Introduction

The Chotěbuz-Podobora hillfort lies 5 km northwest of the town of Český Těšín. The hillfort is situated on featureless plateaus, which were formed by the fluvial sediments of the main terrace of the Olše River and then covered by loess loams. The terrace hangs over the concave bank of the Olše River.

The hillfort was built by Lusatian Culture people (10th to 5/4th century BC). The Slavonic settlement is dated from the 8th century to the first part of the 11th century AD (Kouřil 1994). The people who lived there controlled a site of strategic importance: the exit through the Moravian Gate, the main European connection between North and South (Kouřil 1997, 2005). Other evidence of a local elite presence at the Chotěbuz-Podobora hillfort at that time is the finding of log house remains and paling structures on the acropolis as well as the finds of militaria (axes, spears, and arrow-heads) and riding equipment (spurs with hooks and discs, stirrups, bits, and buckles). The spurs in particular can be considered to be a significant emblem of the Slavonic mounted units, as a statutory symbol and feature of free men and warriors supporting the local leader. In addition to these militia, an abundance of jewellery was also found (silver-plated earrings, rings and beads). The finding of two limb bones of a greyhound-like dog at the acropolis is also associated with the aforementioned features.

The bones of the greyhound-like dog were located together with other osteological material originating from the dense cultural layer found in the probe pit S-44 at the acropolis at a depth of 100–145 cm below the surface (Nývltová Fišáková 2007). This layer relates to the older phase of Slavonic settlement which is archaeologically dated to the second half of the 8th century and 9th century (Kouřil 1994). The discovered skeletal remains of the greyhound-like dog were dated at 830 ± 49 AD cal. (AMS radiocarbon dating Poz-20508; calibration using IntCal13 dataset – Reimer et al. 2013). This age clearly correlates with the aforementioned archaeological assumptions. The test pit was located in the northernmost part of the acropolis, in the best-protected section of the whole fortification. The limb bones are long, slim, and flattened front-to-back (Text-fig. 1). The dog’s height is calculated as 70 cm according to Harcourt’s (1974) method (Nývltová Fišáková 2007, 2010).
the maximum length and the minimum width falls into the variation width of the ratio for greyhounds, mainly Polish greyhounds (Nývltová Fišáková 2007, 2010, fig. 8).

The greyhound has been considered an exceptional animal since the very earliest times (Sullivan 1999, Morris 2001, Branigan 2004). During the Middle Ages these dogs could only have been owned by nobles and rulers (Kholová 1987, Stuchlík and Cisařovský 1992, Sullivan 1999, Branigan 2004, Cisařovský 2008). The greyhound was one of the heraldic symbols of kings and nobles in both England and France (Sullivan 1999, Branigan 2004), as well as in Czech lands (Mysliveček 1993, Buben 2003). This is the oldest evidence of a greyhound in the Czech Republic. Until this discovery, the oldest finding was the mummified English greyhound from the castle in Ždánice (Nývltová Fišáková, 2007).

The main aim of this study was to analyze 17 excavated bones to test the hypothesis that these were the oldest greyhound remains within the Czech territory. We chose to analyse mitochondrial DNA (mtDNA) because of its special features: it is situated the outside nucleus in mitochondria, there are many copies while there are usually only two copies of each chromosome. Circular mtDNA is also more stable than linear nDNA, thus increasing the chance to obtain analyzable results even from degraded (ancient) DNA.

The first published complete mitochondrial DNA (mtDNA) genome of the domestic dog (Canis familiaris) contained 16,727 bp (base pairs) with the control region (CR) spanning positions 15,458–16,727 (1,270 bp) (Kim et al. 1998). While the dog mtDNA genome closely resembles the mtDNA genome of other mammals, the dog (and other related canids) mtDNA CR differs due to the presence of a 10 bp repeating unit (5´-GTACACGT(A/G)C-3´) that begins at base 16,130 and varies in number and sequence both within and among individuals. The CR of dog mtDNA, like that of human mtDNA, has been the subject of a number of studies investigating variation among individuals (Vila et al. 1997, Kim et al. 1998, Webb and Allard 2009). Previous studies revealed more than 100 single nucleotide polymorphisms (SNPs) within the mtDNA CR of the domestic dog.

Materials and methods

The ancient material was found at Chotěbuz-Podobora site. These bones were osteologically very similar to greyhound-like dogs – long and generally slim in comparison with other dogs of a similar height (Text-fig. 1).

For comparison with recent domestic dog, we took a buccal swab from a greyhound whose ancestors were born in 1840 in Great Britain. We also took swabs from 15 recent dogs (greyhound-like dogs) from the dog exhibition in Brno 2010 as suitably representative samples for verification of our results. Sample 1 is Romanov z Palatinu Morava (male), sample 2 is Body z Murky (male) (both are Russian barsoi). Sample 3 is Limia Lykon (female, afghan greyhound). Sample 4 is Glenoak Junah (female, afghan greyhound). Sample 5 and 6 are Lajjina and Pussy Bohemia Genao (both Sloughi). Sample 7 is Arabian Sloughi Djawida EJJI Fatira of Love ARAB BEAUTY. MAYSUN Boehmia Genao is sample 8, whose grand-father was a Moroccan Sloughi. Sample 9 Takellaut N8FA BADAKA is a Sloughi-Azawakh, whose ancestor was from the Sahara. Sample 10 is a Polish greyhound Baldwin-Xanthusia (male). Uncle Teddy (male) and Hellega (female) are whippets, samples 11 and 12. Sample 13 is deerhound Greyrory’s Draeura (female). Rendel (male) and Rosarka (female) are samples 14 and 15 are also barsois.

Isolation of DNA from ancient bone

Forensic DNA isolation methods were used which are a combination of the isolating methods described in the publications of Baron et al. (1996) and Yang et al. (1997). First we isolated DNA from the bone. The surfaces of both small bone fragments were mechanically cleaned with sandpaper and 96% ethanol was used to remove foreign substances, such as bacteria, fungi, or other foreign DNA. Usage of a more efficient bleaching liquid could be severely damaging for old bone. The next step of preparation for isolation of DNA was cutting the bone into thin slices with
Isolation of DNA from buccal swabs of recent greyhounds

DNA from the buccal swabs of 15 recent dogs was isolated with a 5% suspension of Chelex (Walsh et al. 1991).

Amplification of mtDNA from isolates of bone and buccal swabs

Oligonucleotide primers used for amplification and sequencing of the canid mtDNA CR were created using a dog reference sequence (GenBank accession no. U96639). Convention nomenclature was used: the primer name indicates the position of the 5’ base. Forward primers were F15.719 (5’-GTAATGTCTCCTTCTCGCTT-3’) and F16.431 (5’-CACGCGCGTAAGACATTAAG-3’). Reverse primers were R16.114 (5’-CCTGAACCATTGACTGAATAG-3’) and R42 (5’-GGCATTTTCAGTGCCTTGCTT-3’) (Gundry et al. 2007, see Text-fig. 2 for the position of primers). We amplified DNA from two parallel isolates of the same bone and we obtained two identical 395 bp sequences with primer pair A (F15.719 and R16.114) and two identical 338 bp sequences with primer pair B (F16.431 and R42) (Text-fig. 3). We amplified DNA from two parallel isolates of the same bone and we obtained two identical 395 bp sequences with primer pair A (F15.719 and R16.114) and two identical 338 bp sequences with primer pair B (F16.431 and R42) (Text-fig. 3). We amplified DNA from two parallel isolates of the same bone and we obtained two identical 395 bp sequences with primer pair A (F15.719 and R16.114) and two identical 338 bp sequences with primer pair B (F16.431 and R42) (Text-fig. 3). We amplified DNA from two parallel isolates of the same bone and we obtained two identical 395 bp sequences with primer pair A (F15.719 and R16.114) and two identical 338 bp sequences with primer pair B (F16.431 and R42) (Text-fig. 3). We amplified DNA from two parallel isolates of the same bone and we obtained two identical 395 bp sequences with primer pair A (F15.719 and R16.114) and two identical 338 bp sequences with primer pair B (F16.431 and R42) (Text-fig. 3). We amplified DNA from two parallel isolates of the same bone and we obtained two identical 395 bp sequences with primer pair A (F15.719 and R16.114) and two identical 338 bp sequences with primer pair B (F16.431 and R42) (Text-fig. 3). We amplified DNA from two parallel isolates of the same bone and we obtained two identical 395 bp sequences with primer pair A (F15.719 and R16.114) and two identical 338 bp sequences with primer pair B (F16.431 and R42) (Text-fig. 3).

Sequencing of mtDNA

Cycle sequencing was performed using a Terminator Ready Reaction Mix (Applied Biosystems) and BigDye Sequencing Buffer (Applied Biosystems) according to the manufacturer’s instructions. The sequencing primers were the same as those listed above. The sequencing reaction (test solution containing 7 μl of the amplified DNA, 3.5 μl primer [at 1 mM] and 9.5 μl Ready Reaction mix) was carried out in a GeneAmp 9700 PCR System thermal cycler and consisted of denaturation for 1 min. at 96 °C, followed by 25 cycles of 10 sec. at 96 °C, 5 sec. at 60 °C, and 4 min. at 60 °C, followed by a holding period at 12 °C until the next step.

Results

We amplified DNA from two parallel isolates of the same bone and we obtained two identical 395 bp sequences with primer pair A (F15.719 and R16.114) and two identical 338 bp sequences with primer pair B (F16.431 and R42) (Text-fig. 3). The blank extraction control was negative. To verify that we had worked with ancient DNA we used primer pairs: F15.719, R16.169 (450 bp), R16.269 (550 bp) and R16.519 (800 bp) that did not yield any amplicon, which is considered a signature of degraded, presumably ancient DNA (Text-fig. 4). We compared the obtained sequences with the corresponding sequences of recent dogs published by Gundry et al. (2007).

In the next step we compared our sequences from ancient bone and sequences of a recent greyhound published by Gundry et al. (2007). The first sequences were obtained by primer pair A (Text-fig. 4) the second sequences by primer pair B (Text-fig. 5).
Text-fig. 3. Electrophoresis after amplification: Electrophoretical analysis of mitochondrial DNA from two parallel bone samples. In the upper part of the figure, are mtDNA sequences amplified by primer pair A (F15.412 and R16.114 (395 bp)), and in the lower part by primer pair B (F16.431 and R42 (338 bp)). Lane 1 and 3 – undiluted samples, 2 and 4 – 10x diluted samples, NC – negative control, L – 100 bp DNA ladder (band size from 100 bp to 1500 bp).

| Recent 1 | CTTATTTACTCCAATCCATTATCATCGACAATGCCACATCTCGACATGAATGACT | 60 |
| Bone 1   | CTTATTTACTCCAATCCATTATCATCGACAATGCCACATCTCGACATGAATGACT | 60 |
| Recent 61 | AATCAGCCCAAGATCCACACACTGTGCTGTCATGACTTTCTTTATTTTTAg | 120 |
| Bone 61  | AATCAGCCCAAGATCCACACACTGTGCTGTCATGACTTTCTTTATTTTTAg | 120 |
| Recent 121 | ggggggAATCTGCTATCACTCATCTACTGACAGGGCAACGGCAACTTTAATCTTATCTTCTT | 180 |
| Bone 121 | GGGGGGAAATCTGCTATCACTCATCTACTGACAGGGCAACGGCAACTTTAATCTTATCTTCTT | 180 |
| Recent 181 | GCCTCAGGAAATGTCAGCTGGGGTCTGATGACTTAAATTCCTCTTGACTGAGCTGTTT | 240 |
| Bone 181 | GCCTCAGGAAATGTCAGCTGGGGTCTGATGACTTAAATTCCTCTTGACTGAGCTGTTT | 240 |
| Recent 241 | ATCCATTATCTATATGATCAGCATAAAATTCAAGG | 278 |
| Bone 241 | ATCCATTATCTATATGATCAGCATAAAATTCAAGG | 278 |

Text-fig. 4. Electrophoresis after amplification: Electrophorical analysis of mitochondrial DNA. mtDNA sequences were amplified by primers F15.412 and R16.169 (450 bp), R16.269 (550 bp), R16.519 (800 bp). Lane 1 are primers F15.412 + R16.169, lane 2 primers F15.412 + R16.269, lane 3 primers F15.412 + R16.519. NC – negative control – water, L – 100 bp DNA ladder (band size from 100 bp to 1500 bp).

Text-fig. 5. Multiple sequence alignment of mtDNA from ancient bone and recent greyhound, (Gundry et al. 2007) primer pair A – F15.719 and R16.114.
greyhounds. As a reference sequence the mtDNA from bone compared with the mtDNA of 15 recent greyhounds in the Czech Republic is used.

Discussion
The evidence of a greyhound in the Slavonic and later in the Great Moravian period within the Czech territory is extraordinary, and it represents the first of this kind ever found in the Czech lands.

We authenticated the ancient DNA using the possibility to amplify long mtDNA fragments (with a length of 450 bp, 550 bp and 800 bp) (Herrmann and Hummel 1994, Hummel 2003). By comparing the sequences from the ancient bone and from the buccal swab with the help of BLAST (Basic Local Alignment Search Tool), we confirmed that the

Text-fig. 6. Comparison of overlapping sequences of mtDNA from recent greyhound and mtDNA from ancient bone – primer pair B – F16.431 and R42.

Text-fig. 7. Comparison of overlapping sequences of mtDNA from mtDNA from BLAST and mtDNA from ancient bone – primer R42 (query is mtDNA from BLAST and subject is mtDNA from bone), position 16,480 is colour-coded.

greyhounds. As a reference sequence the mtDNA from bone compared with the mtDNA of 15 recent greyhounds in the Czech Republic is used.

Table 1. Comparison of mtDNA sequences from recent greyhounds and mtDNA from bone with primers F15.719 and R16.114. Mutations are described in the upper line, samples are marked as 1 to 15.
sequences of mitochondrial DNA came from a dog (99% identical). By comparing our data with Gundry et al.’s (2007), we confirmed that our PCR amplicons with primer pairs A and B are located on canine mtDNA.

The sequences from both the recent greyhound from N. Ireland and the ancient bone obtained by using PCR with primer pair A were identical. However, the sequences amplified with primer pair B contained four deletions and one substitution (Text-fig. 6). We suggest that the first two deletions are at positions 184 and 185, the third at 203 and the fourth at position 204 in the recent dog sequence in comparison with the ancient bone sequence. One substitution is at position 194 where nucleotide C is replaced by T. These differences probably arose during 11 centuries of dog breeding (Herrmann and Hummel 1994).

We compared our sequences with the reference database presented by Gundry et al. (2007) which contains dog mtDNA CR sequences, variable SNPs, and haplotypes of 125 American domestic dog breeds and three wild canids. In particular we compared the SNPs at positions which are

### Table 1. continued

| Primer pair | F15719 | R16114 | CUS | ASC |
|-------------|--------|--------|-----|-----|
| A           | AGGAGTAA | TCTCAAAA | TACTTGGT | CTTGATAT |
| B           | AGGAGTAA | TCTCAAAA | TACTTGGT | CTTGATAT |
| C           | AGGAGTAA | TCTCAAAA | TACTTGGT | CTTGATAT |
| D           | AGGAGTAA | TCTCAAAA | TACTTGGT | CTTGATAT |
| E           | AGGAGTAA | TCTCAAAA | TACTTGGT | CTTGATAT |
| F           | AGGAGTAA | TCTCAAAA | TACTTGGT | CTTGATAT |

### Table 2. Comparison of mtDNA sequences from recent greyhounds and mtDNA from bone with primers F16.431 and R42, mutations are described in the upper line, samples are marked as 1 to 15.

| Primer pair | F15719 | R16114 | CUS | ASC |
|-------------|--------|--------|-----|-----|
| 1           | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 2           | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 3           | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 4           | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 5           | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 6           | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 7           | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 8           | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 9           | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 10          | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 11          | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 12          | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 13          | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |

### Table 2. continued

| Primer pair | F15719 | R16114 | CUS | ASC |
|-------------|--------|--------|-----|-----|
| 1           | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 2           | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 3           | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 4           | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 5           | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 6           | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 7           | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 8           | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 9           | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 10          | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 11          | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 12          | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |
| 13          | ACTCATGTC | ATCTATTATA | CACTTAT | TGTCCCGCCA |

American domestic dog breeds and three wild canids. In mtDNA CR sequences, variable SNPs, and haplotypes of 125 presented by Gundry et al. (2007) which contains dog mtDNA.
specific for greyhounds and we discovered that one greyhound SNP is different. This part of bone sequence from position 16,461 to 16,490 is 3'-GAAAACCCCC TTACCCCCG TAAACTCATG-5'. At position 16,480 is nucleotide A which is specific for the greyhound. Nucleotide G at this position is specific for the Yorkshire Terrier or American Eskimo Dog and American Spitz, which are in the same group. All sequences established with primers set B, which determined the area 16,461 to 16,490 had nucleotide G at position 16,480. These findings seem to be a misleading result but we should note that our comparison was between different dogs from two different areas (Europe and North America). In the article was described only one greyhound sequence and this is probably the reason for the discrepancy (Zajc et al. 1997). Our recent greyhounds are demonstrably greyhounds and we found guanine at position 16,480. The same nucleotide was located at the same position in the sequence of the ancient bone found at the hillfort Chotěbuz-Podobora. After the previous comparison of ancient mtDNA with recent mtDNA from the N. Ireland greyhound we found that they are so similar that we are persuaded our ancient biological material comes from a greyhound and in view of the impressive history of this breed, there was a high probability that it was unthinkable to interbreed greyhounds with any other breed of dogs.

Furthermore, we compared the sequences of mtDNA from bone with sequences from recent greyhounds from the dog exhibition in Brno 2010, as can be seen in Tables 1 and 2. Compared to the ancient mtDNA deletions, insertions and substitutions were evident. The differences are shown in Tables 1 and 2. Some of them are marked by one, two or three stars because they appear interesting from an evolutionary point of view. We presume that for example a deletion of one or two A nucleotides (in the table it is shown as *) is the result of molecular evolution and more A nucleotides signify that the individual is evolutionarily older with respect to the mtDNA. Sample 8 is MAYSUN Bohemia Genao, whose grandfather was a Moroccan Sloughi, and is the only individual in which we observed the addition of a group of two T nucleotides (shown as **) in their mtDNA. The sloughi (sample 6) has a triplet of T substituted by a triplet of C. All changes found constitute the passage of time. The more changes compared to ancient DNA, the younger the specimen is. With respect to the foundations of paleogenetics, we can talk about a dog's "Eves", the rare changes signify different molecular "Eves", different places of development (Horai et al. 1995).

According to historical sources, greyhounds are a very ancient breed that has been prized by nobility for thousands of years. They can reach speeds of over 65 km/h being thus the fastest dog breed in the world. The true origin of the greyhound is unsure, but drawings of findings from the Çatal Hüyük site in Turkey (6000 BC), the finding of a greyhound-like dog in a funereal vase in the town of Fusa in Iran (4200 BC) or in rock art in Tassili (dated at 5000 – 2000 BC) indicate that the greyhound is indeed one of the oldest breeds of dog (Bazin 1959, Brentjes 1969, Branigan 2004). The greyhound-like dog came to the Chotěbuz-Podobora site almost certainly as an exclusive article from the territory of present-day Poland, probably as a gift to the local lord, or as loot. The presence of the greyhound-like dog coincides with other exceptional findings from Chotěbuz-Podobora which are dated from the second half of the 8th century and the first third of the 9th century. These indirectly confirm the assumption of inhabitants of privileged rank at the hillfort in the Great Moravian period (Kouřil 2007).

Conclusion

Genetic comparison of the dog bone from Chotěbuz-Podobora hillfort with genetic material from a modern greyhound has shown that the bone found most probably belonged to this breed.

Since ancient times the greyhound has been an exceptional animal which was only ever owned by nobles and rulers. The presence of a greyhound at the hillfort supports the theory that at the turn of the 8th and 9th centuries AD the fort was occupied by elite forces under the command of a local leader, who guarded the entrance to the Moravian Gate. It also implies that trade or political links existed with the region which is now Poland.

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