

Taxonomic identification of sample by DNA

Sample ID: CGG_6_000415_M

Samples collected: 10/04/2016

Locality: Fyn

Number of samples: 1

Type of material: saliva from roe deer meat

DNA investigation: DNA was successfully isolated from the sample using Blood and Tissue kit (Qiagen) following manufacturer's recommendations. Mitochondrial DNA was amplified amplified for two mitochondrial fragments with the primers H16498_R-L15995_F (370bp, universal primer) and WDLoopL-WDLoopH254 (250 bp, dog/wolf specific primer). The universal primer pair amplified the prey (roe deer). The WDLoop primer pair amplified DNA from a carnivore, *Canis lupus*. The consensus sequence yields a BLAST id of 100% for *Canis lupus*.

Consensus sequence for the primer pair WDLoopL-WDLoopH254 (CGG_6_000415_M):
 TTCCCTGACACCCCTACATTCAATATTGAATCACCCCTACTGTGCTATGTCAGTATC
 TCCAGGTAAACCCTTCTTCCCTCCCCTATGTACGTCGTGCATTAATGGTTGCCCAT
 GCATATAAGCATGTACATAATATTACATTCTACATAGGACATATTAACACTCAATCTC
 ATAATTCACTGATCTATCAAACAGTAATGCATATCACTTAGTCCAATAAGGGC
 TTAATCACCA

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Canis lupus isolate OW17 D-loop, partial sequence; mitochondrial
 Sequence ID: [KF723525.1](#) Length: 431 Number of Matches: 1

Range 1: 8 to 247 GenBank Graphics					▼ Next Match	▲ Previous Mat
Score	Expect	Identities	Gaps	Strand		
444 bits(240)	5e-121	240/240(100%)	0/240(0%)	Plus/Plus		
Query 1	TTCCCTGACACCCCTACATTCAATATTGAATCACCCCTACTGTGCTATGTCAGTATCTC			60		
Sbjct 8	TTCCCTGACACCCCTACATTCAATATTGAATCACCCCTACTGTGCTATGTCAGTATCTC			67		
Query 61	CAGGTAAACCCTTCTTCCCTCCCCTATGTACGTCGTGCATTAATGGTTGCCCATGCAT			120		
Sbjct 68	CAGGTAAACCCTTCTTCCCTCCCCTATGTACGTCGTGCATTAATGGTTGCCCATGCAT			127		
Query 121	ATAAGCATGTACATAATATTACATTCTACATAGGACATATTAACCTCAATCTCATAATT			180		
Sbjct 128	ATAAGCATGTACATAATATTACATTCTACATAGGACATATTAACCTCAATCTCATAATT			187		
Query 181	ACTGATCTATCAAACAGTAATCAAATGCATATCACTTAGTCCAATAAGGGCTAATCACCA			240		
Sbjct 188	ACTGATCTATCAAACAGTAATCAAATGCATATCACTTAGTCCAATAAGGGCTAATCACCA			247		

Figure 3. Screenshot for the BLAST search. Query refers to our sequences and Sbjct is the sequence in the database. The best match for our samples is the sequence ID: KF723525 belonging to a wolf.

Consensus sequence for the primer pair H16498_R- L15995_F (CGG_6_000415_M):
 GGGGGCAGATAACGATGTTGACAAGGACAGATTGACTTAATGCGCTATGTACGA
 TCAATAACATAATGTGCTATGTACTATTAACTTAATGGACTTGCTTATAAGCAT
 GGGGCATATAATATAATGTACTATTACATATTATGTCTTAATACATTAACCTTAAT
 GTACTATAAGCGCTTAAGTTATATCGTACATACGTTGCTACATGCTGTGAGTTA
 ATTATATGTAGCGTATATAATTGGGTATTGAGTACAAAAGTGAATTAAATTACTGAA
 GATTTAAAAAAATTAATACTGGTAAAGTTCTGGTTTTGGAACATATTAAATAAC
 GTATCAGGAAATAGTTAATTAGAACT

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Capreolus capreolus partial D-loop sequence
 Sequence ID: [AM279273.1](#) Length: 746 Number of Matches: 1

Range 1: 88 to 456 GenBank Graphics					▼ Next Match	▲ Previous
Score	Expect	Identities	Gaps	Strand		
671 bits(363)	0.0	367/369(99%)	0/369(0%)	Plus/Minus		
Query 1	GGGGGCAGATAACGATGTTGACAAGGACAGATTGACTTAATGCGCTATGTACGA				60	
Sbjct 456	GGGGCAGATAACGATGTTGACAAGGACAGATTGACTTAATGCGCTATGTACGA				397	
Query 61	TAACATAATGTGCTATGTACTATTAACTTAATGGACTTGCTTATAAGCATGGGCAT				120	
Sbjct 396	TAACATAATGTGCTATGTACTATTAACTTAATGGACTTGCTTATAAGCATGGGCAT				337	
Query 121	ATAATATAATGTACTATTACATATTATGTCTTAATACATTAACCTTAATGTACTATAAG				180	
Sbjct 336	ATAATATAATGTACTATTACATATTATGTCTTAATACATTAACCTTAATGTACTATAAG				277	
Query 181	CGCATTAAAGTTATATCGTACATACGTTGCTACATGCTTGTGAGTTAATTATATGTAGCG				240	
Sbjct 276	CGCATTAAAGTTATATCGTACATACGTTGCTACATGCTTGTGAGTTAATTATATGTAGCG				217	
Query 241	TATATAATTGGGTATTGAGTACAAAAGTGAATTAAATTACTGAAGATTTAAAAAAATTTA				300	
Sbjct 216	TATATAATTGGGTATTGAGTACAAAAGTGAATTAAATTACTGAAGATTTAAAAAAATTTA				157	
Query 301	ATACTGGTAAAGTTCTGGTTTTGGAACTATATTAAATAACGTATCAGGAATAGTTA				360	
Sbjct 156	ATACTGGTAAAGTTCTGGTTTTGGAACTATATTAAATAACGTATCAGGAATAGTTA				97	
Query 361	ATTAGAACT 369					
Sbjct 96	ATTAGAACT 88					

Figure 4. Screenshot for the BLAST search. Query refers to our sequences and Sbjct is the sequence in the database. The best match for our samples is the sequence ID: AM279273 belonging to a roe deer.

Taxonomic identification of sample by DNA

Sample ID: CGG_6_000415_S

Samples collected: 10/04/2016

Locality: Fyn

Number of samples: 9

Type of material: swabs

DNA investigation: DNA was successfully isolated from the sample using DNA Blood and Tissue kit (Qiagen) with some modifications from manufacturer's recommendations (protocol can be shared on request). Mitochondrial DNA was amplified for two mitochondrial fragments with the primers H16498_R-L15995_F (370bp, universal primer) and WDLoopL-WDLoopH254 (254 bp, dog/wolf specific primer). The universal primer pair amplified the prey (roe deer). The WDLoop primer pair amplified DNA from a carnivore, *Canis lupus familiaris*. The consensus sequence yields a BLAST id of 100% for *Canis lupus familiaris*. These results are consistent for the 9 analyzed swabs. We are able to identify two different haplotypes in our sequences. Sequences from swabs 1,4,6 and 8 present the same haplotype (haplotype A). While sequences from swabs 2,3,5,7 and 9 can be grouped together on a different haplotype (haplotype B).

Consensus sequence for the primer pair H16498_R- L15995_F (CGG_6_000415_S):

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CCAATTATACGCTACATATAATTAACTACACAAGCATGTAGCAACGTATGTACGA
TATAACTTAATGCCTTATAGTACATTAAGTTAATGTATTAGGACATAATATGTATA
ATAGTACATTATATTATGCCCATGCTTATAAGCAAGTCCATTAAGTCATTAAATA
GTACATAGCACATTATGTTATTGATCGTACATAGCGCATTAAAGTCAAATCTGTCCTT
GTCAACATGCGTATCCC GCCCCTAGATCACGAGCTTAATCACCATGCCCGTGAAA
CCAGCAACCCGCTTGGCAGGGATCCCTCTCGCTCCGGGCCATAAACATCGTGGGG
GTAGCTATTAATGAACCTTATCAGACA
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Capreolus capreolus haplotype Cc22 D-loop, complete sequence; mitochondrial
Sequence ID: KF724423.1 Length: 924 Number of Matches: 1

Range 1: 91 to 460					GenBank	Graphics	▼ Next Match	▲ Previous
Score	Expect	Identities	Gaps	Strand				
684 bits(370)	0.0	370/370(100%)	0/370(0%)	Plus/Plus				
Query 1	CCAATTATACGCTACATATAATTAACTACACAAGCATGTAGCAACGTATGTACGA			60				
Sbjct 91	CCAATTATACGCTACATATAATTAACTACACAAGCATGTAGCAACGTATGTACGA			150				
Query 61	AACTTAATGCCCTTATAGTACATTAAGTTAATGTATTAGGACATAATATGTATAATAGTA			120				
Sbjct 151	AACTTAATGCCCTTATAGTACATTAAGTTAATGTATTAGGACATAATATGTATAATAGTA			210				
Query 121	CATTATATTATATGCCCATGCTTATAAGCAAGTCCATTAAGTCATTAATAGTACATAGC			180				
Sbjct 211	CATTATATTATATGCCCATGCTTATAAGCAAGTCCATTAAGTCATTAATAGTACATAGC			270				
Query 181	ACATATGTTATTGATCGTACATAGCGATTAAGTCAAATCTGTCCTTGTCAACATGCGT			240				
Sbjct 271	ACATATGTTATTGATCGTACATAGCGATTAAGTCAAATCTGTCCTTGTCAACATGCGT			330				
Query 241	ATCCCCCCCCCTAGATCAGCAGCTTAATCACCATGCCGGTGAACCCAGCAACCCGGTTG			300				
Sbjct 331	ATCCCCCCCCCTAGATCAGCAGCTTAATCACCATGCCGGTGAACCCAGCAACCCGGTTG			390				
Query 301	GCAGGGATCCCTCTTCCTCGTCCGGGCCATAAATCGGGGGTAGCTATTAAATGAACCT			360				
Sbjct 391	GCAGGGATCCCTCTTCCTCGTCCGGGCCATAAATCGGGGGTAGCTATTAAATGAACCT			450				
Query 361	TTATCAGACA 370							
Sbjct 451	TTATCAGACA 460							

Figure 5. Screenshot for the BLAST search. Query refers to our sequences and Sbjct is the sequence in the database. The best match for our samples is the sequence ID: KF724423 belonging to a roe deer.

Consensus sequence for haplotype A for the primer pair WDLoopL-WDLoopH254

(CGG_6_000415_S):

ATTCCTGACACCCCTACATTCATATATTGAATCACCCCTACTGTGCTATGTCAGTAT
CTCCAGGTAAACCCTCTCCCCTCCCTATGTACGTCGTGCATTAATGGTTGCCCA
TGCATATAAGCATGTACATAATATTATACCTTACATAGGACATATTAACACTCAATCT
CATAATTCACTGATCTATCAACAGTAATCGAATGCATATCACTTAGTCCAATAAGGG
CTTAATCACCATGCCTCGAGAAC

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Canis lupus familiaris control region, partial sequence; mitochondrial
Sequence ID: [KT946984.1](#) Length: 404 Number of Matches: 1

Range 1: 51 to 304 GenBank Graphics

Score	Expect	Identities	Gaps	Strand
470 bits(254)	9e-129	254/254(100%)	0/254(0%)	Plus/Minus

▼ Next Match ▲ Previous M

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Query 1   ATTCCTGACACCCCTACATTCATATATTGAATCACCCCTACTGTGCTATGTCAGTATCT  60
Sbjct 304  ATTCCTGACACCCCTACATTCATATATTGAATCACCCCTACTGTGCTATGTCAGTATCT  245
Query 61   CCAGGTAAACCCTCTCCCCTCCCTATGTACGTCGTGCATTAATGGTTGCCCAATGCA  120
Sbjct 244   CCAGGTAAACCCTCTCCCCTCCCTATGTACGTCGTGCATTAATGGTTGCCCAATGCA  185
Query 121  TATAAGCATGTACATAATATTATACCTTACATAGGACATATTAACCTCAATCTCATATT  180
Sbjct 184  TATAAGCATGTACATAATATTATACCTTACATAGGACATATTAACCTCAATCTCATATT  125
Query 181  CACTGATCTATCAACAGTAATCGAATGCATATCACTTAGTCCAATAAGGGCTTAATCACC  240
Sbjct 124  CACTGATCTATCAACAGTAATCGAATGCATATCACTTAGTCCAATAAGGGCTTAATCACC  65
Query 241  ATGCCCTCGAGAAC  254
Sbjct 64   ATGCCCTCGAGAAC  51

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Figure 6. Screenshot for the BLAST search. Query refers to our sequences and Sbjct is the sequence in the database. The best match for our samples is the sequence ID: KT946984 belonging to a dog.

Consensus sequence for haplotype B (CGG_6_000415_S):

TTCCCTGACACCCCTACATTCATATATTGAATTACCCCTACTGTGCTATGTCAGTATC
TCCAGGTAAACCCCTCTCCCCTCCCCTATGTACGTCGTGCATTAATGGTTGCCCAT
GCATATAAGCATGTACATAATATTATATCCTACATAGGACATATTAACACTCAATCTC
ATAGTTCACTGATCTATCACAGTAATCGAATGCATATCACTTAGTCCAATAAGGGC
TTAACACCATGCC

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Canis lupus familiaris isolate Cf_ws9 mitochondrion, complete genome
Sequence ID: [KM061594.1](#) Length: 16730 Number of Matches: 1

Range 1: 15454 to 15697					GenBank	Graphics	▼ Next Match	▲ Previous Match
Score	Expect	Identities	Gaps	Strand				
451 bits(244)	3e-123	244/244(100%)	0/244(0%)	Plus/Plus				
Query 1	TTCCCTGACACCCCTACATTCATATATTGAATTACCCCTACTGTGCTATGTCAGTATCTC		60					
Sbjct 15454	TTCCCTGACACCCCTACATTCATATATTGAATTACCCCTACTGTGCTATGTCAGTATCTC		15513					
Query 61	CAGGTAAACCCCTCTCCCCTCCCCTATGTACGTCGTGCATTAATGGTTGCCCATGCAT		120					
Sbjct 15514	CAGGTAAACCCCTCTCCCCTCCCCTATGTACGTCGTGCATTAATGGTTGCCCATGCAT		15573					
Query 121	ATAAGCATGTACATAATATTATCCTACATAGGACATATTAACACTCAATCTCATAGTTC		180					
Sbjct 15574	ATAAGCATGTACATAATATTATCCTACATAGGACATATTAACACTCAATCTCATAGTTC		15633					
Query 181	ACTGATCTATCAACAGTAATCGAATGCATATCACTTAGTCCAATAAGGGCTTAATCACCA		240					
Sbjct 15634	ACTGATCTATCAACAGTAATCGAATGCATATCACTTAGTCCAATAAGGGCTTAATCACCA		15693					
Query 241	TGCC 244							
Sbjct 15694	TGCC 15697							

Figure 7. Screenshot for the BLAST search. Query refers to our sequences and Sbjct is the sequence in the database. The best match for our samples is the sequence ID: KM061594 belonging to a dog.