

Human Bone Paleoproteomics Utilizing the Single-Pot, Solid-Phase-Enhanced Sample Preparation Method to Maximize Detected Proteins and Reduce Humics

Timothy P. Cleland*

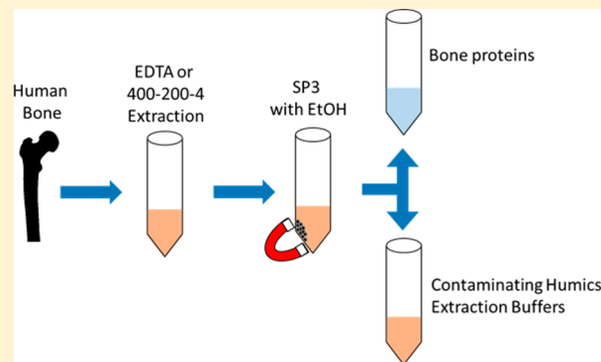
Museum Conservation Institute, Smithsonian Institution, Suitland, Maryland 20746, United States

Supporting Information

ABSTRACT: Sample preparation has become an important part of bone proteomics and paleoproteomics and remains one of the major challenges to maximizing the number of proteins characterized from bone extractions. Most paleoproteomic studies have relied on in-solution digestion with the inclusion of filter-aided sample preparation (FASP) as effective methods to detect the proteome. However, neither of these are optimal because few proteins have been detected utilizing only in-solution digestion and the molecular weight cutoff of FASP may miss remaining fragments of proteins in fossil bone. The recently developed single-pot, solid-phase-enhanced sample preparation (SP3) overcomes these issues by not relying on molecular weight while still controlling where the proteins are digested. Here, historical human bones were extracted

with either 500 mM tetrasodium EDTA or 400 mM ammonium phosphate dibasic, 200 mM ammonium bicarbonate, 4 M guanidine HCl and digested with the SP3 method. Across all samples, 78 ± 7 (400-200-4) and 79 ± 17 (EDTA) protein accessions were identified, including previously difficult to detect proteins such as osteopontin. SP3 also effectively removed 90% or more of the coextracting humic substances (based on reduced absorbance) from extracted proteins. The utility of SP3 for maximizing the number of protein detections in historical bones is promising for future paleoproteomic studies.

KEYWORDS: SP3, bone, human, EDTA, paleoproteomics, sample preparation, osteopontin, collagen I, humics, femur



INTRODUCTION

Sample preparation remains one of the most challenging aspects in bone proteomics and paleoproteomics because the hydroxyapatite mineralization necessitates special considerations that other tissue types do not require to access the proteome. Recent exhaustive studies have characterized the extractome of bone protein extraction methodology,^{1,2} however, few postextraction preparation methods have been used and/or evaluated. Many bone proteomics studies have relied on the use of dialysis approaches^{3–9} to remove the demineralization and/or solubilization buffers. Dialysis is relatively slow (i.e., can take days to complete) and is inefficient at removing one of the most commonly utilized demineralizing agents, ethylenediaminetetraacetic acid (EDTA).¹⁰ Alternatively, protein precipitation has had limited usage to remove salts before digestion for bone proteomics.^{1,11} Precipitation has been used less because of protein loss and difficulties resolubilizing the precipitated proteins,¹² especially for the large amount of collagen I in bone protein extracts. For the issues of dialysis and protein precipitation to be overcome, buffer exchange and concentration using molecular weight spin filters has become an attractive alternative. An additional benefit to this approach is the ability to digest directly on the filter (i.e., filter-aided sample preparation (FASP)¹³) after

removal of extraction buffer. FASP has been shown to be effective at detecting bone and other mineralized proteomes.¹⁴ For paleoproteomics, the FASP method has been refined by reducing the molecular weight cutoff from 30,000 to 10,000 to reduce loss of proteins that have been fragmented during diagenesis.^{15,16} Despite its shown effectiveness, FASP is likely not an ideal approach for paleoproteomics because it will lose the smallest protein fragments remaining from the most degraded paleoproteomic samples and has been shown to be less effective for small amounts of protein (e.g., 1 μ g).¹⁷ Three additional approaches have been applied in paleoproteomics, including direct protein solubilization and digestion,¹⁸ direct digestion of demineralized bone,¹⁹ and a digestion-free approach.²⁰ The direct protein solubilization and digestion approach does not use any additional cleanup prior to digestion, reducing the likelihood of protein loss. With direct digestion of demineralized bone, access to insoluble components including highly glycosylated collagens becomes possible, leading to an increase in protein identification. Lastly, recent applications of a digestion-free approach have allowed detection of proteins into deep time. This method relies on

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protein fragmentation as a response to diagenesis and reduces the opportunity to detect competing contamination (e.g., keratin). Despite the directness of these three methods, none reduce or remove contaminating humic substances.

In contrast to the FASP method, a novel paramagnetic bead approach with carboxylate-modified surfaces (single-pot solid-phase-enhanced sample preparation (SP3)) has recently been developed²¹ and refined^{17,22} for bottom up sample preparation. SP3 has been hypothesized to function in a process similar to hydrophilic interaction liquid chromatography (HILIC) with binding of the proteins in an organic phase and elution in aqueous phase.²¹ On extant proteins, SP3 has been shown to be effective on small amounts of protein¹⁷ as well as in a variety of sample buffers.^{21,22} One major benefit to SP3 over FASP is that it is not molecular weight dependent and can thus capture a wide variety of proteins and protein fragments. Additionally, SP3 uses a combination of hydrophobic and hydrophilic beads to maximize the types of proteins that are captured regardless of their hydrophobicity.^{17,21,22} Here, the SP3 method is applied to historical human bone proteins to evaluate the ability of the method to (1) detect complex proteomes from two different extraction types and (2) remove humic/contaminating brown substances with no additional treatment. Differences in protein composition between the extraction methods are also discussed.

MATERIALS AND METHODS

Human Bone and Protein Extraction

Nine raw, previously powdered human bones (six femora, two tibiae, and one fibula; Table 1) from North America were

Table 1. Human Bone Powder Weights Used for 400-200-4 and EDTA Extraction

sample identifier	burial date	bone type	EDTA weight (mg)	400-200-4 weight (mg)
GLO-099-2A	1862	left femur	15.8	15.0
GLO-099-2C	1862	left femur	21.2	18.3
6CT58-5-AMM03	1830–1850	left femur	5.9	4.4
6CT58-5-AMM05	1830–1850	left femur	6.3	2.5
6CT58-5-AMM11	1830–1850	left femur	16.8	17.0
6CT58-5-AMM13	1830–1850	right femur	15.1	16.8
51KEYWORDH-CC-08	1850–1899	right tibia	12.7	11.3
7NCE98A-WOODVILLE-06	1790–1850	right femur	28.0	24.8
44HE950-RADFORD-5	1775–1875	left tibia	7.7	6.9

analyzed. Previously powdered bone was chosen to minimize additional destructive sampling of these human remains. Collection access was granted by Smithsonian National Museum of Natural History Skeletal Biology Program. Between 2.5 and 28 mg of powder was homogenized in 600 μ L of either 400 mM ammonium phosphate dibasic, 200 mM ammonium bicarbonate, 4 M guanidine HCl (400-200-4)^{4,23} or 500 mM tetrasodium EDTA using a BeadRuptor Elite (Omni International) for 4 cycles at 6.95 m/s for 45 s with 5

min dwell at -20 °C. Although not strictly necessary to extend the incubation time of the 400-200-4 method beyond the homogenization cycle,⁵ both extraction types were incubated for 66 h at 37 °C with shaking for total incubation consistency. After incubation, both extractions were centrifuged at 18,500 rcf for 10 min and then the supernatant was decanted and saved. Protein concentration for all supernatants was determined by Pierce Protein Plus (Bradford) Assay. Ten micrograms of protein from 400-200-4 or 100 μ L of EDTA were taken for SP3 without any additional treatment.

SP3

Before binding to the magnetic beads, proteins were reduced and alkylated for 10 min at 95 °C in 10 mM TCEP and 40 mM chloroacetamide.²⁴ Proteins from both extractions were then digested following the refined SP3 method in Moggridge et al.²² (March 2018 revision from https://github.com/chruhuges/SP3_JPR-2018). Briefly, a 1:1 mixture of hydrophilic and hydrophobic Sera-Mag Speedbeads (GE Healthcare) was washed three times²¹ in 18.3 M Ω Milli-Q (Millipore) water with incubation on a 3D-printed magnetic stand²⁵ for two minutes between washes. Beads were stored in Milli-Q water until use. To the reduced and alkylated bone protein solutions were added 10 μ L of the prepared beads and a volume of ethanol equal to the total volume (sample + bead volume + reducing/alkylating volume) and allowed to incubate for 10 min at room temperature. After incubation, beads were placed on the magnetic rack for 2 min to collect the beads, and the supernatant was removed and discarded. The beads were subsequently washed three times with 80% ethanol and incubated on the magnetic rack for 2 min, and the supernatant was discarded.

The beads were subsequently resuspended in 90 μ L of 50 mM ammonium bicarbonate and 10 μ L of 0.04 μ g/ μ L Promega sequencing-grade modified trypsin (1:25 trypsin:protein). All samples were incubated overnight at 37 °C; then, peptide supernatants were collected for final C18 stage tip desalting.²⁶ In short, three 3 M Empore C18 disks were cut and placed in 200 μ L tips. The tips were wetted with 100% methanol and then 80% acetonitrile with 0.1% formic acid and equilibrated with 0.1% formic acid; samples were added and then washed with 0.1% formic acid. All peptides were eluted with 80% acetonitrile with 0.1% formic acid and then dried under a vacuum. Final peptide samples were resuspended with 0.1% formic acid to a final concentration of 0.5 μ g/ μ L.

Evaluation of Humic Removal by SP3

A subset of 400-200-4 and EDTA extracts (51KEYWORDH-CC-08, 6CT58-5-AMM13, 7NCE98A-WOODVILLE-06) were treated with SP3 to measure the reduction in potential humic coloration following a protocol similar to Qian and Hettich.²⁷ Briefly, SP3 was applied, and then the pretreatment, post-SP3 binding, and post-2 h digestion were measured using a Biotek Epoch microplate spectrophotometer from 250 to 700 nm. An averaged value blank was subtracted from all absorbance values, and plots were generated in RStudio 1.1.383 from 300 to 700 nm to account for interference from the 96-well plate between 250 and 300 nm.

LC-MS/MS

Peptides (0.5 μ g) were trapped on a 3 cm in-house packed Thermo BioBasic C18 column (75 μ m i.d., 5 μ m particle size) followed by separation on a 20.1 cm in-house-packed Thermo BioBasic C18 column analytical column using a Dionex

Ultimate 3000 at 300 nL/min using the following gradient: 0–8 min 2% B, 8–98 min 2–55% B, 98–100 min 55–90% B, 100–103 min 90% B. Buffer A is 0.1% Optima-grade formic acid in Optima-grade water, and buffer B is 99.9% Optima-grade acetonitrile with 0.1% Optima-grade formic acid. Peptides were nanosprayed into a ThermoScientific LTQ Orbitrap Velos and analyzed with the following parameters: MS1: 60k resolution, 100 ms ion time, 1E6 AGC target; MS2: top 8 peaks fragmented with higher energy collisional dissociation (HCD; 30% NCE), 15k resolution, 5 m/z isolation width, 250 ms AGC, SE5 AGC target. Randomization for injection order was generated using random.org. EDTA and 400-200-4 samples were randomized together and run in at least triplicate.

Data Analysis

The data analysis was performed using MetaMorpheus version 0.0.285, available at <https://github.com/smith-chem-wisc/MetaMorpheus>, against XML databases of the human proteome (Uniprot; downloaded April 4, 2018) and cRAP database (Uniprot, July 23, 2018).^{28–31} The following calibration settings were used: protease = trypsin; maximum missed cleavages = 2; minimum peptide length = 5; maximum peptide length = unspecified; initiator methionine behavior = variable; fixed modifications = carbamidomethyl of C, carbamidomethyl of U; variable modifications = oxidation of M; max modificationss per peptide = 3; max modification isoforms = 4096; precursor mass tolerance = 15 ppm; product mass tolerance = 25 ppm. The combined search database contained 20431 nondecoy protein entries including 230 contaminant sequences. The following G-PTM-D settings were used: protease = trypsin; maximum missed cleavages = 2; minimum peptide length = 5; maximum peptide length = unspecified; initiator methionine behavior = variable; max modification isoforms = 4096; fixed modifications = carbamidomethyl of C, carbamidomethyl of U; variable modifications = oxidation of M; G-PTM-D modifications count = 335 (including deamidation of NQ and hydroxyproline); precursor mass tolerances = 5 ppm around the masses of the 340 modifications; fragment mass tolerance = 20 ppm. The final database search used the following settings: protease = trypsin; maximum missed cleavages = 2; minimum peptide length = 5; maximum peptide length = unspecified; initiator methionine behavior = variable; fixed modifications = carbamidomethyl of C, carbamidomethyl of U; variable modifications = oxidation of M; max modificationss per peptide = 3; max modification isoforms = 4096; precursor mass tolerance = 5 ppm; fragment mass tolerance = 20 ppm; report PSM ambiguity = true. All PSMs, peptides, and proteins were filtered at 1% FDR. Contaminants (keratins and keratin-associated proteins, lysozyme c, trypsin), decoy proteins, and peptide standards (cytochrome *c* digest) were filtered in Microsoft Excel 2016 after the initial FDR filter. Statistical analysis was performed in RStudio 1.1.383 or in Microsoft Excel 2016. For comparison, the data set from Sawafuji et al.³² was searched using Metamorpheus version 0.0.286 using similar parameters to the above.

RESULTS AND DISCUSSION

SP3

Initial SP3 attempts followed the protocols in Hughes et al.²¹ and Sielaff et al.,¹⁷ but these were found to be suboptimal because of the immiscibility between both 400-200-4 or EDTA

buffers and acetonitrile, which required large dilution volumes to produce miscibility (Figure S1). As a result of the acetonitrile immiscibility, the recently refined ethanol SP3 protocol²² was applied, and it was found to be effective for both of these buffers without additional dilution (Figure S1). For the ethanol SP3 protocol, at a 1% FDR following removal of contaminants and decoys, a total of 101 protein accessions were detected between the two extractions across all samples. A nonsignificant difference ($p = 0.78$) of 78 ± 7 and 79 ± 17 protein accessions were detected for the nine 400-200-4 and EDTA extracts, respectively. Individually (Tables S1–S3; Figure 1), significantly more protein accessions were detected for the EDTA extraction than for 400-200-4 extraction for GLO-099-2A ($p < 0.01$), GLO-099-2C ($p < 0.05$), and 6CTS58-5-AMM11 ($p < 0.05$), and significantly more protein accessions were detected for 400-200-4 than EDTA for 44HE950-RADFORD-5 ($p < 0.001$) and 7NCE98A-WOODVILLE-06 ($p < 0.01$). For target peptides within 1% FDR

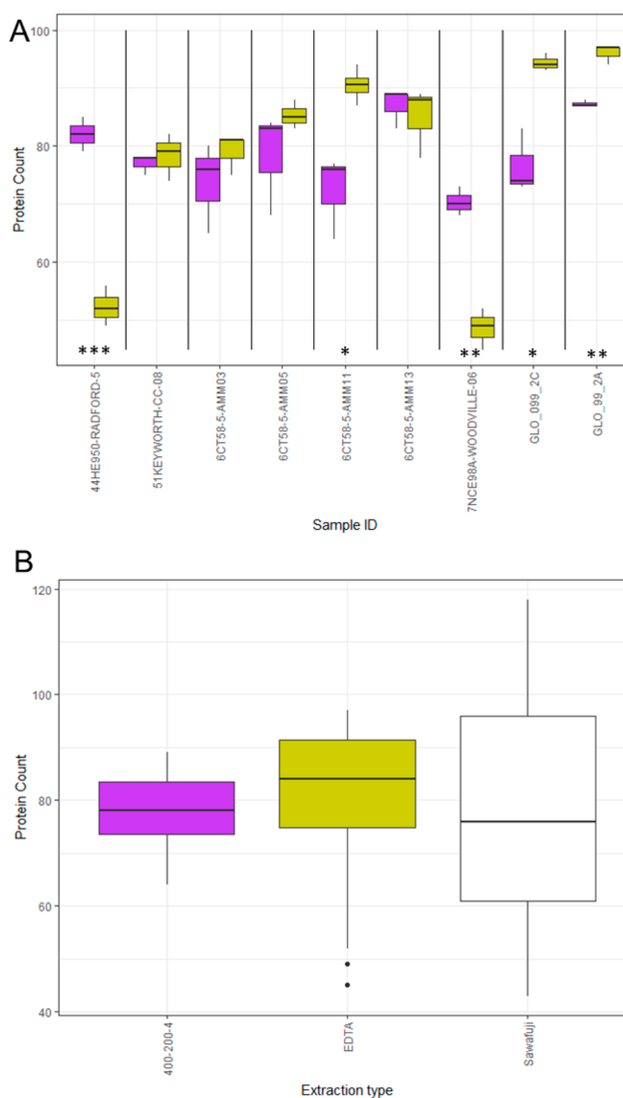


Figure 1. (A) Distribution of protein counts between samples. Purple is 400-200-4, and yellow is EDTA. *** $p < 0.001$; ** $p < 0.01$; * $p < 0.05$. (B) Boxplot comparing protein counts among 400-200-4, EDTA, and Sawafuji et al.³² Boxes indicate the median, first, and third quartile, and the whiskers indicate the minimum and maximum values.

(Table S1), an average of 471 ± 128 peptides was detected for all nine samples with nonsignificant differences ($p = 0.29$) of 453 ± 150 and 490 ± 101 peptides detected for 400-200-4 and EDTA, respectively. A significant difference in the number of peptides detected was observed for 6CT58-5-AMM03 ($p < 0.05$), 7NCE98A-WOODVILLE-06 ($p < 0.05$), and 44HE950-RADFORD-5 ($p < 0.0001$) (400-200-4 > EDTA) and 6CT58-5-AMM11 ($p < 0.0001$) (EDTA > 400-200-4). Before contaminants were removed, 118,329 PSMs, 3975 peptides, and 154 protein groups were detected at a 1% FDR across the 55 raw files.

Comparing the results from the human bone extractions here to recently published human paleoproteomes from Japan,³² insignificant differences for both the peptide count (EDTA: $p = 0.63$; 400-200-4: $p = 0.76$) and protein accessions (EDTA: $p = 0.96$; 400-200-4: $p = 0.97$) were found (Figure 1B). The closeness in protein count suggests that both EDTA and 400-200-4, combined with SP3, are effective at extracting similar numbers of proteins to the on-pellet digestion used by Sawafuji et al.³² The diversity and number of proteins found after SP3 shows that this sample preparation method overcomes previous limitations of both EDTA and 400-200-4. For EDTA, SP3 removes a larger percentage of the salt than dialysis or molecular weight spin columns while reducing the amount of protein loss. The previous applications of 400-200-4 in extant⁴ and fossil remains⁵ only had limited numbers of protein accessions (extant = 12; fossil = 3) detected after in-solution digestion. Again, the major increase in identifications using SP3 on 400-200-4 (6.5–26 \times than previously reported) may be derived from greater removal of the buffer components or that proteins were lost to the dialysis membrane during processing previously. SP3 provides an environment that is conducive to digestion of these bone protein extracts regardless of the original buffer composition.

Removal of Humics

Removal of humic substances that coextract during bone protein extraction has been identified as a challenge in paleoproteomics since at least 2007⁷ when they were mentioned as being critical to remove before detection of peptides was possible. As a result of this realization and other mass spectrometry studies showing that humics lead to ion suppression,^{27,33} efficient removal of humics with limited or no protein loss is potentially critical to future paleoproteomic studies, especially as samples become either older or more densely modified by humics. For reduced humics in these human protein extracts to be evaluated, a subset of the nine samples (i.e., 51KEYWORDH-CC-08, 6CT58-5-AMM13, 7NCE98A-WOODVILLE-06) was chosen to represent the diversity of observed coloration/darkness (Figure 2A). 51KEYWORDH-CC-08 was visually the lightest (Figure 2A) of these selected samples with maximum raw absorbances of 0.27 (EDTA) and 0.15 (400-200-4) at 300 nm. 6CT58-5-AMM13 and 7NCE98A-WOODVILLE-06 (Figure 2A) were both much darker and represent the darkest samples in the entire set. 6CT58-5-AMM13 has maximum raw absorbances of 1.2 (EDTA) and 0.97 (400-200-4), and 7NCE98A-WOODVILLE-06 has maximum raw absorbances of 1.4 (EDTA) and 0.85 (400-200-4) at 300 nm. The raw extracts (Figures S2 and S3) and post-SP3 (Figure 2B, C) binding supernatants follow similar trends across the wavelengths, whereas the post-digestion ammonium bicarbonates, regardless of original extraction buffer, are close in intensity to buffer controls

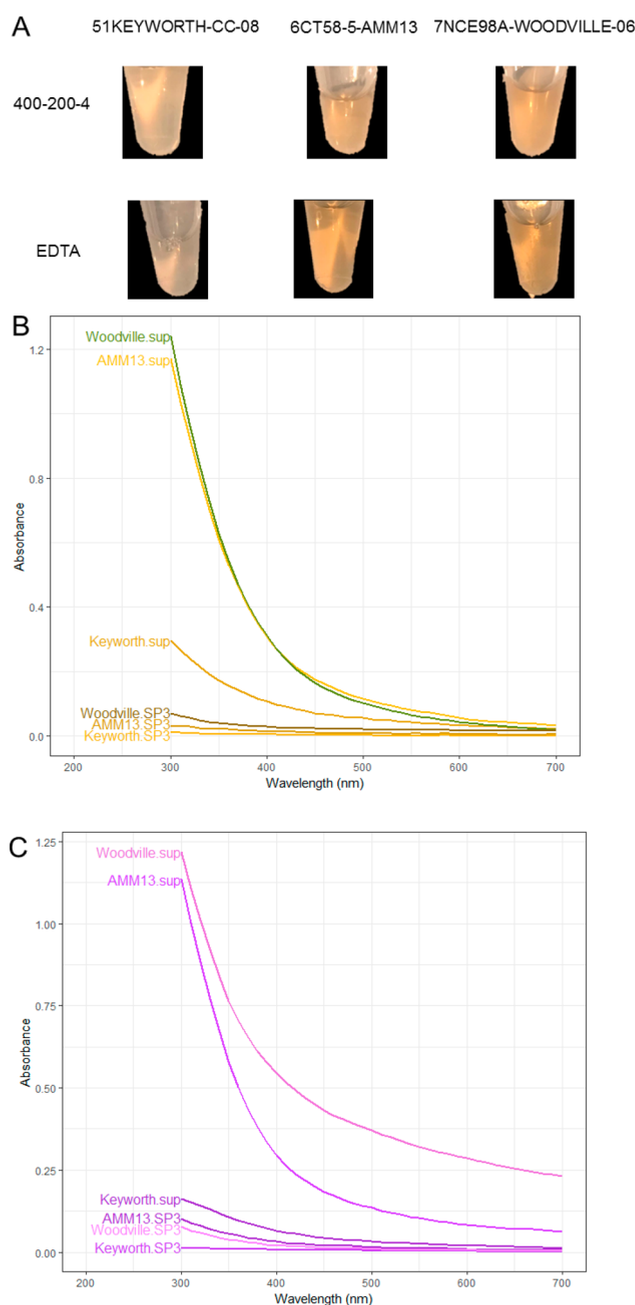


Figure 2. (A) Extractions used for evaluation of humic removal showing differences in coloration. Absorbance plots for EDTA (B) and 400-200-4 (C) showing the decrease in humics signal postdigestion. Absorbance curves labeled .sup indicate supernatant after SP3 binding and .SP3 indicate postdigestion absorbance.

(EDTA: 51KEYWORDH-CC-08 4.8%, 6CT58-5-AMM13 2.9%, 7NCE98A-WOODVILLE-06 4.9% of the maximum intensity at 300 nm; 400-200-4: 51KEYWORDH-CC-08 10.3%, 6CT58-5-AMM13 10.5%, 7NCE98A-WOODVILLE-06 9.1%). The 90+% reduction in humics likely explains the higher number of protein identifications than previously shown for EDTA or 400-200-4 because the ion suppression effects of the humics are removed/reduced. Future investigations of the SP3 approach on much darker extractions will show the effectiveness of this method for removing these contaminating molecules and further support that their removal is necessary to characterize the bone paleoproteome. Additionally, unlike

the recently proposed precipitation and molecular weight spin filtration to remove humics,²⁷ SP3 is rapid and separates the protein content from the humics in a single step in a matter of minutes with no required modification of the sample pH beyond the original buffer pH.

Differences in Extractomics between Buffer Types

For clarity with respect to differences between the EDTA and 400-200-4 extraction, results from all individuals were combined regardless of their burial conditions or burial age. Between the two extractions, 43 protein accessions were significantly greater ($p < 0.05$) in intensity for EDTA than for 400-200-4, and 20 protein accessions were significantly greater in intensity for 400-200-4 than for EDTA ($p < 0.05$). Three main groups of proteins were found to be significantly more concentrated in the EDTA fraction than 400-200-4. These EDTA enriched proteins are (1) small leucine-rich proteoglycans (SLRPs; e.g., lumican, biglycan, decorin) that are associated with collagen fibrillogenesis,³⁴ (2) mineral-associated and mineralization proteins (e.g., pigment epithelium-derived factor (PEDF), alpha-2-hs-glycoprotein (A2HSG), osteomodulin (OMD), osteopontin (SPP1), SPARC, bone sialoprotein 2 (BSP2)³⁴), and (3) various growth factors including insulin-like growth factor 1 (IGF-1), its associated binding protein (insulin-like growth factor binding protein 5), insulin-like growth factor 2 (IGF-2), and transforming growth factor beta-1 (TGF- β 1). In contrast to the EDTA, 400-200-4 has significantly greater relative intensity of collagens I, II, III, and XI. Even though there are significant differences in intensity between the extractions, the majority of the proteins overlap between them (100 of 101 overlap). Consistent with previous bone proteomic and paleoproteomic studies using EDTA and 400-200-4 using other sample preparation methods,^{1,4,19,32} SLRPs were detected from all nine samples, suggesting that SP3 is also effective for buffer removal and digestion of these collagen-associated extracellular proteins.

Despite the increased relative intensity in the EDTA fraction, the mineral-associated proteins were consistently detected in both extraction types, reflecting the original hypothesized⁴ advantages of the 400-200-4 method and its ability to separate mineral proteins from the hydroxyapatite. Unlike the original description of 400-200-4 using in-solution digestion that was only able to detect osteocalcin from the mineral-associated group, the inclusion of SP3 has allowed for detection of osteopontin from all nine individuals in both the EDTA and 400-200-4 extractions (Figure 3). Osteopontin is critical for mineralization and microdamage control³⁵ but has previously been difficult to detect by shotgun proteomics because it is acidic and highly phosphorylated (as many as 36 phosphorylations³⁶). The consistent detection of osteopontin is in contrast to the recent exhaustive extractomics study¹ that was only able to detect it from HCl extraction but not EDTA. Additional enhancements to the SP3 protocol to include an additional phosphopeptide enrichment will show levels of phosphorylation to osteopontin and other phosphorylated bone proteins, and it will show the levels of lability of phosphorylation in deep time.

IGF-1, IGF-2, and TGF- β are associated with bone growth, maintenance, and cell proliferation in bone and are necessary for long-term bone health and remodeling.^{37,38} These growth factors were consistently detected in both EDTA and 400-200-4 (with limited detection of IGF-1), but previous investigations using in-solution for digestion of EDTA^{1,19} or 400-200-4⁴

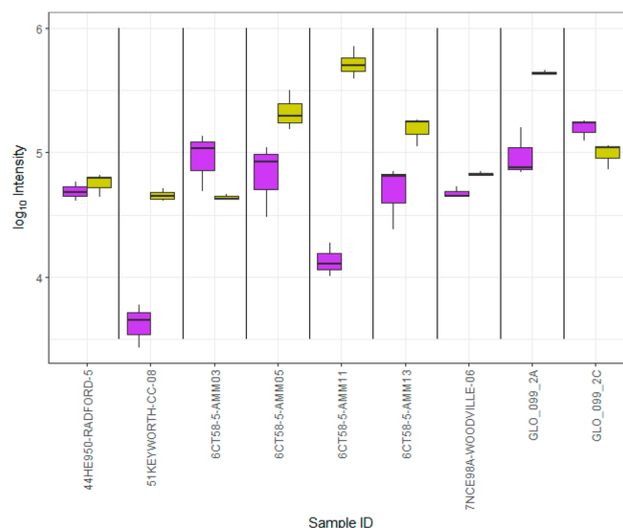


Figure 3. Boxplots of osteopontin intensity by sample and extraction type (purple boxplots: 400-200-4; yellow boxplots: EDTA). Boxes indicate the median, first, and third quartile, and the whiskers indicate the minimum and maximum values.

extractions from extant bone were unable to detect these important proteins. This suggests that SP3 allows digestion of proteins that are at a much lower concentration than collagen I that would have less interaction during in-solution digestion as a result of collagen I interference (EDTA: Collagen I $75.63 \pm 10.08\%$ of TIC, TGF- β $0.04 \pm 0.04\%$, IGF-2 $0.07 \pm 0.06\%$, IGF-1 $0.004 \pm 0.001\%$; 400-200-4: Collagen I $82.59 \pm 3.8\%$, TGF- β $0.01 \pm 0.005\%$, IGF-2 $0.01 \pm 0.007\%$, IGF-1 $0.0003 \pm 0.001\%$). Future assessment of protein concentration using mass spectrometry or other nonmass spectrometry methods between individuals will require the use of only one method despite the considerable overlap between EDTA and 400-200-4. This may be especially true in assessment of protein-based bone quality that requires the examination of particular mineralization proteins.^{39–41}

In addition to differences in mineral-associated proteins, SLRPs, growth factors, and collagens with significantly different intensities between EDTA and 400-200-4, periostin and sclerostin were detected in both extractions but were not found to be significantly different in intensity. Periostin and sclerostin are important proteins for mechanical sensing and bone repair and maintenance that have been targets for osteoporotic treatment and fracture risk assessment.^{42–45} Despite the similar intensities between EDTA and 400-200-4 extracts, sclerostin was more consistently detected in the EDTA extraction, suggesting that a combination of these extraction methods may be required to study these proteins in both extant and paleoproteomic studies.

For the differences between the two extraction methods to be illustrated further, a 2-fold difference cutoff was applied, and 37 and 14 protein accessions were found to be both significant ($p < 0.05$) and 2-fold higher in intensity (Figure 4) for EDTA and 400-200-4, respectively. Like the gross statistical analysis, mineralization proteins, growth factors, and a subset of the SLRPs are more than twice as concentrated in the EDTA extraction. The collagens generally remain after filtering in the 400-200-4, but collagen I α 1 does not pass this threshold. This suggests that 400-200-4 preferentially extracts collagen I α 2 over collagen I α 1. On the basis of these results, future applications examining the mineralization and mineral-

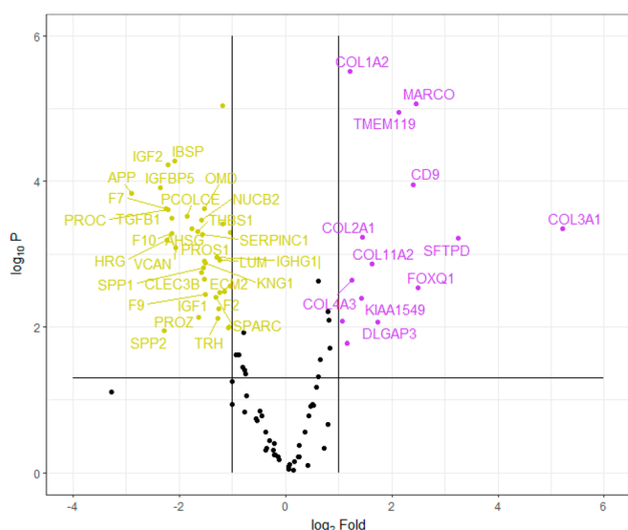


Figure 4. Volcano plot of differences between the EDTA (yellow/left) and 400-200-4 (purple/right) with greater than 2-fold intensity differences.

associated proteins would benefit from using EDTA extraction methodology over 400-200-4 to maximize the amount recovered. In contrast, and supporting what was observed by Schroeter et al. that EDTA is less effective than other methods at extracting collagens,¹ 400-200-4 should be used for studies aiming to maximize collagens recovered from bone.

CONCLUSIONS

SP3 is an effective sample preparation for historical bone that is capable of separating humics while effectively generating peptides from a large diversity of collagenous and non-collagenous bone proteins. Despite this study focusing on historical bone, SP3 is applicable to extant bone proteomics (e.g., protein-based fragility studies, biomarkers) and can result in deep characterization of bone proteomes from a variety of normal and pathological states. Because it efficiently removes EDTA, it is an ideal sample preparation method for bone proteomics and paleoproteomics. Although the goal of the study was to show the efficacy of SP3 for use in bone proteomics and paleoproteomics, ongoing research will compare SP3 to other sample preparation methods, including FASP. Future applications of this methodology to progressively older bone samples will show how effective this approach is into deep time. Finding a large number of proteins from historical and fossil bone will generate a more complete understanding of the biology and paleobiology of the tested organisms. Possible and those already realized, but that may benefit from SP3, outcomes of having more proteins include more complex molecular evolutionary evaluations than morphology or collagen I alone (e.g., ref 20), characterization of diet (e.g., refs 15 and 46), tracing changes in disease state within ancient populations (e.g., ref 47) or across time, and detection of complex post-translational modification changes across species evolution.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jproteome.8b00637.

Table S1, Protein accessions, peptide, and peptide spectral match counts for each sample filtered at 1% FDR; Table S2, Protein groups exported from Metamorphus; Table S3, Protein groups filtered at 1% FDR with decoys and contaminants removed (XLSX)

Figure S1, Acetonitrile immiscibility between EDTA and 400-200-4; Figure S2, Absorbance plots for humic substances of EDTA raw extraction and controls; and Figure S3, Absorbance plot for humic substances of 400-200-4 raw extraction and controls (PDF)

AUTHOR INFORMATION

Corresponding Author

*E-mail: clelandtp@si.edu.

ORCID

Timothy P. Cleland: 0000-0001-9198-2828

Notes

The author declares no competing financial interest.

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