Sequence analysis of the cytochrome c oxidase subunit 1 gene of Sarcoptes scabiei isolated from goats and rabbits in East Java, Indonesia

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Abstract

Aim: This study aimed to sequence the Cytochrome c oxidase (COX-1) gene sequence from mitochondrial DNA of Sarcoptes scabiei isolated from Lamongan goats and Mojokerto rabbits, align it with DNA isolated from Zi’gong rabbit (GenBank accession No. EU256389.1), and produce a phylogenetic analysis of S. scabiei COX-1 gene.

Materials and Methods: S. scabiei mites were obtained from goats and rabbits, and DNA was extracted using QIAamp DNA Mini Kit. The forward and reverse primer sequences were designed based on the DNA sequence of an S. scabiei COX-1 gene isolated from the Zi’gong rabbit (5’-TCTTAGGGGCTGATTAGTAG-3’ and 5’-AGTTCCCTCTACAGTTCCAC-3’, respectively). To confirm sequencing output, the sequence resulting from the reverse primer was inverted and aligned to the sequence from the forward primer using Clone Manager Professional Version 9 for Windows (Scientific & Educational Software; http://www.scied.com). This alignment was subsequently used to build a phylogenetic tree, using the Neighbor-Joining method, in the MEGA6 program (https://www.megasoftware.net/).

Results: Polymerase chain reaction (PCR) products from S. scabiei isolates from Lamongan goats and Mojokerto rabbits produced bands of around 290 bp with 2% agarose gel electrophoresis. Comparing the DNA sequences of the S. scabiei COX-1 gene with those isolated from various goats and Lamongan rabbits showed 99% homology.

Conclusion: PCR products of the S. scabiei COX-1 gene isolated from Lamongan goats and Mojokerto rabbits were around 290 bp long. The sequences had more than 99% homology. The sequences of the COX-1 gene of S. scabiei from Lamongan goats and Mojokerto rabbits were relatively close to the sequence of the gene in S. scabiei obtained from various hosts according to National Center for Biotechnology Information data.

Keywords: cytochrome c oxidase-1, East Java, goat, Indonesia, rabbit, Sarcoptes scabiei.

Introduction

Scabies an infectious skin disease produced by the Sarcoptes scabiei mite affects both humans and animals. Outbreaks in domestic animals and wild mammals cause morbidity, mortality, and huge economic losses [1,2]. Scabies is very contagious and produces pruritic dermatitis, alopecia, hyperkeratosis, and crust formation [3,4]. Histopathological changes, manifest as lesions, parakeratosis, acanthosis, congestion, inflammation, and cell degeneration [5]. Scabies causes global health problems as an infectious disease which appears and re-appears [6]. Scabies outbreaks have been reported in industrialized countries, and problems caused by the disease in developing countries are increasing [7]. There are problems due to drug resistance, drug residue, and toxicity from extensive use of acaricides, especially in developing countries [8]. The S. scabiei mite has a range of hosts. Although there are no differences in the S. scabiei mite’s morphology found on different animal species, they generally show low cross-infectivity [9]. Experiments show no transmission of scabies from dog to rat, marmot, pig, cow, cat, sheep, or goat [10,11]. Research was also conducted by Harumal et al. [12] using the S. scabiei antigen 1 from cloning S. scabiei var. hominis in rabbits. The result was less protective after challenge, but the rabbits did not develop crusting.

There are ongoing genetic studies of S. scabiei. Analysis of a 450bp nucleotide ribosomal marker on the second internal transcribed spacer (ITS2) of S. scabiei such as dog, cow, fox, wombat, and dromedary found no differences [13]. There is still limited, research on developing molecular diagnostic tools for scabies because it is difficult to isolate the Sarcoptes
mite. Microscopic examination of skin scrapings shows a positive result in only about 30-50% of scabies cases with crusta papula [8]. There is no immunodiagnostic test or commercial subunit vaccine available for scabies. Some research has focused on producing a recombinant protein from S. scabiei, aimed investigating the host’s immune response, to develop a subunit vaccine [1,14]. Based on field observations, the number of scabies cases in goats and rabbits in Indonesia increasing; however, there have been no case reports because the condition has been treated with ivermectin and acaricide.

The Cytochrome c oxidase (COX-1) gene is often used by researchers for genetic characterization of S. scabiei from animals and humans. The COX-1 fragments are often used for DNA barcoding to differentiate between species. These fragments rarely experience amino acid substitutions, but silent mutations often occur. The COX-1 fragment is useful for reconstructing phylogenetic diversity in evolutionary branches below the species level. Mitochondria undergo rapid evolution, but there are parts of it, such as the COX-1 fragment that experience low evolution, rates and can, thus, be used as genetic characters [9,15,16].

This study aimed to sequence the COX-1 gene sequence from mitochondrial DNA of S. scabiei isolated from Lamongan goats and Mojokerto rabbits, align it with DNA isolated from Zi’gong rabbit (GenBank accession No. EU256389.1), and produce a phylogenetic analysis of S. scabiei COX-1 gene.

Materials and Methods
Ethical approval
The research complied with the ethics guidelines for using experimental animals and was approved by the Ethics Commission of the Faculty of Veterinary Medicine, Universitas Airlangga, Indonesia, No: 630-KE.

Isolation of S. scabiei mites
S. scabiei mite samples were collected from ten rabbits and six goats infected with scabies with severe clinical symptoms, including skin thickening, crust formation, and hair loss around their eyes, ears, mouths, and legs. The goats were from husbandries at Lamongan and the rabbits from Mojokerto, East Java, Indonesia. About 250 mg of crusted skin was scraped with a sterile scalpel from the goats’ and rabbits’ eyes, washed in Petri dishes with phosphate-buffered saline (PBS), and filtered to remove skin debris. The washing process was repeated 3 times. As a final washing step, the crusted skin was transferred to a 1.5 ml microcentrifuge tube, washed with 1 ml PBS by vortexing, and centrifuged at 3,000 revolutions/min (rpm) for 10 min. The filtrate was discarded, and the clean, crusted skin was stored at −20°C until used [2,4].

Extraction of mitochondrial DNA
DNA extraction was performed using QIAamp DNA Mini Kits (Qiagen, Hilden, Germany), following the manufacturer’s protocol as follows: 20 µl of Qiagen protease (20 mg/ml) was added to a microcentrifuge tube with a 200 µl suspension of S. scabiei; and 180 µl of buffer tissue lysis (ATL buffer) was then added and the tube was centrifuged at 8,000 G for 3 min, vortexed for 15 s, and incubated at 56°C for 24 h. Then, 200 µl of buffer lysis (AL buffer) was added to the tube and vortexed for 15 s; 200 µl of 96% ethanol was added to the tube and vortexed for 15 s. The materials were removed, placed into a column tube, and centrifuged at 8,000 G for 1 min to pellet the DNA. Next, 500 µl of washing buffer 1 (AW1 buffer) was added to the pellet and centrifuged at 8,000 G for 1 min; 50 µl of AW2 buffer was then added to the pellet, and the tube was centrifuged at 13,000 G for 3 min. The tube was removed and changed; then 50 µl of elution buffer (AE buffer) was added to the final DNA pellet. After incubation at 15-25°C for 1 min, the contents were centrifuged at 8,000 G for 1 min to collect the DNA [15,17].

DNA amplification and sequencing
A 290-bp DNA fragment of S. scabiei COX-1 was amplified by polymerase chain reaction (PCR) using the isolated genomic DNA as a template. The sequences of the forward and reverse primers were designed using the program Primer3Plus[18] based on the DNA sequence of an S. scabiei COX-1 gene isolated from the Zi’gong rabbit (GenBank accession No. EU256389.1). The forward primer was 5’-TCTTAGGGGCTGGATTTAGTATG-3' and the reverse primer was 5’-AGTTCTCTACCAGTTCCAC-3’. PCR volumes were 20 µl, consisting of 10 µl Master Mix (Thermo Scientific), 1 µl each of forward and reverse primers, 6 µl distilled water, and 2 µl DNA template. PCR amplification was performed in an iCycles IQ (Biorad) with an initial denaturation at 95°C for 5 min, followed by 35 temperature cycles, each consisting of a 30 s denaturation at 95°C, a 60 s annealing at 50°C, and a 60 s extension at 72°C. After 35 cycles, there was a final extension for 5 min at 72°C. The amplified PCR product was electrophoresed on a 2% agarose gel in 1× Tris-borate-EDTA buffer with pH 8.3 and 0.03 µl ethidium bromide. The gel was visualized under ultraviolet light [15,19].

The PCR product was then purified according to the protocol of the BigDye XTerminator Purification Kit (Thermo Scientific, USA) and double-sequenced with the PCR forward and reverse primers using an ABI PRISM 310 Genetic Analyzer (Applied Biosystems).

Bioinformatics analysis
To confirm the sequencing output, the sequence resulting from the reverse primer was inverted and aligned to that from the forward primer using the program Clone Manager Professional 9 Version 9 for Windows (Scientific & Educational Software; http://www.scied.com) using the DNA sequence with
GenBank accession No. EU256389.1 mentioned above as a reference. The confirmed COX-1 sequences of the Mojokerto rabbits and Lamongan goats were deposited in GenBank (https://www.ncbi.nlm.nih.gov/genbank/) under access codes MH077557 (Lamongan goat) and MH077558 Mojokerto rabbit.

For comparison with other related DNA sequences, the COX-1 DNA fragment sequence from the Mojokerto rabbit (GenBank accession: MH077558) was input as a query search with the program nucleotideBLAST. All hits with a nucleotide identity of 80% or higher, along with all sequences resulting from this investigation, were aligned using the ClustalW2 program [20]. This alignment was subsequently used to build a phylogenetic tree with the Neighbor-Joining method, in the MEGA6 program (https://www.megasoftware.net/) [21,22], with a COX-1 sequence from Megaselia spp. (GenBank accession No. KT103510.1) as an outer group.

Results

S. scabiei mites were isolated from Lamongan goats and Mojokerto rabbits, with clinical symptoms: Thickening of the skin, crust formation, and hair loss on the areas around the eyes, ears, mouths, and legs. The results of 2% agarose gel electrophoresis of COX-1 DNA from S. scabiei, amplified with the primer pairs described above, show the PCR products were around 290bp long, indicating bands at positions between 200 bp and 300 bp markers (Figure-1).

Sequence analyses confirmed that the PCR products were a partial CoDing Sequence (CDS) of the COX-1 gene (Figure-2). Sequence comparison showed that the partial CDS of the S. scabiei COX-1 gene isolated from Lamongan goats is homologous to that from Mojokerto rabbits, with an identity of 99%. Out of 290 nucleotides aligned, only one nucleotide was different. An adenine was found in the CDS of the S. scabiei COX-1 gene isolated from Lamongan goats, in place of guanine in that position in the sequence from Mojokerto rabbits. The guanine residue is conserved in other S. scabiei COX-1 gene. A BLASTN analysis against the National Center for Biotechnology Information (NCBI) nucleotide collection (nr) database reveals about 19 S. scabiei isolates from animal species from various countries, which were similar identical at nucleotide lengths of 290 bp.

The phylogenetic tree of the COX-1 gene of S. scabiei mites from Lamongan goats and Mojokerto rabbits was constructed using the MEGA6 program’s Neighbor-Joining method. This tree shows that the gene is relatively close to 19 S. scabiei isolates obtained from the NCBI nucleotide database with their accession numbers [20] (Figure-3). Phylogenetic analysis of the COX-1 nucleotide sequences revealed three clades of S. scabiei: (1) A large, unresolved clade, including sequences from Lamongan goats and Mojokerto rabbits, and sequences obtained from the GenBank of S. scabiei from various hosts and regions; (2) a clade of S. scabiei from Australian wombats and S. scabiei isolate B1; and (3) a clade of S. scabiei type hominis from Australia and Megaselia spp. from Canadian insects, used as an outer group in this analysis (Figure-3).

Discussion

The sequence alignment has shown that the partial CDS of S. scabiei COX-1 gene isolated from Lamongan goats is almost identical to that from Mojokerto rabbits. Out of 290 nucleotides aligned, only one nucleotide was different, an adenine was found in the CDS of the S. scabiei COX-1 gene isolated from Lamongan goats in place of guanine in that sequence from Mojokerto rabbits. The guanine residue is conserved in other S. scabiei COX-1 genes. The COX-1 gene has conserved region and variable region, and differences are possible in the variable region. However, this mutation is most likely silent [9,16].

Based on the alignment results, the COX-1 mitochondrial S. scabiei gene has a high level of identity among isolates. Possibly, the S. scabiei infecting Lamongan goats was the same as the one infecting Mojokerto rabbits. This observation indicates that S. scabiei isolates from Lamongan goats and Mojokerto rabbits are the same species, and the COX-1 mitochondrial S. scabiei did not evolve. Makouloutou et al. [9] have stated that the S. scabiei causes scabies in most Japanese wild animals, similar to that in wild animals on other continents.

Research into S. scabiei mites’ genetics is still underway. Some research based on studies using the second internal transcribed spacer (ITS2) of the ribosomal RNA gene. The ITS2 gene has a conserved region and a variable region, at the genus level, it has the same conserved region, but the variable region will be different depending on the species. Taxonomically, S. scabiei has different varieties based on the host’s origin and shows cross-infectivity to a low degree between hosts, because each mammal has a

Figure-1: Polymerase chain reaction products of Sarcoptes scabiei from goat-Lamongan (S1) and rabbit-Mojokerto (S2), negative control (K), marker (M).
Figure-2: Sequence alignment of the polymerase chain reaction products of the cytochrome c oxidase-1 gene of *Sarcoptes scabiei* from Lamongan goats and Mojokerto rabbits with data from GenBank.

Figure-3: Phylogenetic analysis of partial CDS of the cytochrome c oxidase subunit 1 gene of *Sarcoptes scabiei*, isolated from several different species. The genes from Lamongan goats and Mojokerto rabbits are introduced in this article, while the other sequences were obtained from the National Center for Biotechnology Information (NCBI) nucleotide database, their accession numbers are shown. All sequences were aligned using ClustalW2, and the cladogram was built using the Neighbor-Joining method [20]. Bootstrap test values, which were resulted from 500 replications are indicated for each branch. The sequences with the NCBI accession number KT103510.1 are from *Megaselia* spp. BOLD-2016 voucher BIOUG06146-B08 and were used as an outlier in this analysis.
specific receptor. S. scabiei morphology from various mammalian hosts and various regions indicates that it is a single species, but with heterogeneous variable region [4,13,23,24]. Erster et al. [25] reported that an analysis of mitochondrial COX-1 did not show a correlation between host preference and genetic identity in S. scabiei from pet, farm, and wild hosts in Israel.

The phylogenetic analysis (Figure-3) shows relatively low bootstrap values, indicating the COX-1 fragment sequences analyzed are rather difficult to resolve. Indeed, the sequences are highly similar. The sequences’ 290bp length may also contribute to these low bootstrap values. However, from the phylogenetic analysis, it is clear that S. scabiei from Lamongan goats and Mojokerto rabbits is relatively close to S. scabiei from various hosts such as swine, dog, rabbit, and Australian and Chinese marsupials. However, Megalacia spp. from Canadian insects is an outer group. Scabies is highly infectious and transmitted through direct contact, so it is possible to transmit it between goats kept in one cage, as well as between rabbits that are kept in one cage with poor sanitation. According to Arlian et al. [10], S. scabiei cross-infestation between different hosts does not occur in wild animals. In our study, S. scabiei mites were isolated from goat farms in the Lamongan area, where each farmer had around five to six animals in one cage, and they showed clinical symptoms of scabies. Similarly, S. scabiei isolated from rabbits showing clinical symptoms of scabies came from rabbit farms in Mojokerto, which 6-10 rabbits are kept in one cage with poor sanitation, so transmission between rabbits is easy. Transmission within a single species is possible because individuals have the same receptor. S. scabiei mites have adapted well to their hosts. Adaptation to the host and geographical segregation will reduce transmission between S. scabiei species. Understanding the transmission of S. scabiei has important implications in epidemiological studies for developing scabies control strategies in animals and humans [24,26,27].

Conclusion

The PCR products of S. scabiei COX-1 gene isolated from Lamongan goats and Mojokerto rabbits were around 290 bp in length. Sequences showed a similarity of more than 99%. The sequences of the COX-1 gen S. scabiei from Lamongan goats and Mojokerto rabbits were relatively close to those of the COX-1 gene of S. scabiei obtained from the NCBI databases from various hosts.

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Competing Interests

The authors declare that they have no competing interests.

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