Sex Determination in Highly Fragmented Human DNA by High-Resolution Melting (HRM) Analysis

Brenda A. Álvarez-Sandoval¹, Linda R. Manzanilla², Rafael Montiel¹*¹

Introduction

Traditionally, sex determination in human remains has been based on the dimorphism between the sexes that is present in the majority of human bones [1]. These studies have been based mainly on cranial and pelvic traits [2–15]. Furthermore, other researchers have reported studies based on, hand and foot bones [16,17], scapula [18,19], long bones [4,20–26], patellae [27], sternum [28–30], fourth rib [28], hyoid bone [30], clavicle [31], metacarpus acusticus internus characteristics [32,33] and dentition [34,35].

Other methods to sex determination have been proposed such as anthropometric measurements of the limbs [36–39], hands [40,41], and from length of index and ring finger, and the index and ring finger ratio [42–45]. Nevertheless, it has been reported that 100% of successful sex determinations by osteological measurements only occur when the skeleton is from an adult, it is complete, it is in good condition of preservation, and the morphometric variability in the population to which it belongs is known [46–48].

Advances in the field of molecular genetics has provided more sensitive methods for sex determination, such as the polymerase chain reaction (PCR) that allow amplification of single molecules of target DNA to analytical quantities. Biological remains such as hair, bone fragments, or teeth, usually contain some amounts of degraded DNA [48–57]; therefore, it is possible to establish an individual sex using genetic tests. Recent studies have confirmed that teeth are more refractory to contamination by exogenous DNA than bones, although bones can also be good candidates for analysis under some circumstances [58].

Some proposed methods have been based on the analyses of genetic markers lying in the Y chromosome [59–61], or in the use of both X-chromosomal and Y-chromosomal STRs [47]. Furthermore, a new method to sex determination using shotgun sequencing has been reported [62], although it might be too expensive for routine application in a large number of samples. Most of these procedures are not sensitive enough, are time-consuming, expensive, and require an important amount of sample.

Despite the wide list of molecular methods proposed for sex determination, the method based on the amplification of the human amelogenin gene (AMEL) is the most widely used. This gene, originally sequenced by Nakahori et al [63,64], is found on both chromosomes X and Y (GenBank Accession number M55418 for AMELX-allele and M55419 for AMELY-allele). There are sequence and size divergences between them. AMELX-allele has a size of 2872 bp and is located on the Xp22.1–Xp22.3 region of the X-chromosome, while the human AMELY-allele has a size of 3272 bp and is located on the Yp11.2 region of the Y-chromosome. Male individuals have two different AMEL alleles, one located on chromosome X and one on the Y chromosome, while female individuals have two identical AMEL alleles located on chromosome X [63–72].

Haas-Rochholz and Weiler [73] reported a total of 19 regions of absolute homology that vary in size from 22 to 80 bp; 5 deletions of 1 to 6 bp located on AmelX-allele; and 5 deletions of 1 to 100 bp located on AmelY-allele.

Abstract

Sex identification in ancient human remains is a common problem especially if the skeletons are sub-adult, incomplete or damaged. In this paper we propose a new method to identify sex, based on real-time PCR amplification of small fragments (61 and 64 bp) of the third exon within the amelogenin gene covering a 3 bp deletion on the AMELX-allele, followed by a High Resolution Melting analysis (HRM). HRM is based on the melting curves of amplified fragments. The amelogenin gene is located on both chromosomes X and Y, showing dimorphism in length. This molecular tool is rapid, sensitive and reduces the risk of contamination from exogenous genetic material when used for ancient DNA studies. The accuracy of the new method described here has been corroborated by using control samples of known sex and by contrasting our results with those obtained with other methods. Our method has proven to be useful even in heavily degraded samples, where other previously published methods failed. Stochastic problems such as the random allele drop-out phenomenon are expected to occur in a less severe form, due to the smaller fragment size to be amplified. Thus, their negative effect could be easier to overcome by a proper experimental design.

Citation: Álvarez-Sandoval BA, Manzanilla LR, Montiel R (2014) Sex Determination in Highly Fragmented Human DNA by High-Resolution Melting (HRM) Analysis. PLoS ONE 9(8): e104629. doi:10.1371/journal.pone.0104629

Editor: David Caramelli, University of Florence, Italy

Received January 14, 2014; Accepted July 15, 2014; Published August 6, 2014

Copyright: © 2014 Álvarez-Sandoval et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Funding: This work was supported by Consejo Nacional de Ciencia y Tecnología (CONACYT; www.conacyt.gob.mx) grant No. CB-2008-01-105481 to Dr. Rafael Montiel. M.Sc. Álvarez-Sandoval is supported by a CONACYT fellowship (Reg. No. 300750). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

* Email: montiell@langebio.cinvestav.mx
Materials and Methods

The study population came from Teopancazco, a multiethnic neighborhood centered in the southern part of Teotihuacan, Central Mexico (200–650 A.D.). Bone and tooth samples were collected in several controlled extensive excavations (1997–2005) and provided by Dr. Linda R. Manzanilla, head of the project “Teotihuacan. Elite and rulership. Excavations in Xalla and Teopancazco” (Last authorizations for the Project: Official letter 401.B (4) 19.2013/36/0579 of the National Institute of Anthropology and History). The controlled extensive excavations, with a 1 meter grid, provided the spatial reference for different activity areas and tridimensional measurement of instruments, raw materials and debris. The sampling process was carried out under strict criteria in order to avoid exogenous contamination. The human burial remains from Teopancazco are kept in the Instituto de Investigaciones Antropológicas (UNAM) under the custody of Dr. Linda R. Manzanilla. The samples are part of the Mexican National Heritage.

A subset of 21 samples (from 21 different individuals buried at Teopancazco) was sent to the aDNA laboratory in LANGEBIO, CINVESTAV, Mexico, in order to be analyzed. Seventeen individuals of unknown sex were analyzed, 14 sub-adults (burials 3, 4, 38, 45, 49, 51, 56, 59, 61, 96, 99, 100, 101 and 105) and 3 adults (burials 46, 89, 108). Moreover, 4 adult individuals of known sex (burials 2, 78, 102 and 105) were used as ancient positive controls. For each individual analyzed, a fragment of bone or a complete teeth, in a good state of preservation (without fissures), was used in the analysis. Preliminary DNA characterization showed a general poor DNA preservation at this archaeological site.

In addition, 2 blood samples donated by the first and third author of this paper [1 male (RM) and 1 female (BAA-S)] were considered as modern positive controls. Both samples were collected under informed consent in order to be considered only as positive controls with no other scientific research objective.

Genetic analysis

1.- Contamination control. All analyses were carried out under strict conditions for the analysis of ancient DNA, in a dedicated ultraclean room, positively pressurized with filtered and UV-irradiated air, with separation of pre- and post-PCR areas. Laboratory equipments were treated with 30% bleach for DNA contamination removal. The use of disposable filter-plugged pipette tips, tubes, protective cloths, hair covers, laboratory gloves, surgical masks, glasses, and shoes protectors to prevent contamination was mandatory. Solutions and buffers were irradiated with ultraviolet light. Negative extraction controls and negative PCR controls were always employed.

2.- DNA extraction. Bone and teeth samples were processed in a type II B2 biological security hood, located inside the ultraclean room. The outer surface of the bone or teeth was UV-irradiated during 15 minutes and a dental drill was used to remove 0.1 g of powdered material. Powdered samples were added to 5 ml of extraction buffer (0.5 M Tris-HCl pH 8.0–8.5, 0.5 M EDTA pH 8.0, 0.5% SDS and 0.5 mg Proteinase K) and incubated at 37°C for 24 hrs, followed by phenol/chloroform DNA extraction [96]. Aliquots of crude DNA extracts were analyzed with an Agilent 2100 Bioanalyzer Expert High Sensitivity DNA Assay to quantify and assess the quality of extracted DNA.

The fresh blood samples (3 ml) were collected in EDTA vacutainer tubes. Total genomic DNA was isolated using a standard non-organic method and diluted to obtain a working concentration of 2.5 ng/μl. All blood samples were processed in a post-PCR laboratory.

3.- Sex determination by High Resolution Melting Analysis. The HRM analysis was based on the melting temperature (Tm) difference of the amplified AMELX-allele.
and AMELY-allele fragments (61 bp for the AMELX-allele and 64 bp for the AMELY-allele) of the human amelogenin gene. Fragments were amplified with the LightCycler 480 Real-Time PCR Instrument using the LightCycler 480 High Resolution Melting Master kit (Roche Applied Science), which contains a saturating fluorescent dye (EvaGreen). The PCR reactions were performed by triplicate (unless indicated) in a total volume of 20 μl containing 2 μl of template DNA (5 ng), 3 mM MgCl₂, 1X conc. Lightcycler® Master-Mix [composed by FastStart Taq DNA Polymerase, reaction buffer, dNTP mix (with dTTP instead of dUTP) and High Resolution Melting Dye] and nuclease-free water (QIAGEN), and 0.2 μM of each of the two primers: Amel_F (5'-CCCTTTGAAGTGGTACCAGAG-3') and Amel_R (5'GGGAATARGAACAAATTGTCTAC-3', R=A or G).

The LightCycler 480 Instrument protocol included a pre-incubation step of 10 min at 95°C. The amplification phase comprised 80 cycles of 15 s, 1 m at 56°C, and 30 s at 72°C. After the PCR step, the High-Resolution Melting analysis was performed measuring the drop of fluorescence signal under the following conditions: 1 m at 95°C, 1 m at 40°C and an increase from 60°C to 90°C at a rate of 1°C/s. The instrument is capable of capturing a large number of fluorescent data points per change in temperature with high precision in order to generate a melting curve chart of sample fluorescence versus temperature using the LightCycler 480 software Ver. 1.5 (Roche Applied Science). The analysis also displays the first negative derivative of these sample curves resulting in a new graph where the melting temperature of each sample appears as a characteristic peak. AMELX-allele and AMELY-allele amplicons produced from both modern male and female DNA templates were evaluated as positive controls and three negative controls with no DNA were evaluated in each amplification.

4.- Cross-verification of the High Resolution Melting Analysis by Sanger Sequencing. Cloning of PCR products is useful to assess levels of both contamination and molecular damage in ancient DNA. Also, in our experience, it is not easy to sequence accurately short DNA fragments (<80 bp) by Sanger sequencing, therefore in these cases it is useful to clone the fragments and use vector’s primer-binding sites for sequencing. For these reasons, the AMELX-allele and AMELY-allele amplicons generated from positive controls and from sex-unknown samples were cloned using a TOPO TA Cloning kit (with pCR 2.1-TOPO vector) and TOP10 chemical Competent cells (Invitrogen) following manufacturer’s instructions. A minimum of 30 colonies were selected from each cloned sample and underwent colony PCR using the M13 primers. Then we sequenced 5 clones per sample that contained the transformed vector with the fragment of interest. DNA sequencing was performed in both directions by dideoxy method using the 3730 XL DNA Analyzer (Applied Biosystems) at the Genomic Services Unit in Langebio, Cinvestav, Mexico.

5.- Cross-verification of the High Resolution Melting Analysis by Agilent 2100 Bioanalyzer Expert High Sensitivity DNA Assay. A 2 μl aliquot of AMELX-allele and AMELY-allele amplicons generated from positive controls and sex-unknown samples were analyzed by Agilent 2100 Bioanalyzer Expert High Sensitivity DNA Assay (the Bioanalyzer assay) with the High Sensitivity DNA Marker 35/10380 bp, using the Agilent 2100 Bioanalyzer (Agilent Technologies Inc.) at the Genomic Services Unit in Langebio, Cinvestav, Mexico.

6.- Cross-verification of the High Resolution Melting Analysis by anthropological data. Human remain samples of 4 adult individuals from Teopancazco (burials 2, 78, 102 and 105) were analyzed by conventional osteological measurements for sex determination at the Institute of Anthropological Research, UNAM, Mexico, under the direction of Linda R. Manzanilla. These samples were used as ancient controls.

Results

DNA preservation

Several attempts to determine the sex of these samples with two previously published methods were unsuccessful. These methods targeted amelogenin gene fragments of 106–112 bp [51] and 70–73 bp [83]. This was a first indication of DNA degradation. Consistent with this, the Bioanalyzer assay showed a high DNA fragmentation pattern, with a main distribution of DNA fragments in the range of 40–100 bp, in all ancient samples analyzed (Figure S1). We therefore considered these samples to be heavily degraded and that represented ideal cases to test our HRM method, which targets smaller amelogenine gene fragments.

Sex determination by High Resolution Melting Analysis

This analysis is based on the melting temperature (Tm) difference of the amplified AMELX-allele and AMELY-allele fragments (61 bp for the AMELX-allele and 64 bp for the AMELY-allele) produced by a 3-bp deletion on the AMELX-allele. After software processing, the curves obtained from male and female individuals showed a clear difference in shape (Figure 1), allowing thus sex identification.

The accuracy of the assay was examined by using a set of 6 control samples of known sex. To assess reproducibility, three replicates of each sample were analyzed. No discrepancy was observed between sex determinations by HRM and the previously known sex status in all control samples and their replicas. Regarding sex-unknown archaeological samples, this method was efficient in all the individuals examined in spite of the observed pattern of DNA degradation. However, for one of the individuals only two replicates were done, because the DNA extract was not enough for the three replicates due to its use in previous experiments.

Cross-verification of the High Resolution Melting Analysis

The accuracy of sex determination by HRM analysis was verified by contrasting results with those obtained with other three methods. The first one was the sequencing of 5 positive clones harboring fragments from the AMELX-allele and AMELY-allele amplified from samples of known sex. The sequences obtained demonstrated the 3-bp insertion on the AMELX-allele in the male samples (Figure 2).

A second confirmation of the HRM analysis was conducted by a Bioanalyzer assay, in which AMELX-allele and AMELY-allele amplicons generated by Real-Time PCR from positive controls and from sex-unknown samples were analyzed. The results showed a differential pattern in terms of amplicons length between male and female individuals (Figure 3).

The third cross-verification was conducted by contrasting the HRM data with previously obtained sex determination by osteological measurements in 4 adult individuals recovered from Teopancazco, Teotihuacan (burials 2, 78, 102 and 105). The sex determination by HRM analysis matched with the sex determined by osteological measurements in all the samples analyzed.
HRM analysis starts with a PCR amplification of the amelogenin gene. The amplification is in the presence of EvaGreen, a saturating dye which presents high fluorescence when bound to dsDNA. Amplification is followed by a High Resolution Melting analysis (HRM) in the LightCycler 480 Real-Time PCR Instrument (Roche Applied Science). A dissociation curve analysis displays the different melting temperature of products from the X and Y chromosome.

doi:10.1371/journal.pone.0104629.g001

Figure 2. Confirmation of the HRM analysis by Sanger Sequencing. AMELX-allele and AMELY-allele fragment (61 and 64 bp, respectively) of positive controls were amplified by real-time PCR, cloned, and sequenced. The sequences obtained demonstrate the 3-bp deletion on the AMELX-allele.

doi:10.1371/journal.pone.0104629.g002
Sex determination in highly degraded samples from human remains has been a challenge in the anthropological and forensic fields. Traditionally, it has been based on the visual examination of the sexual dimorphism present in the majority of human bones and dentition [1–45]. Nevertheless, these kinds of methods are not appropriate if the skeletons are sub-adult, incomplete or highly damaged [46–48]. For this reason, in the last years a large number of methods for sex determination based on molecular biology have been proposed [48–57]. The most widely used is based on the human amelogenin gene (AMEL) [63–72] because this analysis can distinguish AMEL alleles from each sexual chromosome due to genetic differences between them, as described previously by Haas-Rochholz and Weiler [73].

At the moment, different primer sets to amplify the amelogenin gene have been reported [49,51,52,63,64,67,68,74–76]. However, the length of the fragments amplified is relatively large which causes this method to be unsuitable for heavily degraded samples. A targeted DNA fragment size of 100 bp might be too long for a successful and reliable analysis in some cases [97,98]; especially considering this is nuclear DNA, more prone to degradation as compared to mitochondrial DNA.

In our study, the ancient samples showed a high DNA fragmentation pattern (between 40–100 bp) as assessed by a Bioanalyzer assay, indicating a general poor DNA preservation. As a result, primer sets published previously [51] were not efficient for sex determination in these samples (data not shown). This prompted us to devise a new method to sex identification analyzing smaller amplicons, suitable for heavily degraded samples. There are other published methods that involve smaller amplicons than ours, but they also involve post-PCR process, such as electrophoresis, Denaturing High Performance Liquid Chromatography (DHIPLC) [78], or pyrosequencing [79], which can cause crossing-over contamination, especially in ancient samples.

Here we propose a new method based on the melting-curve analysis reported firstly by Ririe et al., [80]. This method is more adequate for samples with extremely poor preservation, as it is more sensitive, while at the same time it avoids crossing-over contaminations, by not requiring any processing after real-time PCR as in other previously published methods [78,79]. Furthermore, it allows detecting in a small amplified DNA fragment, the 3 bp deletion in the AMEL-X allele.

The use of EvaGreen in our method is relevant because it is stable both under PCR conditions and during routine storage and handling. Furthermore, no PCR interference is produced when used at low concentration, thus permitting a more robust PCR signal as well as a sharper and stronger DNA melt peak in melting-curve analyses [96,99]. In fact, the melting-curve obtained by our method shows a clear difference in shape between male (2 peaks) and female individuals (single peak) (Figure 1).

Ancient or degraded samples are prone to the allele drop-out phenomenon, consisting in amplification failure of only one of the two alleles in a locus, mainly the largest one [100], indicating an inverse relationship between amplicon fragment and amplification failure. Therefore, a reduction of allele drop-out frequency could be expected by targeting smaller fragment sizes. On the other hand, to circumvent the problem it has been recommended to analyze several replicas of the same sample [92]. Therefore, we conducted the analysis by triplicate in all samples but one. Results from the three replicas were successfully obtained in 16 samples out of 17 samples analyzed. The exception was a sample for which DNA was finished at the second replica. However a consistent result was obtained from the two replicas for this sample.

Our method was cross-verified by three analyses. HRM results and those obtained by (i) sequencing several cloned amplicons from the AMELX-allele and AMELY-allele, (ii) a Bioanalyzer assay, and (iii) previous anthropological data [101], were all concordant.

**Figure 3.** Confirmation of the HRM analysis by a Bioanalyzer assay. AMELX-allele and AMELY-allele fragment (61 and 64 bp, respectively) of positive controls were amplified by real-time PCR and then analyzed by the Agilent 2100 Bioanalyzer Expert High Sensitivity DNA Assay. The results show a differential pattern in terms of amplicons length. (A) Male positive control. (B) Female positive control. doi:10.1371/journal.pone.0104629.g003
This HRM method was efficient to determine the sex of 16 individuals of unknown sex recovered from Teopancazco, Troitihuacan. Fourteen of them were infants and three samples were from adult individuals with sex reported as no consistent by anthropological data. This illustrates the usefulness of this method to sex identification in samples in which other previously published molecular methods [51,53] failed (see DNA preservation section above).

A variety of methodologies to sex determination have been reported previously, but are not sensitive enough on samples presenting extensive degradation of genetic material, as is the case of Mesoamerican populations subjected to warmer conditions in relation to more northern ancient populations. Our method demonstrated to be efficient, rapid and sensitive enough for sex determination in human remains showing a general poor DNA preservation. Finally, the method can be used in a high-throughput manner to analyze a relatively large number of samples in one run, at a relatively low cost. For example, including replicates and controls, 21 samples can be analyzed in a 96-well PCR plate and 04 samples in a 384-well PCR plate.

Supporting Information

Figure S1 DNA fragmentation pattern in a sample recovered from Teopancazco. Main distribution of DNA fragments is in the range of 40–100 bp. (TIF)

Acknowledgments

We thank Hilda E. Ramos-Aboites and Christian E. Martinez-Guerrero for technical support and Marcus Moore for language editing of the paper. This work was supported by Conacyt Nacional de Ciencia y Tecnologia (CONACYT; www.conacyt.gob.mx) grant No. CR-2008-01-105411 to Dr. Rafael Montiel, M.Sc. Alvarez-Sandoval is supported by a CONACYT fellowship (Reg. No. 300750). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Author Contributions

Conceived and designed the experiments: BAA-S. Performed the experiments: BAA-S. Analyzed the data: BAA-S LRM RM. Contributed reagents/materials/analysis tools: LRM RM. Wrote the paper: BAA-S RM.

References

1. Rogers R, Saunders S (1994) Accuracy of Sex Determination Using Morphological Traits of the Human Pelvis. J Forensic Sci 39(4): 1047–1056.
2. Kalnem J, Rathburn TA (1996) Sex determination by discriminant function analysis of the parietal portion of the temporal bone. J Forensic Sci 41: 865–867.
3. Burris BG, Harris EF (1998) Identification of race and sex from palate dimensions. J Forensic Sci 43: 959–965.
4. Albanese JA (2003) Metric method for sex determination using the hipbone and the femur. J Forensic Sci 48(2): 263–273.
5. Teke HY, Duran S, Canturk N, Canturk G (2007) Determination of gender by measuring the size of the maxillary sinuses in computerized tomography scans. Surg Radiol Anat 29(1): 9–13.
6. Gapert R, Black S, Last J (2009) Sex determination from the foramen magnum: discriminant function analysis in an eighteen and nineteenth century British sample. Int J Legal Med 123: 25–33.
7. Francesinquí Junior L, Francesinquí MA, De La Cruz BM, Pereira SD, Ambrosano GM, et al. (2007) Identification of sex using cranial base measurements. J Forensic Odontostomatol 25(1): 7–11.
8. May H, Peled N, Dar G, Cohen H, Abjab J, et al. (2011) Hypothesis from frontal Acanthoc criminala: criteria for sexing and aging a skeleton. Int J Legal Med 125: 669–673.
9. Spradley MK, Junta RL (2011) Sex estimation in forensic anthropology: skull versus postcranial elements. J Forensic Sci 56: 289–296.
10. Wescott DJ, Moore-Jansen PH (2011) Metric variation in the human occipital bone: forensic anthropological applications. J Forensic Sci 46: 1159–1163.
11. Garamond R, Rissech C (2007) Determination of sex from juvenile crania by means of discriminant function analysis. J Forensic Sci 57: 24–34.
12. Krishan K, Kanchan T, Passi N, DiMaggio J (2012) Heel-Ball (HB) index-12. Krishan K, Kanchan T, Passi N, DiMaggio J (2012) Heel-Ball (HB) index examination of the relationship between the HB index and sex. J Forensic Sci 57: 172–173.
13. Stein M, Becker PJ, L’Abbe EN, Scholtz Y, Myburgh J (2011) Analysis of the repeatability of pubic and ischial measurements. Forensic Sci Int 214(1–3): 210.e1–4.
14. Uhman AT, Al-Rawi NH, Al-Timimi JF (2012) Evaluation of foramen magnum in gender determination using helical CT scanning. Dentomaxillofac Radiol 41: 197–202.
15. Kanchan J, Gupta A, Krishan K (2013) Estimation of sex from mastoid triangle Acraniform diagnostic analysis. Forensic Sci Int Leg Med 20(7): 853–856.
16. Case DT, Ross AH (2007) Sex determination from hand and foot bone lengths. J Forensic Sci 52(2): 264–270.
17. Harris SM, Case DT (2012) Sexual dimorphism in the tarsal bones: implications for sex from Japan. J Forensic Sci 57: 295–303.
18. Scholtz Y, Steyn M, Pretorius E (2010) A geometric morphometric study into the sexual dimorphism of the human scapula. Homo 61(4): 235–270.
19. Dabhis GR, Moore-Jansen PH (2010) A method for estimating sex using metric analysis of the scapula. Forensic Sci Int Leg Med 20(1): 148–151.
20. Berrizbeitia EL (1989) Sex determination with the head of the radius. J Forensic Sci 34: 1206–1213.
21. Aye VO (2010) Determination of sex by arm bone dimensions. Forensic Sci Int 201(1–3): 111.e1–3.
22. Mall G, Hubig M, Buttner A, Kuznik J, Penning R, et al. (2001) Sex Determination and Estimation of Stature from the Long Bones of the Arm. Forensic Sci Int 117: 25–30.
23. Purkait R (2005) Triangle identified at the proximal end of femur: a new determinant. Forensic Sci Int 147(2–3): 135–139.
24. Rieseck G, Schafer M, Malgosia A (2000) Development of the femur e determinant in age determination. J Forensic Leg Med 18(3): 1–9.
25. Rogers TL (2009) Sex determination of adolescent skeletons using the distal humerus. Am J Phys Anthrop 140(1): 143–148.
26. Gonçalves D, Thompson TJU, Cunha E (2013) Osteometric sex determination on burned human skeletal remains. J Forensic Leg Med 20(7): 906–911.
27. Introna F, De Vella G, Campobasso CP (1999) Sex Determination by Discriminant Function Analysis of Patella Measurements. Forensic Sci Int 95(1): 39–45.
28. Ramadã SU, Turkmen N, Dolgun NA, Gökharmın D, Menezesc RG, et al. (2010) Sex determination from measurements of the sternum and fourth rib using mulitcalc computed tomography of the chest. Forensic Sci Int 197(1–3): 120.e1–5.
29. Bongiovanni R, Spradley MK (2012) Estimating sex of the human skeleton based on metrics of the sternum. Forensic Sci Int 219: 290.e1–7.
30. Kindschuh SC, Dupras TL, Cowgill LW (2010) Determination of sex from the hyoid bone. Am J Phys Anthropol 143(2): 279–284.
31. Rogers NL, Rusenoy L, McCormick WF (2009) The rhomboid fossa of the clavicle as a sex and age estimator. J Forensic Sci 45(1): 61–67.
32. Graw M, Waid J, Ahlbrecht M (2005) Course of the metacristatus internus as a criterion for sex differentiation. Forensic Sci Int 147: 113–117.
33. Masotti S, Scuci-Leonelli E, Gualdi-Russo E (2013) Cremated human remains: is measurement of the lateral angle of the metacristatus internus a reliable method of sex determination? Int J Leg Med 127(5): 1039–1044.
34. Macaluso PJ (2011) Investigation on the utility of permanent maxillary molar cusps area for sex estimation. Forensic Sci Med Pathol 7: 233–247.
35. Lain R, Taylor J, Cooker S, Craig P, Graham J (2009) Comparative dental anatomy in disaster victim identification: lessons from the 2009 Victorian Bushfires. Forensic Sci Int 205: 36–39.
36. Ozden H, Balci Y, Demiray C, Turgut A, Ertugrul M (2005) Stature and sex estimation using foot and shoe dimensions. Forensic Sci Int 147: 181–184.
37. Mudgill R, Kaur R, Menezes RG, Kanchan T, Garg RK (2008) Foot index: is it a tool for sex determination? J Forensic Leg Med 15(4): 223–226.
38. Zeybek G, Ergur I, Demiray C (2008) Stature and gender estimation using foot measurements. Forensic Sci Int 181(1–3): 54.e1–5.
39. Amatundur D (2010) Estimation of sex from the dimensions of foot, footprints, and shoe. Anthropoz 41(3): 21–29.
40. Jowaheer V, Agnihotri AK (2011) Sex identification on the basis of hand and foot measurements in Indo-Mauritian population: a model based approach. J Forensic Leg Med 17(4): 173–176.
41. Kanchan T, Krishan K (2011) Anthropometry of hand in sex determination of dismembered remains - a review of literature. J Forensic Leg Med 18: 14–17.
42. Kanchan T, Kumar GP, Menezes RG (2008) Index and ring finger ratio - a new sex determinant in south Indian population. Forensic Sci Int 181(1–3): 53.e1–4.
43. Kanchan T, Kumar GP (2010) Index and ring finger ratio - a morphologic sex determinant in South Indian children. Forensic Sci Med Pathol 6: 255–260.
45. Krishan K, Kanchan T, Asha N, Kaur S, Chatterjee PM, et al. (2013) Estimation of sex from index and ring finger in a North Indian population. J Forensic Leg Med 20(5): 471–479.

46. Rodríguez JV (1994) Introducción a la Antropología forense. Análisis e identificación de restos óseos humanos. Anaconda ed. Bogotá, Colombia, 139 p.

47. Schmidt D, Hummel S, Herrmann B (2003) Brief Communication: Multiplex X/Y-PCR Improves Sex Identification in aDNA Analysis. Am J Phys Anthropol 121(4): 337–341.

48. Oskarsson E, Andréfouët S, Humphrey L, Goétherström A (2011) Further developments in molecular sex assignment: a blind test of 18th and 19th century human skeletons. J Archaeol Sci 38(6): 1326–1330.

49. Akane A, Shiono H, Matsubara K, Nakahori Y, Seki S, et al. (1991) Sex identification of forensic specimens by polymerase chain reaction (PCR): two alternative methods. Forensic Sci Int 49(1): 81–86.

50. Hummel S, Herrmann B (1991) Y-Chromosome-Specific DNA amplified in ancient human bone. Naturwissenschaften 78: 266–267.

51. Sullivan KM, Mannucci A, Kimpton CP, Gill P (1993) A rapid and quantitative DNA test: fluorescence-based PCR analysis of X-Y homologous gene amelogenin. Biotechniques. 15: 636–641.

52. Faerman M, Filon D, Kahila G, Greenblatt C, Smith P, et al. (1995) Sex identification of archaeological human remains based on amplification of the X and Y amelogenin alleles. Gene 167(1–2): 327–332.

53. Stone AC, Milner GR, Pabio S, Stonencking M (1996) Sex determination of ancient human skeletons using DNA. Am J Phys Anthropol 99: 231–238.

54. Goétherström A, Liden K, Ahlstrom T, Kallersjo M, Brown TA (1997) Osteology, DNA and sex identification: morphological and molecular identifications of five neolithic individuals from Ajvide, Gotland. Int J Osteoarchaeol 7: 71–81.

55. Lassen C, Hummel S, Herrmann B (2000) Molecular sex identification of stillborn and neonate individuals (“Traufkinder”) from the burial site Aegerten in the 11th century AD. J Archaeol Sci 37(12): 1131–1140.

56. Meyer E, Wiese M, Bruchhaus H, Claussen M, Klein A (2000) Extraction and amplification of authentic DNA from ancient human Remains. Forensic Sci Int 112: 87–90.

57. Gibbon V, Paximidas M, Srkij G, Ruff P, Penny C (2009) Novel methods of molecular sex identification from skeletal tissue using the amelogenin gene. Forensic Sci Int Genet 3: 74–79.

58. Pilić E, Molić A, Serpico C, Achilli A, Lancioni H, et al. (2013) Monitoring and Y amelogenin alleles. Gene 167(1–2): 327–332.

59. Taberlet P, Friffin S, Goossens B, Questiau S, Manceau V, et al. (1996) Identification of neolithic human skeletons. Anthropol Anthropol 58(1): 1–8.

60. Skoglund P, Storå J, Goétherström A, Jakobsson M (2013) Accurate sex identification of ancient human remains using DNA shotgun sequencing. J Archaeol Sci 40(12): 4477–4482 doi:10.1016/j.jas.2013.07.004.

61. Nakahori Y, Hamano K, Iwaya M, Nakagome Y (1991) Sex identification by molecular analysis of DNA extracted from ancient human bone. Naturwissenschaften 78: 266–267.

62. Velarde-Félix JS, Molina-Benítez CE, Soloárez-Rosas C, Cázares-Salazar SG, Rendón-Aguilar H, et al. (2008) Identifi cación del sexo mediante análisis de ADN procedente de huesos antiguos. Rev Mex Patol Clin 55(1): 17–20.

63. Friffin S, Goossens B, Questiau S, Manceau V, et al. (1996) Reliable genotyping of samples with very low DNA quantities using PCR. Nucleic Acids Res 24: 3109–3114.

64. Chang YM, Burgoyne LA, Both K (2003) Higher failures of amelogenin sex test in an Indian population group. J Forensic Sci 48: 1309–1313.

65. Tschentscher F, Frey U, Bajnaiwosti T (2000) Amelogenin sex determination by pyrosequencing of short PCR products. Int J Legal Med 122: 333–353.

66. Langenker RE, Mies RB, Wittwer CT (1997) Continuous fluorescence monitoring of rapid cycle DNA amplification. Biotechniques 22: 130–131.

67. Witteveen CT, Herrmann MG, Moss AA, Rasmussen RP (1997) Continuous fluorescence monitoring of rapid cycle DNA amplification. Biotechniques 22: 130–131.

68. Alonso A, Martín P, Albarcán C, García P, Primorac D, et al. (2003) Specific quantification of human genomes from low copy number DNA samples in forensic and ancient DNA studies. Croat Med J 44(5): 273–280.

69. Santos FR, Pandya A, Tyler-Smith C (2003) Demonstration of preferential binding of SYBR Green I to specific DNA fragments in real-time multiplex PCR. Nucleic Acids Res 31(22): 136.

70. Geschke J, Amy J, Martin J, Hooijer H, Smits M, et al. (2006) Multiplexing of ancient DNA using real-time PCR: preliminary results. Sex Determination in Highly Fragmented Human DNA

71. Goëtherström A, Liden K, Ahlstrom T, Kallersjo M, Brown TA (1997) Osteology, DNA and sex identification: morphological and molecular identifications of five neolithic individuals from Ajvide, Gotland. Int J Osteoarchaeol 7: 71–81.

72. Lassen C, Hummel S, Herrmann B (2000) Molecular sex identification of stillborn and neonate individuals (“Traufkinder”) from the burial site Aegerten in the 11th century AD. J Archaeol Sci 37(12): 1131–1140.

73. Skoglund P, Storå J, Goétherström A, Jakobsson M (2013) Accurate sex identification of ancient human remains using DNA shotgun sequencing. J Archaeol Sci 40(12): 4477–4482 doi:10.1016/j.jas.2013.07.004.

74. Nakahori Y, Hamano K, Iwaya M, Nakagome Y (1991) Sex identification by polymerase chain reaction using X-Y homologous primer. Am J Med Genet 49(4): 472–473.

75. Nakahori Y, Takenaka O, Nakagome Y (1991) A human X-Y homologous region encodes amelogenin. Genomics 9: 264–269.

76. Lau EC, Mohandas TK, Shapiro LJ, Slavkin HC, Snead ML (1989) Human and mouse amelogenin loci are the sex chromosomes. Genomics 4: 162–168.

77. Taberlet P, Friffin S, Goossens B, Questiau S, Manceau V, et al. (1996) Reliable genotyping of samples with very low DNA quantities using PCR. Nucleic Acids Res 31(22): 136.

78. Giglio S, Simon PT, Saint CP (2003) Demonstration of preferential binding of SYBR Green I to specific DNA fragments in real-time multiplex PCR. Nucleic Acids Res 31(22): 136.

79. Tschentscher F, Frey U, Bajnaiwosti T (2000) Amelogenin sex determination by pyrosequencing of short PCR products. Int J Legal Med 122: 333–353.

80. Ririe KM, Rasmussen RP, Wittwer CT (1997) Product differentiation by analysis of DNA melting curves during the polymerase chain reaction. Anal Biochem 245: 154–160.

81. Witteveen CT, Herrmann MG, Moss AA, Rasmussen RP (1997) Continuous fluorescence monitoring of rapid cycle DNA amplification. Biotechniques 22: 130–131.

82. Witteveen CT, Herrmann MG, Moss AA, Rasmussen RP (1997) Continuous fluorescence monitoring of rapid cycle DNA amplification. Biotechniques 22: 130–131.

83. Andreasson H, Allen M (2003) Rapid quantification and sex determination of ancient human bone. Naturwissenschaften 80: 687–691.

84. Alonso A, Martín P, Albarcán C, García P, Primorac D, et al. (2003) Specific quantification of human genomes from low copy number DNA samples in forensic and ancient DNA studies. Croat Med J 44(5): 273–280.

85. Gagneux P, Boesch C, Woodruff DS (1997) Microsatellite scoring errors associated with nonmicrosat scoring genotyping based on nuclear DNA amplified from shed hair. Mol Biol 6: 861–863.

86. Santos FR, Pandya A, Tyler-Smith C (1998) Reliability of DNA-based sex tests. Nat Genet 18: 103.

87. Lattanzi W, D'Aciamo MC, Lenato GM, Chini G, Vaglio G, et al. (2000) A large intersitial deletion encompassing the amelogenin gene on the short arm of the Y chromosome. Hum Mol Genet 9(7): 1201–1207.

88. Taberlet P, Friffin S, Goossens B, Questiau S, Manceau V, et al. (1996) Reliable genotyping of samples with very low DNA quantities using PCR. Nucleic Acids Res 34(19): 5189–5194.