeated. Subsequently, the silica pellet was air dried at room temperature for 15min, resuspended in 100μl 1xTE and after 8min incubation at RT, the

silica was collected by centrifugation at maximum speed (16,000 g). The supernatant, representing the extract, was transferred into a fresh tube. 1:10 and 1:50 dilutions were prepared immediately after the extraction, and a standard PCR was performed using these two dilutions before the extracts were frozen.

4.3 Modified silica method using GuSCN binding solution—method C

This extraction method is exactly the same as described above (4.2), except that the 5M NaCl was exchanged by the binding solution used in the initial silica method (L2; see 1a.1). 1:10 and 1:50 dilutions were prepared immediately after extraction, and a standard PCR was performed using these dilutions before the extracts were frozen.

4.4 Phenol chloroform (3)—method D

Extractions were performed as described under 1a.3 using 500 mg bone powder and 25ml 0.5M EDTA for decalcification. The remaining protein pellet was digested in 10ml extraction buffer at 53°C. The volume of phenol and Ready Red was adjusted to 10ml each and the volume of washing with 1xTE was, in total, $650\mu l$. The remaining extract was $\sim 100\mu l$ as for all other methods. 1:10 and 1:50 dilutions were prepared immediately after extraction, and a standard PCR was performed using these two dilutions before the extracts were frozen.

4.5 Ethanol precipitation (4)—method E

500 mg of sample powder were digested in 1.1ml extraction buffer composed of 0.1M EDTA (pH 8.0), 0.5% N-lauroylsarcosine, 0.25mg/ml proteinase K at 37°C overnight.

After centrifugation, the approximately 750μl of supernatant were divided into 3 aliquots à 250μl (the remaining liquid was bound to the bone powder). To each 250μl aliquot, 250μl 4M NH₄-Acetat, 3.5μl Blue Dextran (1μg/μl) and 500μl 96% ethanol were added and this mixture vortexed. After incubation for 10min at -80°C, the DNA was precipitated at 13,200rpm for 15min, dried at 56°C for 10min and dissolved in 33.3μl 1xTE for each of the 3 parts, resulting in 100μl per sample. 1:10 and 1:50 dilutions were prepared immediately after extraction and a standard PCR was performed with these two dilutions before the extracts were frozen.

4.6 User-developed protocol: Isolation of genomic DNA from compact bone using the DNeasy Tissue Kit (Qiagen, Germany)—method F

These extractions were performed as described under 1a.4 using 500mg powder and 18ml 0.5M EDTA for decalcification. Washing the protein pellet with ddH₂O was omitted. 3.8ml buffer ATL and 200µl proteinase K (10mg/ml) were added to the pellet and incubated for about 2 hours at 55°C until the pellet was dissolved. The volume of both buffer AL and ethanol absolute was increased to 4ml and no carrier was added. This mixture was added step by step (each 650µl maximum) to the columns and centrifuged for 1min after each addition of buffer. The volume of the elution buffer AE was increased to 100µl. 1:10 and 1:50 dilutions were prepared immediately after extraction and a standard PCR was performed with these two dilutions before the extracts were frozen.

5 Quantitative PCR and standard PCR

5.1 Real time quantitative PCR

A probe based real time quantitative PCR assay was designed using sequences from *Ursus arctos* (AF303110), *U. maritimus* (NC003428) and *U. americanus* (Y08520) and the PrimerExpress software (Applied Biosystems). It is located in the 12S region of the mitochondrial genome and is 110bp long, including primers. As no information about cave bear 12S sequences were found in the database, the nucleotide sequences of both primers and probe were sequenced in four cave bear samples from three different caves, all showing the same sequence. The assay, furthermore, was designed not to amplify human contamination, as the forward primer contains two mismatches (5) at the 3'end and the probe has, in total, 7 mismatches to the corresponding human sequence. DNA from the above mentioned *Ursus* species should also not be detected, as all have a minimum of one mismatch to the probe. However, it was never tested if DNA from these species could be detected using this assay. Sequences of primer and probe were as follows:

CB TM 12S F 5'-AAA ATG CCC TTT GGA TCT TAA T-3'

CB TM 12S R 5'-ACT GCT GTA TCC CGT GGG-3'

CB TM 12S Probe 5'-VIC-CGA TTT GAA GGA GCG GGC A-TAMRA-3'

As the quantitative PCR assay is probe-based, potential nonspecific amplification products are not measured, as they would lack a sequence complementary to the probe. Thus, even if amplification of nonspecific products occurred, it would not influence the results. We used eleven dilution steps (between 25 and 50,000 copies) of a PCR product about 399bp in length amplified from a cave bear extract from Gamssulzen cave, Austria,

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as template DNA in the standard. Each dilution step was amplified in triplicate for the same 110 bp fragment as the ancient DNA extracts alongside the samples. The copy number of the standard was estimated via UV absorption using a spectrophotometer (Nanodrop ND-1000).

 $5\mu l$ of the extracts or dilutions of extracts were used for each quantitative PCR in a final volume of $20\mu l$ consisting of 1x TaqMan® Buffer A, 4.5mM MgCl₂, 0.4mM dUTP, 0.2mM dATP, dCTP and dGTP each, 1U / 2U AmpliTaq® Gold DNA Polymerase (1U was used for the "initial comparisons" and the "detergent" test, all subsequently performed real time quantitative PCR runs were performed using 2U), $0.3\mu M$ each forward and reverse primer and $0.2\mu M$ probe. Initial denaturation and activation of AmpliTaq® Gold were run for 5min at 95° C, followed by 50 to 60 cycles at 95° C for 15sec and 60° C for 45sec.

Standard curves were generated from the dilution series, and copy numbers of the extracts were extrapolated from the equation of the standard curve. R² of standard curves were between 0.96 and 0.99, with 73% of all runs having values above 0.98.

5.2 Standard PCR after final extraction

After finishing each final extraction protocol, a standard PCR using bear specific primers (CBL164 and CBH221 from (1)) was performed, amplifying a 103bp long fragment of the mitochondrial HVR I. 5μl of 1:10 and 1:50 dilutions were used in a final volume of 20μl consisting of 1x GeneAmp® PCR Buffer II , 4mM MgCl₂, 625μM of each dNTP, 2.5U AmpliTaq Gold® and 500nM of forward and reverse primer, respectively. Initial denaturation and activation of AmpliTaq Gold® were run at 94°C for 4min, followed by

60 cycles 94°C for 20sec, 52°C for 30sec, 72°C for 30sec and a final elongation at 72°C for 10min.

6 Inhibition test

To assess the degree of inhibition originating from the different extraction methods in the final comparison, a quantitative human mitochondrial PCR assay (6) was spiked with 5µl of a 1:10 dilution of the extracts. In a 20µl reaction the concentrations of the various ingredients were as follows: 1x TaqMan buffer A, 4.5 mM MgCl₂, 200 pM probe, 300 pM forward and reverse primer, 2U AmpliTaq Gold® and (except for the negative controls and the standard) 30 pg of human DNA. The standard dilution was done in quadruplicate from the same human DNA, beginning with 10,000 pg down to 1pg in 10 fold dilutions, and was amplified alongside the inhibition test. The quantitative PCR was performed on a Stratagene Mx3000p (Stratagene, The Netherlands), with 10min activation of AmpliTaq Gold® at 95°C, followed by 45 cycles at 95°C for 15sec and 60°C for 1min. In total, 23 positive controls (without spiking with ancient DNA) were performed.

Two different approaches for testing for inhibition were used. The first approach calculates the inhibition from the differences of the mean of all positive PCR controls (containing ~ 30pg human DNA), and the mean of the six measurements per sample for each extraction method using the absolute amount of DNA measured. If this difference deviated more than two times the standard deviation from the mean of the positive controls, we scored the extract as inhibitory. To understand the degree of inhibition, these differences were divided by the mean value of the positive controls. This procedure

produced the inhibition of each sample and method as percentages (i.e. 90% inhibition indicates that the copy number measured was only 10% of the mean copy number measured for the positive controls).

As it is assumed that inhibition influences the efficiency of the PCR, so that there is less than doubling of the target sequence per cycle, thereby rendering the slope of the qPCR curve to be less steep compared to uninhibited PCR, the second approach calculates the inhibition from the efficiencies of each individual qPCR. The efficiencies of each individual PCR were calculated using the software LinRegPCR (7). Next, we calculated the difference between the mean of the efficiencies from all positive controls (and the standards), and the mean of the six measurements per sample and extraction method. If the differences were larger than two standard deviations of the positive controls, inhibition was assumed. Again, to get an idea of the degree of inhibition, the differences were divided by the mean of the positive controls, and the result was treated as inhibition percentages for each sample and method.

7 Measuring PCR performance

7.1 Presence vs. absence of BSA

Five extracts from different cave bear samples were tested in triplicate with the same 103bp fragment mentioned above (5.2), in both the presence and absence of a final volume of 1mg/ml BSA (bovine serum albumin) per PCR reaction using different amounts of AmpliTaq Gold® (between 1U and 1/16U). 5μl of 1:10 dilution of extract were used in a final volume of 20μl using 1x GeneAmp® PCR Buffer II, 2.5mM MgCl₂, 200μM of each dNTP, 200nM of forward and reverse primer, respectively, both with and

without 1mg/ml BSA and varying concentrations of AmpliTaq Gold® (1U, 0.5U, 0.25U, 0.125U, 0.0625U) using the temperature profile described above (5.2).

7.2 Efficacy of different Taq DNA Polymerases

7.2.1 General consideration

If not stated otherwise (e.g. in case of pre-mixed solutions), in the following tests, the same dilution of one cave bear sample, 1U of the respective Taq DNA polymerase, 0.2µM of each primer and 200µM each dNTP were always used.

7.2.2 SUPER TAQ (HT-Biotechnology LTD, UK) + TaqStart™ Antibody (BD Bioscience, USA)

As this polymerase is not a hot start enzyme and the handling in clean rooms is much easier with hot start polymerases not requiring ice, this polymerase was mixed 2:1 with a monoclonal antibody that binds to and inactivates Taq DNA polymerases below 70°C. PCR was performed under the following conditions: 1x PCR Buffer, 2.375mM MgCl₂ (MgCl₂ from the PCR buffer is included). The temperature profile of the PCR was: initial denaturation 94°C for 2min, 60 cycles 94°C for 20sec, 50°C for 30sec, 72°C for 30sec, and final extension at 72°C for 10min.

7.2.3 AmpliTaq Gold® (Applied Biosystems, USA)

PCR was completed using 1x GeneAmp® PCR buffer II and 2.5mM MgCl₂ in a final volume of 20μl, according to a temperature profile of 94°C for 5min, 60 cycles 94°C for 20sec, 50°C for 30sec, 72°C for 30sec and a final extension of 72°C for 10min.

7.2.4 Platinum® *Taq* DNA Polymerase (Invitrogen, USA)

PCR was performed using 1x PCR Buffer and 1.5mM MgCl₂ in a final volume of 20µl and according to a temperature profile of 94°C for 2min, followed by 60 cycles at 94°C for 20sec, 50°C for 30sec, 72°C for 30sec and final extension at 72°C for 10min.

7.2.5 Platinum® *Taq* DNA Polymerase High Fidelity (Invitrogen)

PCR was completed using 1x High Fidelity PCR Buffer and 2mM MgSO₄ in a final volume of 20μl according to the following temperature profile: 94°C for 2min, 60 cycles 94°C for 20sec, 50°C for 30sec, 68°C for 30sec and final extension at 70°C for 10min.

7.2.6 Platinum® PCR SuperMix (Invitrogen)

This is a ready to use mixture of polymerase, buffer and dNTP's. To adjust the amount of polymerase (1U) for all other experiments, the MgCl₂ concentration in the final volume of 50μl was 1.5mM and each dNTP 200μM. Initial denaturation was done at 94°C for 2min, followed by 60 cycles at 94°C for 20sec, 50°C for 30sec, 70°C for 30sec and a final extension at 72°C for 10min.

7.2.7 AccuPrimeTM Taq DNA Polymerase (Invitrogen)

PCR was executed using 1x AccuPrime PCR Buffer I (already including MgCl₂, resulting in a final concentration of 1.5mM) in a final volume of 20μl and according to the following temperature profile: 94°C for 2min, 60 cycles 94°C for 20sec, 50°C for 30sec, 68°C for 30sec and a final extension at 70°C for 10min.

7.2.8 BD AdvantageTM 2 Polymerase Mix (BD Bioscience, USA)

PCR was performed using 1x Advantage 2 SA PCR Buffer (resulting in a final concentration of 2mM MgCl₂) in a final volume of 20µl and according to the following temperature profile: initial denaturation at 94°C for 2min, 60 cycles 94°C for 20sec, 50°C for 30sec, 68°C for 30sec and a final extension at 70°C for 10min.

7.2.9 SuperTaq^{TM*} Plus (Ambion, USA)

PCR was completed using 1x Long PCR Buffer and 1.5mM MgSO₄ in a final volume of 20μl with a temperature profile as follows: initial denaturation at 94°C for 2min, followed by 60cycles at 94°C for 20sec, 50°C for 30sec, 68°C for 30sec and a final extension at 70°C for 10min.

7.2.10 HotMaster™ Taq DNA Polymerase (Eppendorf, Germany)

PCR was executed using 1x HotMaster Taq Buffer with 25mM Mg²⁺ (resulting in an end concentration of 2.5mM Mg²⁺) in a final volume of 20µl and according to a temperature profile with initial denaturation at 94°C for 2min, followed by 60 cycles at 94°C for 20sec, 50°C for 30sec, 65°C for 30sec and a final extension at 70°C for 10min.

7.2.11 MasterTaq® Kit (Eppendorf)

PCR was completed using 1x Taq Buffer with Mg²⁺ (resulting in an end concentration of 1.5mM Mg²⁺) and 1x TaqMaster PCR Enhancer in a final volume of 20μl and according to the following temperature profile: initial denaturation at 94°C for 2min, followed by

60 cycles at 94°C for 20sec, 50°C for 30sec, 72°C for 30sec and a final extension at 72°C for 10min.

7.2.12 BDTMTaq Full Hot Start DNA Polymerase Enzyme Mix (BD Bioscience)

PCR was completed using 1x Taq Full PCR Buffer (resulting in 2mM MgCl2) in a final volume of 20µl and according to the following temperature profile: 94°C for 2min, followed by 60 cycles at 94°C for 20sec, 50°C for 30sec, 68°C for 30sec and a final extension at 72°C for 10min.

Results & Discussion

To 1b):

After digestion of the powder at the same temperature and time, it is noteworthy that the powder was almost completely digested in the extraction buffer using the silica method, whereas almost no change of the powder volume could be observed for the proteinase K digestion buffer of the DNA IQ Kit. This difference could be explained by the overall smaller amount of buffer or the lower concentration of EDTA in the IQ extraction buffer (only 50mM vs. 450mM for the silica extraction buffer), both of which are too small to demineralize (or decalcify) the bone material. As the amount of proteinase K is the same in both buffers, proteinase K is unlikely to be the limiting factor.

To 2.3):

Apart from EDTA and sarcosyl, which were tested in separate experiments, the initial extraction buffer contained proteinase K, DTT, PTB and PVP. The later four reagents were tested in all possible combinations in a single experiment. Thus, for each reagent, we had eight different buffers containing the reagent and eight corresponding buffers lacking the reagent (Table S5). We compared both all means for the corresponding pairs from a certain reagent and the individual relative values for each sample from each buffer pair. Using a paired t-Test, only proteinase K showed a statistically significant improvement for both comparisons (P<0.01 and P<0.05, respectively). DTT showed a tendency of a positive trend (average performance with DTT = 0.86, without = 0.72), but neither comparison was statistically significant (P=0.08 in both cases). PTB showed no effect in either direction (P>0.1 in both cases; see also table S5), while PVP even had an

adverse effect with statistical significance for the comparison of the means (P<0.05) and close to statistical significance for the individual values (P=0.07). Due to these results, we retained proteinase K in the buffer and performed a more in-depth evaluation of the effects of reducing agents.

Supplementary Tables:

Sample ID	Type	Origin	Country	¹⁴ C age	Dating No.	Citation
233	bone	Potocka zijalka	Slovenia	26,900 +/- 110 BP	Beta - 143240	(1)
243	bone	Vindija layer H	Croatia	42,660 +/- 900 BP	Beta - 156101	(1)
322	bone	Ramesch cave	Austria	-		
323	bone	Ramesch cave	Austria	-		
330	bone	Winden cave	Austria	-		
576	bone	Gailenreuth	Germany	-		
748	tooth	Wildkirchli	Switzerland	-		
751	tooth	Ossenyaga cave	Russia	-		
891	bone	Gamssulzen cave	Austria	44,160 (+1400/-1190)	KIA25287	(8)
1050	bone	Harz	Germany	-	-	-

Table S1: Samples used in this study. Laboratory internal sample ID, type of material, cave in which the sample was found (origin), country, ¹⁴C age, dating number and citation are given.

	323	330	748	Sample ID				
Extraction	(Copy Number per gram (Standard Deviation)						
Method	F	Relative to Best Me	ethod	Average				
Silica	0	1,953 (1,933)	206 (504)					
Omod	0	1	0.07	0.36				
Centricon	0	0	2,894 (3,314)					
Centricon	0	0	1	0.33				
Phenol	0	399 (978)	363 (414)					
Chloroform	0	0.2	0.13	0.11				
DNeasy	16 (38)	0	173 (316)					
tissue kit	0.03	0	0.06	0.03				
All-tissue	72 (177)	19 (47)	599 (771)					
DNA-kit	0.12	0.01	0.21	0.11				
DNA IQ	626 (626)	13 (32)	2,872 (2,558)					
system	1	0.01	0.99	0.67				

Table S2a: Results of quantitative PCR comparing different extraction methods using 40 mg of sample powder. Absolute copy numbers per gram of powder (standard deviation in parenthesis) are shown together with relative numbers compared to the best performing method in this test series, calculated separately for each sample. This is the same table as Table 1a, including standard deviations for the absolute copy numbers.

		323			330			748			891		Sa	ample ID	
Extraction		Copy Number per gram (Standard deviation)													
Method		Relative to Best Method								,	Average				
	original	1:10	total	original	1:10	total	original	1:10	total	original	1:10	total	original	1:10	total
Silica	0	0	0	0	70 (53)	35 (51)	36 (19)	151 (145)	93 (115)	22 (100)	11,065 (1,164)	5,555 (5,809)			
Omou	0	0	0	0	1	0.21	0.2	1	1	0.01	1	1	0.05 *	0.75	0.4
DNA IQ	98 (140)	48 (117)	73 (131)	330 (382)	0	165 (318)	178 (203)	0	89 (173)	2,142 (260)	420 (171)	1,281 (927)			
System	1	1	1	1	0	1	1	0	0.96	1	0.04	0.23	1	0.26	0.63

Table S2b: Results of quantitative PCR comparing the two best performing extraction methods using 200 mg of sample powder. Absolute copy numbers per gram of powder (standard deviations are given in parentheses) are shown for the pure extract and a 1:10 dilution. Relative numbers for pure extract, dilution and total were separately calculated for each sample, and were compared to the best performing method in this test series. This is the same table as Table 1b, including standard deviations for the absolute copy numbers.

	330	748	891	Sample ID
Surfactant		y Number per gr (Standard Deviation)	am	
	Rela	ative to Best Metl	hod	Average
	94	11,496	585,456	
1% Tween® 20	(33)	(3,105)	(158,150)	
	0.07	0.67	1	0.58
	982	13,266	443,401	
1% Triton®-X 100	(354)	(4,057)	(69,532)	
170 THOME 70 100	0.78	0.78	0.76	0.77
	7	4,893	65,566	
0.5% N-lauryl-sarcosine	(10)	(4,695)	(815)	
,	0.01	0.29	0.11	0.17 *
	0	5,207	89,645	
0.5% SDS	U	(1,113)	(14,791)	
	0	0.30	0.15	0.15 *
	1,263	1,839	13,453	
1% CTAB	(1,786)	(635)	(4,327)	
	1	0.11	0.02	0.38
	249	3,577	277,575	
2 M Urea	(7)	(995)	(10,349)	
	0.20	0.21	0.47	0.29 *
	23	10,322	355,536	
2 M GuCl	(1)	(2,706)	(23,555)	
	0.02	0.60	0.61	0.41
	72	17,108	450,806	
without	(88)	(213)	(3,736)	
	0.06	1	0.77	0.61

Table S3. Results of quantitative PCR comparing different surfactants in the extraction buffer. Absolute copy numbers per of gram powder (standard deviations are given in parenthesis) are shown together, with relative numbers separately compared to the best performing method for each sample in this test series. This is the same table as Table 2, including standard deviations for the absolute copy numbers.

	233	748	891	Sample ID
Temperature and	Cop			
Duration of Incubation	Rela	ative to Best Me	thod	Average
56°C	0	5,771 (3,142)	113,569 (10,851)	
1h	0	0.65	0.41	0.35
37°C	120 (170)	8,868 (6,114)	97,507 (27,340)	
1h	0.06	1	0.35	0.47
37°C	0	7,508 (2,805)	172,780 (51,569)	
Overnight	0	0.85	0.63	0.49
Room Temperature	92 (131)	6,779 (4,120)	275,869 (19,782)	
Overnight	0.05	0.76	1	0.6
Room Temperature	1,876 (2,654)	1,443 (2,041)	139,473 (18,298)	
4 Days	1	0.16	0.51	0.56
Room Temperature	356 (342)	2,129 (2,934)	128,746 (53,787)	
4 Days (+ PK)	0.19	0.24	0.47	0.3

Table S4. Results of quantitative PCR comparing different times and temperatures of incubation of samples in extraction buffer. Absolute copy numbers per gram of powder (standard deviations are given in parentheses) and relative numbers compared to the best performing method in this test series are shown. This is the same table as Table 3, including standard deviations for the absolute copy numbers.

		233	748	891	Sample ID
Buffer	DTD		y Number per gr Standard Deviation)	ram	
Components	PTB	,	tive to Best Met	hod	Average
	+	16,714 (9,729)	5,552 (184)	438,896 (30,686)	
EDTA –	·	1.00	0.44	1.00	0.81
EDIA —		7,758 (4,842)	12,660 (11,673)	240,806 (3,547)	
	-	0.46	1.00	0.55	0.67
		3,797	2,952	383,168	0.01
	+	(805)	(2,433)	(77,086)	
EDTA		0.99	0.42	1.00	0.81
PVP		3,817	6,955	299,251	0.01
	_	(5,398)	(4,217)	(99,495)	
		1.00	1.00	0.78	0.93
		12,394	3,681	568,282	
	+	(5,676)	(5,206)	(79,271)	
EDTA _		0.69	0.35	1.00	0.68
DTT		17,894	10,536	244,619	
DII	-	(729)	(8,379)	(32,337)	
		1.00	1.00	0.43	0.81
		14,505	14,433	563,521	
EDTA _ PK	+	(481)	(5,544)	(78,607)	
		0.34	1.00	1.00	0.78
		42,504	29	402,450	
	_	(26,406)	(40)	(177,654)	
		1.00	0.00	0.71	0.57
		11,173	9,772	523,813	
EDTA	+	(6,503)	(1,860)	(48, 100)	
PVP —		0.92	1.00	1.00	0.97
DTT		12,122	5,766	238,216	
DII	-	(8,640)	(4,157)	(75,879)	
		1.00	0.59	0.45	0.68
		48,846	676	447,990	
EDTA	+	(21,563)	(925)	(32,968)	
PVP		1.00	0.04	0.93	0.66
DTT		28,113	17,724	482,233	
PK	-	(5,656)	(13,628)	(276,241)	0.00
		0.58	1.00	1.00	0.86
		23,295	8,553	277,632	
EDTA	+	(1,457)	(5,978)	(71,779)	0.75
PVP —		0.70	0.56	1.00	0.75
PK		33,105	15,344	238,616	
	-	(366)	(8,884)	(120,162)	0.05
		1.00	1.00	0.86	0.95
		42,690	2,641	381,364	
EDTA	+	(3,141)	(2,896)	(35,063)	0.50
DTT —		0.93	0.29	0.53	0.59
PK		45,722	8,953	715,096	
	-	(5,710) 1 00	(6,838)	(174,680) 1 00	1.00
		1.00	1.00	1.00	1.00

Table S5a: Results of quantitative PCR comparing each possible extraction buffer composition in pairwise comparisons to presence (+) or absence (-) of PTB. Absolute copy numbers per gram powder (standard deviations are given in parenthesis) and relative numbers compared to the best performing method in this test series are shown. In three out of eight cases the presence, and in five out of eight cases the absence of PTB performed better in the pairwise comparisons. However, neither the individual nor the summed differences are statistically significant after a paired t-Test (P>0.1). PTB: N-phenacyl thiazolium bromide (2.5mM); PVP: Polyvinylpyrrolidone (1%); DTT: Dithiothreitol (50mM), PK: proteinase K (0.25mg/ml).

		233	748	891	Sample ID
Buffer	PVP	Cop			
Components	PVP	Rela	Average		
		3,817	6,955	299,251	
FDTA —	+	(5,398) 0.49	(4,217) 0.55	(99,495) 1.00	0.68
EDTA —		7,758	12,660	240,806	0.00
	-	(4,842)	(11,673)	(3,547)	
		1.00	1.00	0.80	0.93
		3,797	2,952	383,168	
EDTA	+	(805)	(2,433)	(77,086)	
EDTA PTB —		0.23	0.53	0.87	0.54
FID		16,714 (9,729)	5,552 (184)	438,896 (30,686)	
	-	1.00	1.00	1.00	1.00
		12,122	5,766	238,216	
	+	(8,640)	(4,157)	(75,879)	
EDTA		0.68	0.55	0.97	0.73
DTT		17,894	10,536	244,619 (32,337)	
	-	(729) 1.00	(8,379) 1.00	(32,337) 1.00	1.00
		33,105	15,344	238,616	1.00
EDTA PK –	+	(366)	(8,884)	(120,162)	
		0.78	1.00	0.59	0.79
	-	42,504	29	402,450	
		(26,406)	(40)	(177,654)	
		1.00	0.00	1.00	0.67
	+	11,173	9,772	523,813	
EDTA		(6,503) 0.90	(1,860) 1.00	(48,100) 0.92	0.94
PTB		12,394	3,681	568,282	0.94
DTT	_	(5,676)	(5,206)	(79,271)	
		1.00	0.38	1.00	0.79
		48,846	676	447,990	
EDTA	+	(21,563)	(925)	(32,968)	
PTB		1.00	0.26	1.00	0.75
DTT — PK		42,690 (3,141)	2,641 (2,896)	381,364 (35,063)	
	-	0.87	1.00	0.85	0.91
		23,295	8,553	277,632	
EDTA	+	(1,457)	(5,978)	(71,779)	
PTB —		1.00	0.59	0.49	0.70
PK		14,505	14,433	563,521	
	-	(481) 0.62	(5,544) 1.00	(78,607) 1.00	0.87
		28,113	17,724	482,233	0.07
EDT.	+	(5,656)	(13,628)	(276,241)	
EDTA DTT —		0.61	1.00	0.67	0.76
PK —		45,722	8,953	715,096	
	-	(5,710)	(6,838)	(174,680)	0.04
		1.00	0.51	1.00	0.84

Table S5b: Results of quantitative PCR comparing each possible extraction buffer composition in pairwise comparisons to presence (+) or absence (-) of PVP. Absolute copy numbers per gram powder (standard deviations are given in parenthesis) and relative numbers compared to the best performing method in this test series are shown. In two out of eight cases the presence, and in six out of eight cases the absence of PVP performed better in the pairwise comparisons. Although no individual comparison is statistically significant, depending on the way of comparison (see above) the addition of PVP results in overall statistically significantly worse results after a paired t-Test (P<0.05 and P=0.07, respectively). PTB: N-phenacyl thiazolium bromide (2.5mM); PVP:

	233	748	891	Sample ID
	Co			
DTT	_			
				Average
+				0.04
				0.94
-				0.81
				0.01
+				
				0.88
_	(5,398)	(4,217)	(99,495)	
	0.31	1.00	1.00	0.77
	12,394	3,681	568,282	
+			(79,271)	
				0.80
-				0.02
				0.92
+				
				1.00
				1.00
-				
	0.93	0.00	0.56	0.50
+	11,173	9,772	523,813	
	(6,503)	(1,860)	(48,100)	
	1.00	1.00	1.00	1.00
	3,797	2,952	383,168	
				- 10 h
				0.46 *
_				
+				0.69
				0.09
_				
				0.70
		17.724		
+	(5,656)	(13,628)	(276,241)	
	0.85	1.00	1.00	0.95
	33,105	15,344	238,616	
-	(366)	(8,884)	(120,162)	
				0.79
+				0.60
				0.62
-	0.34	1.00	1.00	0.78
	- + - +	## 17,894 17,894 (729) 1.00 7,758 - (4,842) 0.43 12,122 + (8,640) 1.00 3,817 - (5,398) 0.31 12,394 + (5,676) 0.74 - (9,729) 1.00 45,722 + (5,710) 1.00 42,504 - (26,406) 0.93 11,173 + (6,503) 1.00 3,797 - (805) 0.34 48,846 + (21,563) 1.00 23,295 - (1,457) 0.48 28,113 + (5,656) 0.85 33,105 - (366) 1.00 42,690 + (3,141) 1.00 14,505 - (481)	## DTT ## Relative to Best Met	17,894

Table S5c: Results of quantitative PCR comparing each possible extraction buffer composition in pairwise comparisons to presence (+) or absence (-) of DTT. Absolute copy numbers per gram powder (standard deviations are given in parenthesis) and relative numbers compared to the best performing method in this test series are shown. In five out of eight cases the presence, and in three out of eight cases the absence of DTT performed better in the pairwise comparisons. Only one pairwise comparison was significantly different after a paired t-Test (*, P<0.05), whereas the summed comparison shows no significant difference (P=0.08 in both comparisons), although the additions of DTT seems to show a slightly positive effect. PTB: N-phenacyl thiazolium bromide (2.5mM); PVP: Polyvinylpyrrolidone (1%); DTT: Dithiothreitol (50mM), PK: Proteinase K (0.25mg/ml).

		233	748	891	Sample ID
Buffer		Co	opy Number per g		
Components	PK		(Standard Deviation)		
Components	FK	Re	elative to Best Me	thod	Average
		42,504	29	402,450	
	+	(26,406) 1.00	(40) 0.00	(177,654) 1.00	0.67
EDTA -		7,758			0.07
25	_	7,756 (4,842)	12,660 (11,673)	240,806 (3,547)	
	_	0.18	1.00	0.60	0.59
		33,105	15,344	238,616	0.00
	+	(366)	(8,884)	(120,162)	
EDTA		1.00	1.00	0.80	0.93
PVP		3,817	6,955	299,251	
	-	(5,398)	(4,217)	(99,495)	
		0.12	0.45	1.00	0.52
		45,722	8,953	715,096	
	+	(5,710)	(6,838)	(174,680)	
EDTA _		1.00	0.85	1.00	0.95
DTT		17,894	10,536	244,619	
		(729)	(8,379)	(32,337)	
		0.39	1.00	0.34	0.58
	+	14,505	14,433	563,521	
EDT A		(481)	(5,544)	(78,607)	0.00
EDTA _		0.87	1.00	1.00	0.96
PTB	-	16,714 (9,729)	5,552 (184)	438,896 (30,686)	
		1.00	0.38	0.78	0.72
			17,724		0.72
	+	28,113 (5,656)	(13,628)	482,233 (276,241)	
EDTA		1.00	1.00	1.00	1.00
PVP -		12,122	5,766	238,216	
DTT		(8,640)	(4,157)	(75,879)	
		0.43	0.33	0.49	0.42 *
		48,846	676	447,990	
EDTA	+	(21,563)	(925)	(32,968)	
PVP _		1.00	0.07	0.86	0.64
DTT		11,173	9,772	523,813	
PTB	-	(6,503)	(1,860)	(48,100)	0.74
		0.23	1.00	1.00	0.74
		23,295	8,553	277,632	
EDTA	+	(1,457) 1.00	(5,978) 1.00	(71,779) 0.72	0.91
PVP -					0.91
PTB	_	3,797 (805)	2,952 (2,433)	383,168 (77,086)	
		0.16	0.35	1.00	0.50
		42,690	2,641	381,364	3.00
	+	(3,141)	(2,896)	(35,063)	
EDTA		1.00	0.72	0.67	0.80
DTT -		12,394	3,681	568,282	
PTB	-	(5,676)	(5,206)	(79,271)	
		0.29	1.00	1.00	0.76

Table S5d: Results of quantitative PCR comparing each possible extraction buffer composition in pairwise comparisons to presence (+) or absence (-) of proteinase K (PK). Absolute copy numbers per gram powder (standard deviations are given in parenthesis) and relative numbers compared to the best performing method in this test series are shown. In seven out of eight cases the presence, and in one out of eight cases the absence of proteinase K performed better in the pairwise comparisons. Only one pairwise comparison was significantly different after a paired t-Test (*, P<0.05). However, in the summed comparison, proteinase K shows a strong positive effect with statistical significance (P<0.01 and P<0.05, respectively). PTB: N-phenacyl thiazolium bromide (2.5mM); PVP: Polyvinylpyrrolidone (1%); DTT: Dithiothreitol (50mM), PK: Proteinase K (0.25mg/ml).

	233	748	891	Sample ID
Condition	Co	py Number per gi (Standard Deviation)		
	Rel	Average		
- 0.05M	77,649 (9,460)	7,454 (2,725)	751,214 (31,530)	
Tris	1	1	1	1
+ 0.05M	66,618 (8,368)	3,548 (2,833)	615,327 (0)	
Tris	0.86	0.48	0.82	0.72

Table S6: Results of quantitative PCR comparing absence (-) and presence (+) of 0.05M Tris (pH 8.0) in the extraction buffer. Absolute copy numbers per gram of powder and relative numbers compared to the best performing method in this test series are shown. The difference between the two methods is not significant, according to a paired t-Test (P<0.05).

	233	748	891	Sample ID
EDTA	C	opy Number per ((Standard Deviation		
Concentration	Re	elative to Best Me	thod	average
0.45 M	1,356 (288)	15,084 (3,372)	1,482,244 (5,761)	
	0.08	1	1	0.69
0.2 M	16,765 (11,959)	13,943 (1,568)	847,416 (3,294)	
	1	0.92	0.57	0.83
0.1 M	1,046 (1,367)	7,390 (1,116)	358,338 (23,661)	
	0.06	0.49	0.24	0.12 *
0.05 M	0	0	62,912 (1,467)	
	0	0	0.04	0.01 *

Table S7: Results of quantitative PCR comparing different EDTA concentrations in the extraction buffer. Absolute copy numbers per gram powder (and standard deviations in parenthesis) and relative numbers compared to the best performing method in this test series are shown. Significantly worse EDTA concentrations compared to the best performing one (0.2M EDTA) are indicated by an asterisk (*; P<0.05, paired t-Test).

	233	748	891	Sample ID
Reducing Agent	Сор			
3 3	Rela	tive to Best N	/lethod	Average
without	86,724 (19,891)	1,105 (211)	621,418 (31,671)	
	1	0.15	0.94	0.7
DTT	20,469 (6,548)	5,220 (5,118)	553,495 (6,512)	
	0.24	0.73	0.83	0.6
5,5'-Dithiobis-	75,274 (4,426)	7,130 (10,083)	599,191 (39,923)	
(2-nitrobenzoic acid)	0.87	1	0.9	0.92
2-Mercapto-	19,743 (4,754)	4,300 (605)	559,805 (52,618)	
ethanol	0.23	0.6	0.84	0.56
2-Mercaptobenzo-	55,768 (5,242)	2,521 (3,565)	505,898 (21,820)	
thiazol sodiumsalt	0.64	0.35	0.76	0.58
2-Naphtalin-	25,151 (1,380)	0	201,472 (73,380)	
thiol	0.29	0	0.3	0.2 *
2-Amino-5-mercapto-	39,119 (6,112)	2,939 (1,981)	662,876 (33,784)	
1,3,4-thiadiazol	0.45	0.41	1	0.62
Thioglycolacid	30,506 (239)	4,397 (2,058)	645,816 (58,180)	
· · · · · · · · · · · · · · · · · · ·	0.35	0.62	0.97	0.46
tort Dadosasthial	21,926 (1,480)	4,373 (4,595)	582,423 (72,908)	
tert-Dodecanthiol	0.25	0.61	0.88	0.58

Table S8: Results of quantitative PCR comparing different reducing agents in the extraction buffer. Absolute copy numbers per gram of powder and relative numbers compared to the best performing method in this test series are shown. Only one reducing agent (2-Naphtalinthiol) performed significantly worse than the best performing one (5,5'-Dithiobis-(2-nitrobenzoic acid)), according to a paired t-Test (*, P<0.05)

	233	748	891	Sample ID
Condition	Со	py Number per g (Standard Deviation		
	Rel	ative to Best Met	thod	Average
Coarse	38 (38)	0	115,092 (11,840)	
1.Extraction	0	0	0.25	0.08 *
Fine	13,803 (2,670)	3,127 (4,422)	455,327 (26,054)	
1.Extraction	1	1	1	1
Coarse	3,960 (2,633)	0	182,488 (16,693)	
2.Extraction	1	0	1	0.67
Fine	0	0	53,084 (203)	
2.Extraction	0	0	0.29	0.1
Coarse	3,998	0	297,580	
Sum	0.29	0	0.59	0.29 *
Fine	13,803	3,127	508,411	
Sum	1	1	1	1

Table S9: Results of quantitative PCR comparing fine powder (obtained via Freezer Mill) with coarse samples (obtained via mortar and pestle). Absolute copy numbers per gram of powder, and relative numbers compared to the best performing method in this test series are shown. Relative values are calculated separately for the first, the second and both extractions together. In the last two rows, the first and second extraction of the "coarse" and "fine" powder, respectively, were summed. A paired t-Test indicates that the coarsely ground samples performed more poorly statistically in the first extraction (*, P<0.05) and if both extractions were taken together. The finely ground samples in the 2nd extraction (re-extraction) were not statistically worse when compared to the coarsely ground samples.

	233	748	891	Sample ID
Salt	Co	opy Number per g (Standard Deviation)		
(Concentration)	Re	elative to Best Met	hod	Average
KCI (saturated,	1,222 (132)	147 (207)	82,566 (10,953)	
about 4.6 M)	0.02	0.01	0.07	0.03 *
GuCl (saturated, about 22	54,956 (3,881)	273 (78)	547,916 (65,902)	
M)	0.99	0.03	0.49	0.50
GuSCN a)	55,522 (14,601)	3,324 (331)	401,612 (10,017)	
(5M)	1	0.34	0.36	0.57
GuSCN b)	6,927 (3,292)	7,920 (5,653)	1,088,570 (58,806)	
(5M)	0.12	0.80	0.96	0.63
NaClO ₄	36,864 (10,873)	5,510 (3,582)	436,263 (14,508)	
(5M)	0.66	0.56	0.39	0.54
NaCl (saturated, about 6	40,266 (335)	9,878 (2,756)	763,410 (15,868)	
M)	0.73	1	0.68	0.80
Nal (saturated, about 12	631 (99)	6,764 (4,375)	1,128,573 (23,459)	
M)	0.01	0.68	1	0.56

Table S10: Results of quantitative PCR comparing different salts for binding DNA to silica. Absolute copy numbers per gram of powder (standard deviation are given in parenthesis) and relative numbers compared to the best performing method in this test series are shown. This is the same table as Table 4, including standard deviations for the absolute copy numbers.

	233	748	891	Sample ID
Age of	Cop	y Number per g (Standard Deviation		
Silica Suspension	Rel	ative to Best Me	thod	Average
1 Week	47,294 (3,169)	1,289 (1,822)	525,707 (74,405)	
	1	0.2	0.93	0.71
2 Weeks	14,817 (6,815)	6,346 (1,347)	547,922 (41,028)	
2 1100KG	0.31	1	0.97	0.76
4 Weeks	34,526 (6,632)	5,861 (2,737)	563,641 (62,129)	
	0.73	0.92	1	0.88
8 Weeks	173 (133)	3,432 (3,320)	448,145 (1,768)	
oooo	0	0.54	0.8	0.45 *
12 Weeks	9,339 (2,154)	1,457 (308)	324,794 (6,405)	
12 1100110	0.2	0.23	0.58	0.34 *
~ 26 Weeks	619 (41)	824 (956)	548,084 (10,809)	
	0.01	0.13	0.97	0.37
> 52 Weeks	46 (65)	0	455,788 (7,191)	
<u></u>	0	0	0.81	0.27 *

Table S11: Results of quantitative PCR comparing the age of the silica suspension ranging from 1 week to 1 year. Absolute copy numbers per gram of powder and relative numbers compared to the best performing method in this test series are shown. Silica suspensions performing statistically worse when compared to the best (4 weeks) are indicated by an asterisk (*, P<0.05).

	233	233 748 891		Sample ID
рН		y Number per s Standard Deviation		
,	Rela	tive to Best Me	ethod	Average
4.0	78,576 (11,754)	12,082 (382)	1,024,274 (20,234)	
1.0	1	1	1	1
6.0	73,574 (3,487)	5,881 (5,608)	770,886 (88,137)	
0.0	0.93	0.49	0.75	0.72
8.0	34,624 (821)	7,095 (1,255)	865,694 (17,101)	
0.0	0.44	0.59	0.85	0.63 *

Table S12: Results of quantitative PCR comparing the pH in the binding buffer (L2, including silica suspension and extraction buffer of the respective sample). Absolute copy numbers per gram of powder and relative numbers compared to the best performing method in this test series are shown. The extraction method using a binding buffer (together with the extraction buffer) at pH 8.0 performed statistically worse as compared to the best (pH 4.0), according to a paired t-Test (*, P<0.05). See also Fig.2.

	233	748	891	Sample ID
Amount of	Co	oy Number per g (Standard Deviation)		
Silica Suspension	Rel	ative to Best Met	thod	Average
5µl	0	791 (1,118)	149,603 (6,356)	
ομι	0	0.12	0.24	0.12 *
10µl	4,583 (177)	3,248 (151)	232,573 (898)	
ισμι	0.08	0.5	0.37	0.32 *
20µl	6,438 (3,115)	2,782 (161)	243,496 (41,193)	
200.	0.12	0.43	0.39	0.31 *
40µl	34,127 (132)	1,757 (2,485)	624,831 (19,308)	
. • .	0.61	0.27	1	0.63
100µl	55,525 (1,073)	6,445 (2,614)	428,116 (11,576)	
.σομι	1	1	0.69	0.9

Table S13: Results of quantitative PCR comparing the amount of silica suspension used in the binding step. Absolute copy numbers per gram of powder and relative numbers compared to the best performing method in this test series are shown. 5, 10 and 20μ l silica suspensions performed statistically worse (*, t-Test, P<0.05), as compared to the best performing method (100μ l).

a)

	748	891	Sample ID
Time of		per per gram Deviation)	
Incubation	Relative to	Best Method	Average
1/4 h	26 (36)	16,940 (7,772)	
	0	0.03	0.02 *
1 h	3 (5)	241,154 (45,191)	
• • •	0	0.5	0.25
3 h	5,773 (7,192)	473,725 (82,187)	
•	1	0.98	0.99
Overnight	8 (11)	484,848 (27,405)	
O vorringine	0	1	0.5

b)				
		748	891	sample ID
-	Time of		er per gram Deviation)	
	Incubation	Relative to I	Best Method	Average
-		1,173	579,012	
	1.5 h	(1,659)	(13,241)	
		0.52	0.62	0.57
-		162	935,797	
	3 h	(229)	(72,696)	
		0.03	1	0.52
-		2,266	627,813	_
	6 h	(2,510)	(17,228)	
		1	0.67	0.84

Table S14: Results of quantitative PCR comparing different incubation durations for binding DNA to silica using 100mg sample, 3M NaCl as binding buffer and adjusting pH to 4.0 (this test was done twice a) and b)). Absolute copy numbers per gram of powder and relative numbers compared to the best performing method in each test series are shown. Only ½ h incubation in a) performed statistically worse, as compared to the best performing method (3h) after a t-Test (*, P<0.05). Sample 748 is extremely variable due to overall low copy numbers, and the results from this sample must be interpreted with caution. If only the higher, and therefore more reliable, numbers from sample 891 are

taken into	account,	the yield	appears to	o level	off a	fter 3	hours,	a reason	able pro	cessing
time.										

	748	891	Sample ID		
Silica		Copy Number per gram (Standard Deviation)			
Type	Relative to I	Average			
"Normal"	4,212 (2,252)	1,182,728 (27,048)			
TTOTTI CIT	0.7	0.99	0.85		
"New"	6,045 (3,397)	1,196,603 (109,317)			
14000	1	1	1		

Table S15: Results of quantitative PCR comparing the setup of the silica suspension. "Normal" was prepared after (2), "New" see supplement text (chapter 3.7). Absolute copy numbers per gram of powder and relative numbers compared to the best performing method in this test series are shown. The averages are not statistically significant according to a t-Test (*, P<0.05).

	748	Sample ID		
Wash	Copy Nu (Stand			
Solution	Relative	Average		
Company	6,045 (3,397)	1,196,603 (109,317)		
"New Wash"	0.6	0.91	0.76	
alternative	10,015 (3,109)	1,127,417 (61,852)		
1	1	0.85	0.93	
alternative	691 (784)	1,320,877 (66,433)		
2	0.07	1	0.54	
alternative	2,846 (4,025)	584,570 (69,147)		
3	0.28	0.44	0.36	
alternative	0	0		
4	0	0	0 *	

Table S16: Results of quantitative PCR comparing commercial and alternative wash solutions. See supplement text for composition (chapter 3.8).

Absolute copy numbers per gram of powder and relative numbers compared to the best performing method in this test series are shown. Wash solutions performing significantly worse statistically, as compared to the best performing one (alternative 1), are indicated with an asterisk (*, t-Test, P<0.05).

	233	243	322	576	748	751	891	1050	Sample ID
				Copy Number					-
Extraction Method				Relative to E	Best Method	1			Average
A	266,572 (76,109)	466 (541)	62 (123)	18,458 (7,046)	2,234 (1,381)	2,454 (1,249)	253,083 (119,662)	1,515 (1,088)	
Silica	0.70	0.27	0.04	0.57	0.37	0.31	1	1	0.53
B Modified Silica - NaCl	15,868 (27,672)	249 (382)	0	4,835 (5,641)	230 (445)	844 (1,602)	24,288 (26,966)	0	
	0.04	0.15	0	0.15	0.04	0.11	0.10	0	0.07 *
C Modified Silica - GuSCN	378,173 (31,423)	1,719 (965)	1,626 (1,514)	32,142 (5,117)	6,097 (3,522)	8,057 (2,281)	56,903 (12,434)	1,116 (719)	
	1	1	1	1	1	1	0.23	0.74	0.87
D Phenol Chloroform	133 (256)	0	0	11,053 (9,550)	563 (984)	3,605 (4,950)	149,439 (124,014)	943 (842)	
	0	0	0	0.34	0.09	0.45	0.59	0.62	0.26 *
E Ethanol Precipitation	6,709 (6,209)	0	0	0	0	0	6,878 (4,478)	0	
	0.02	0	0	0	0	0	0.03	0	0.01 *
F Qiagen Kit	0	0	0	317 (464)	145 (288)	2,054 (520)	103,646 (59,696)	112 (132)	
	0	0	0	0.01	0.02	0.26	0.41	0.07	0.10 *

Table S17: Results of quantitative PCR comparing six different extraction techniques using eight ancient cave bear samples each in duplicate. Copy numbers were calculated from three measurements of two dilutions of both extracts, per sample per method (12 data points per method altogether). Absolute copy numbers per gram of powder and relative numbers compared to the best performing method in this test series are shown. Extractions methods performing significantly worse as compared to the best one (C – modified Silica – GuSCN), according to a t-Test are indicated by an asterisk (*, P<0.05).

	233	243	322	576	748	751	891	1050	Sample ID	Extraction Control
		Inhib	Average							
Extraction Method		Inhibitio	Average							
A	0.25	3.62	-0.75	-0.14	-3.06	-6.64	-5.91	-3.19	-1.98	-2.56
Silica	-2.96	-1.56	-1.50	-2.12	-1.23	-0.95	-0.64	-4.89	-1.98	-5.79
В	100.00	49.24	100.00	99.16	65.34	18.20	84.91	100.00	77.11	3.49
Modified Silica - NaCl	100.00	19.79	100.00	74.19	19.66	18.00	32.77	100.00	59.80	0.21
С	6.40	-2.34	1.43	-4.73	-7.80	-0.65	-0.80	-6.97	-1.93	-12.05
Modified Silica - GuSCN	-0.57	-2.25	-0.50	0.49	0.78	-0.86	1.31	3.28	0.21	0.73
D	99.95	100.00	51.16	80.32	10.34	2.30	94.13	92.63	66.35	1.66
Phenol Chloroform	77.16	100.00	15.11	14.97	-1.47	-1.04	37.54	26.20	34.02	0.19
E	29.83	-4.38	91.07	-4.22	-3.56	-4.09	5.00	-3.92	13.22	1.96
Ethanol Precipitation	5.33	1.44	26.08	0.89	1.54	-1.34	1.05	2.34	4.67	1.89
F	100.00	100.00	99.98	99.99	70.80	21.94	60.69	99.98	81.67	-2.32
Qiagen Kit	100.00	100.00	25.32	59.68	5.03	1.23	1.46	23.89	40.44	-0.67

Table S18: Results of the inhibition test performed as described under 6 (Supplement: Material and Method) Ancient DNA extracts were used to spike a quantitative human PCR assay for each extraction in triplicate. The difference of the mean from six measurements (3 qPCR's for each of the two extractions per method) per sample and the mean of the modern human DNA without spiking was calculated as a percentage (upper row for each method – Inhibition in % (derived from the amount of DNA)). Inhibition in % derived from the efficiency of each PCR (lower row and shaded) was calculated as a percentage of the difference between the

mean from six different measurements (3 qPCR's for each of the two extractions per method) per sample and the mean from the efficiencies of the modern DNA without spiking. Numbers in red indicate an inhibition higher than the double standard deviations of the modern human DNA samples without spiking (for inhibition in % derived from the amount of DNA it is >11,42 % and for inhibition derived from the efficiency of each PCR it is > 9,12%).

References

- 1. Hofreiter, M., G. Rabeder, V. Jaenicke-Despres, G. Withalm, D. Nagel, M. Paunovic, G. Jambresic and S. Pääbo. 2004. Evidence for reproductive isolation between cave bear populations. Curr Biol 14:40-43.
- 2. **Boom, R., C.J. Sol, M.M. Salimans, C.L. Jansen, P.M. Wertheim-van Dillen and J. van der Noordaa.** 1990. Rapid and simple method for purification of nucleic acids. J Clin Microbiol *28*:495-503.
- 3. **Leonard, J.A., R.K. Wayne and A. Cooper.** 2000. Population genetics of ice age brown bears. Proc Natl Acad Sci U S A *97*:1651-1654.
- 4. **Kalmar, T., C.Z. Bachrati, A. Marcsik and I. Rasko.** 2000. A simple and efficient method for PCR amplifiable DNA extraction from ancient bones. Nucleic Acids Res 28:E67.
- 5. **Kwok, S., D.E. Kellogg, N. McKinney, D. Spasic, L. Goda, C. Levenson and J.J. Sninsky.** 1990. Effects of primer-template mismatches on the polymerase chain reaction: human immunodeficiency virus type 1 model studies. Nucleic Acids Res *18*:999-1005.
- 6. Walker, J., A.D.J. Hedges, B.P. Perodeau, K.E. Landry, N. Stoilova, M.E. Laborde, J. Shewale, S. Sinha et al. 2005 Multiplex polymerase chain reaction for simultaneous quantitation of human nuclear, mitochondrial, and male Y-chromosome DNA: application in human identification. Analytical Biochemistry 337: 89-97.
- 7. **Ramakers, C., J.M. Ruijter, R.H.L. Deprez and A.F.M. Moorman.** 2003 Assumption-free analysis of quantitative real-time polymerase chain reaction (PCR) data. Neuroscience Letters *339*: 62-66.
- 8. Noonan, J. P.,M. Hofreiter, D. Smith, J.R. Priest, N. Rohland, G. Rabeder, J. Krause, J.C. Detter, S. Pääbo and E.M. Rubin. 2005 Genomic sequencing of Pleistocene cave bears. Science 309 (5734): 597-9.