Identification and Characterization of \textit{Staphylococcus aureus} 16S rRNA gene isolated from different Food Specimens from South Indian Region

Moneddu Kiran Kumar$^1$, Charu Tyagi$^2$, Arjun Sahu$^3$, Nalini Desai$^2$, Jayanand Manjhi$^1$, Kakarla Chandra Mohan$^5$, Yugandhar P. Reddy$^3$, Santosh Kumar Tiwari$^4$, Lomas Kumar Tomar$^{1,4*}$, Varun Kumar Sharma$^{1,4*}$

$^1$Department of Biotechnology & Microbiology, School of Sciences, Noida International University-NIU, Gautam Budh Nagar-201308, Uttar Pradesh, India

$^2$Department of Biotechnology, VSPG (PG) College, CCS University, Meerut-250004, Uttar Pradesh, India

$^3$Department of Zoology, TheAdoni Arts and Science College, Adoni, Kurnool, Andhra Pradesh, India

$^4$Centre for Liver Research and Diagnostics, Deccan College of Medical Sciences, Kanchanbagh, Hyderabad 500 058, Telangana, India

$^5$Department of Zoology, KVR Government Degree College for Women, Kurnool, Andhra Pradesh, India

\textbf{ABSTRACT}

\textit{Staphylococcus aureus} \((S.\ aureus)\) associated food-borne diseases have global impact on human health. Genome wide analyses have shown that \textit{S. aureus} contains specific endotoxin expressing gene and produce toxic proteins which is responsible for food contamination. Appropriate detection of pathogens is one of the major tools to avoid infection rate and reduce the health and socio-economic burden to human being. In addition, inappropriate handling the specimens, misdiagnosis and limited standard medical support could directly influence the infection rate.

The objective of this study was to identify \textit{S. aureus} from different food specimens from Hyderabad, India. A total of 70 random bacterial nutrient agar medium pure plates were made based on different morphological appearance of bacterial colonies. Preliminary identification of \textit{S. aureus} based on standardized morphological method showed specific golden yellow colonies. Biochemical assay also verified bacterial specimens. Furthermore, molecular characterization was performed on the basis of polymerase chain reaction (PCR) and sequencing of 16S rRNA gene of \textit{S. aureus}. Newly sequenced 16S rRNA gene sequences showed 100% homology to \textit{S. aureus}, analyzed using NCBI-BLAST tool.

The phylogenetic analysis and nucleotide base composition studies performed using 39 sequences of 16S rRNA gene from different isolates of \textit{Staphylococcus}, including \textit{Staphylococcus aureus}. For the purpose, 16S rRNA gene sequences were retrieved from the NCBI in FASTA format. The phylogenetic analysis was performed using Maximum Likelihood method and revealed the relationships and percent similarity of \textit{Staphylococcus aureus} 16S rRNA gene.

**Keywords:** Food-borne diseases; 16S rRNA gene; Maximum Likelihood; Phylogenetic analysis; \textit{Staphylococcus aureus}.

\textbf{INTRODUCTION}

Food-borne associated disease is one of the major health problem around the world, including food-borne intoxications and food-borne infections\(^1,2\). As per reference to Foodborne Disease Burden Epidemiology Reference Group (FERG) Report published in 2010, it has been estimated 3-5 billion people affected to food-borne associated disease and nearly 1.5 million deaths annually assumed worldwide\(^3\). Food-borne associated diseases mainly caused by a wide variety of pathogen, including bacterial species such as \textit{Salmonella} spp., \textit{Vibrio} spp., \textit{Clostridium} spp., \textit{Campylobacter} spp., \textit{Salmonella} spp., \textit{Campylobacter} spp., and \textit{Staphylococcus} spp\(^4\).

\textit{Staphylococcus aureus} is one of the major causative agent of food-borne associated disease and most prompt sporadic food-borne diseases are linked to \textit{Staphylococcus aureus}, in the different region of the world\(^4,5\). \textit{Staphylococcus aureus} mainly found in food stuff which grows and produce endotoxins result contaminating the food material\(^6,7\). The ubiquitous \textit{Staphylococcus} species are observed to found a wide variety of food materials, including vegetarian and non-vegetarian items, range from raw, cooked and ready-to-eat...
foods stuff, which increase the risk for all consumers and can affect financial burden to the society. The higher survival probability range of Staphylococcus aureus have been observed that it can survive in low to high temperature range and can grow in different pH range from acidic to alkaline medium. A limited laboratory practices are available to evaluate and the confirmation of Staphylococcus aureus for the commercial diagnostic purpose. Additionally, misinterpretation, lack of standard medical attention and mishandling the specimens are also increase the risk of misinterpretation Staphylococcus aureus prevalence.

The genome of Staphylococcus aureus is approximately 2.3 Million base pair (bp) which mainly consists housekeeping genes, set of virulence genes and other genes required for growth and survival. Staphylococcus aureus has been isolated about seven decade ago from human specimen, and from the first isolation to till date, it is a one of the most extensively studied bacterial strain by various research groups from different countries around the world. It has been observed that pathogenesis of Staphylococcus aureus control mainly by group of thirty genes, including basic survival gene and virulent genes.

The bacterial genome wide sequencing analysis approaches have been developed tool that effectively efficiently help to recognize the specific bacterial strains, including 16S rRNA gene sequencing. Genome sequence analysis also provides a platform to compare genome diversity as well as help to understand the evolutionary association in intra- and inter-species populations. Interestingly, for the molecular detection of Staphylococcus aureus can be easily performed by amplifying the 16S rRNA and other endotoxin virulent gene sequence. In the current study, primarily, we focused to screen and identify Staphylococcus aureus strain from different food materials, including processed/canned food, home sterilized food and unsterilized/raw food using microbial and biochemical assay. Further, we aimed to validate and characterize 16S rRNA gene amplification and DNA sequencing method. We have also performed the multiple sequence alignment and phylogenetic tree to identify the genetic variation and evolutionary relationship between different isolate of Staphylococcus aureus.

### MATERIALS AND METHODS

#### Collection food specimen

The study was carried out over a 6-months period from the region of Hyderabad, Telangana State of India. Food material samples were collected from the local food stores, road-side hawkers and home using sterile sample collection method, from the selected regions. Based on the availability food materials, specimens were classified in three independent categories, such as: (1) Processed and canned food (2) Home sterilized food and (3) Raw and unsterilized food

#### Preparation of the bacterial culture

Collected specimens were subjected for microbial isolation using basic plating technique on nutrient agar medium. The samples were separately macerated/mashed and added to 10ml of sterile distilled water and homogenously mixed. 0.1ml of this sample solution was used for spreading onto the nutrient agar plates under laminar airflow conditions. The plates were incubated in an inverted position for overnight at 37°C.

#### Screening and selection of bacterial specimen

From the 18 master plates obtained a total of 70 random bacterial nutrient agar medium pure plates were made based on different morphological appearance of their colonies. The names of collected food samples and the number of pure plates obtained from them are shown in the table 1.

### Table 1: Detail of collected Food samples and quantity utilized for the current study.

| S. No. | Food material          | (n) | Food material          | (n) | Food material          | (n) |
|-------|------------------------|-----|------------------------|-----|------------------------|-----|
| 1     | Milk                    | 4   | Idli                   | 4   | Dosa                   | 4   |
| 2     | Canned sweet            | 4   | Steamed Rice           | 5   | Sweet                  | 6   |
| 3     | Instant curry (Veg)     | 5   | Homemade Curry (Veg)   | 3   | Noodles                | 6   |
| 4     | Instant curry (Non-Veg) | 6   | Homemade Curry (Non-Veg)| 3 | Bhaji                  | 4   |
| 5     | Canned juice            | 5   | Coconut Paste (Chutney)| 5   | Pani Poori             | 6   |
| 6     | Instant Idly batter     | 5   | Fresh juice            | 2   | Road-side fruit juice  | 4   |

n = number of samples

#### Preliminary identification of Staphylococcus species.

All bacterial specimen culture were primarily subjected for gram staining along with the morphological characterization of bacterial populations, mainly Staphylococcus species. We have adopted standardized morphological method to identify the Staphylococcus colonies with golden yellow color and watery consistency. All identified colonies were further subjected for depth morphological observation using grams staining method.

#### Biochemical identification method.

On the basis of preliminary morphological observation, selected colonies of Staphylococcus species were subjected for biochemical assay. We have performed basic biochemical assay such Catalase, Coagulase, Methyl Red Voges Proskauer, Mannitol and DNase Tests.
Molecular characterization of *Staphylococcus* species

Morphological and Biochemical assay based positive samples of *Staphylococcus* species were further processed for molecular based method using PCR based assay. Details are given below:

Genomic DNA Extraction from the selected bacterial cultures

DNA extraction from bacterial colonies were performed using Sodium dodecyl sulfate (SDS)-based method\(^5\), \(^6\). The bacterial cell suspension was treated with a lysis buffer containing SDS, Trisaminomethane Hydrochloric acid (Tris HCl) and Ethylene-diaminetetra-acetic acid (EDTA). The cell debris and other impurities were removed in several steps sequentially with simultaneous centrifugation. The Genomic DNA were precipitated using chilled ethyl alcohol. The precipitated DNA was collected as a pellet by centrifugation. The pellet was dissolved in TE buffer and stored at 4°C until next use.

After extraction of the genomic DNA from bacterial colonies, we aimed to verify the *Staphylococcus aureus* by 16S rRNA sequence using PCR method, which is highly sensitive, compared to the most common microbial culture and staining techniques. For the purpose, we aimed to design a specific set of primers to amplify the 16S rRNA sequence.

*Staphylococcus aureus* 16S rRNA sequence retrieval and primer designing

The *Staphylococcus aureus* strain ATCC 12600 16S ribosomal RNA, complete sequence (NR_118997.2) was retrieved from the National Center for Biotechnology Information (NCBI; http://www.ncbi.nlm.nih.gov) databases in FASTA format. Primer designing marker was designed using Primer 3 (V.0.4.0) online tool (http://bioinfo.ut.ee/ primer3-0.4.0/primer3/).

Amplification of 16S rRNA gene using PCR

For the PCR reaction, total reaction volume was 50 μl containing 5 μl of DNA template, 10 pmol of each primer (forward primer and reverse primer, purchased from Sigma-Aldrich, Hyderabad) 200 μmol of each deoxyribonucleoside triphosphate per liter, 1.5 mmol of MgCl\(_2\) per liter, 1.5 mmol of Tris-HCl (pH 8.8) per liter, 50 mmol of KCl per liter, and 0.1% Triton X-100.

Sequence analyzing of 16S rRNA gene

The PCR product was purified the PCR product using QIAquick gel extraction kit (Qiagen, Germany). 15μl of the purified DNA was sequenced by use of the ABI Prism DNA sequencing kit, Big Dye Terminator Cycle Sequencing (version 3.0), and ABI Prism 310 genetic analyzer (Applied Biosystems, USA). Comparison of the sequence in a reference database was performed by using an identification program based on selection of the longest recursive matches for optimal alignment of the compared sequences. The reference database sequences retrieved and combined from NCBI GenBank. The final sequence comparisons to the best matches were done manually.

Local sequence alignment

Basic Local Alignment Search Tool (BLAST) was performed for the different isolates of *Staphylococcus aureus* 16S ribosomal RNA gene sequence retrieved from NCBI to identify the homology or similarity its relatives in different isolated of *Staphylococcus aureus* using the online NCBI-BLAST (https://blast.ncbi.nlm.nih.gov/Blast.cgi). This software takes the data in FASTA format and produces the BLAST table.

Phylogenetic analysis

Phylogenetic analysis of *Staphylococcus aureus* 16S ribosomal RNA gene sequence retrieved from NCBI through Maximum likelihood methods and Phylogenetic trees were constructed by the software showing the ancestral relationship among the sequences. The Maximum Likelihood phylogenetic tree give different clusters showing their evolution relationship with each other and tree reveals different clade showing their evolutionary relationship within different isolates of *Staphylococcus aureus*. The sequences which lie in the same cluster are closely related.

RESULTS AND DISCUSSION

**Sequence and Discussion**

*Staphylococcus aureus* strain ATCC 12600 16S ribosomal RNA, complete sequence was retrieved from the NCBI in FASTA format. The sequence of the gene (NR_118997.2) is as the following:

>NR_118997.2 Staphylococcus aureus strain ATCC 12600 16S ribosomal RNA, complete sequence.

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**Title:** Molecular characterization of *Staphylococcus* species

**Abstract:**

This study aimed to characterize the molecular identity of *Staphylococcus* species isolated from clinical samples. Genomic DNA was extracted from bacterial colonies using a SDS-based method. The extracted DNA was used for amplification of the 16S rRNA gene using specific primers designed using Primer 3 software. The amplified DNA was sequenced using the ABI Prism 310 genetic analyzer. The sequence data was aligned using Basic Local Alignment Search Tool (BLAST) and phylogenetic analysis was performed using Maximum likelihood methods. The results showed the identity of the isolated *Staphylococcus* species and their relationship with other known species. This study provides a molecular basis for the identification of *Staphylococcus* species, which can be useful for diagnostic and research purposes.

**Keywords:** *Staphylococcus*, 16S rRNA gene, BLAST, Phylogenetic analysis.
Specific set of primers of *Staphylococcus aureus* 16S rRNA DNA sequence

Specific primer of *Staphylococcus aureus* 16S rRNA sequence was designed using Primer 3 (V.0.4.0) online tool, as depicted in Figure 1. We obtained a set of primer for 16S rRNA sequence of covering 570 nucleotide sequence length. The length of forward primer/sense primer is 127 nucleotide (highlighted with red color) and reverse primer/antisense primer length is 705 nucleotide (highlighted with blue color). We observed primers are specific by using NCBI, primer blast and no primer-primer dimer. Primers were observed with no helping structure. We also check self-complementary alignment of a specific primer using Oligo Calc (Oligonucleotide properties calculator). Details of the primers sequences as well as the PCR programs and protocol are provided in Table 1.

![Figure 1: Staphylococcus aureus 16S rRNA gene Primer3 Output](image)

**Table 1**: Detail of primers sequences used to amplify *Staphylococcus aureus* 16S rRNA gene using PCR.

| Gene       | Primer type | Sequences          | Primer length | PCR Product size |
|------------|-------------|--------------------|---------------|-----------------|
| 16S rRNA   | Forward     | AACCTACCTATAAGACTGGG | 20nt          | 570 bp          |
| PCR Primer |             |                    |               |                 |
|            | Reverse     | CATTTCACCCTACACATGG | 20nt          |                 |
Amplification of *Staphylococcus aureus* 16S rRNA gene using PCR

Prepared PCR reaction mixtures were prepared as mentioned in material and method section. Amplification of 16S rRNA gene was performed in a PCR Thermo Cyclers [MJ Research PTC 200] for 30 cycles by using the following parameters: denaturation at 95°C for 60 second, annealing at 54°C for 45 second, and extension at 72°C for 2 min. The cycles were preceded by a denaturation step at 95°C for 5 min, followed by an extension step at 72°C for 4 min (Figure 2).

**Figure 2:** Adopted PCR condition to amplify the 16S rRNA gene sequence.

Visualization of PCR products through agarose gel electrophoresis

After amplification by PCR, the 570 bp of PCR products of *Staphylococcus aureus* 16S rRNA gene obtained were run on an agarose gel 1%, at 100-120V. The gel was stained with ethidium bromide. The PCR products were then visualized under UV light in Tran illuminator (Safe Imager 2.0 Blue-Light Transilluminator, ThermoFisher, Scientific, MA, USA).

**Figure 3:** Agarose gel electrophoresis (1%) represent amplification of 16S rRNA gene using PCR.

Local sequence alignment

As mentioned in material and method session, we have sequenced 3 isolate of *Staphylococcus aureus* 16S rRNA and different isolates of *Staphylococcus aureus* 16S rRNA gene sequences were retrieved from the NCBI (https://www.ncbi.nlm.nih.gov/nuccore) in FASTA format, and performed local sequence alignment using online NCBI BLAST tool ([https://blast.ncbi.nlm.nih.gov/Blast.cgi](https://blast.ncbi.nlm.nih.gov/Blast.cgi)). After performing BLAST, the NCBI BLAST tool produced BLAST table (list of the aligned sequence) showing the description of the gene, accession numbers, percent similarity, e-value, etc (Table 2).
Table 2: BLAST table of 16s RNA of Staphylococcus aureus

| S. No | Description | Max Score | Total Score | % Identity | Ref No./Accession |
|-------|-------------|-----------|-------------|------------|-------------------|
| 1.    | 16S ribosomal RNA New Sequence 1 | 1068 | 1057 | 100 | New Seq 1 |
| 2.    | 16S ribosomal RNA New Sequence 2 | 1068 | 1068 | 100 | New Seq 2 |
| 3.    | 16S ribosomal RNA New Sequence 3 | 1068 | 1068 | 100 | New Seq 3 |
| 4.    | Staphylococcus aureus strain sam2 16S rRNA gene, | 1068 | 1068 | 100 | New Seq 4 |
| 5.    | Staphylococcus aureus strain OS 16S rRNA gene, | 1068 | 1068 | 100 | New Seq 5 |
| 6.    | Staphylococcus aureus strain S1 16S rRNA gene, | 1068 | 1068 | 100 | New Seq 6 |
| 7.    | Staphylococcus aureus strain RB3 16S rRNA gene, | 1068 | 1068 | 100 | New Seq 7 |
| 8.    | Staphylococcus aureus strain ATCC 12600 16S rRNA, | 1068 | 1068 | 100 | New Seq 8 |
| 9.    | Staphylococcus aureus strain PDS17 16S rRNA gene, | 1068 | 1068 | 100 | New Seq 9 |
| 10.   | Uncultured bacterium clone ncd154d09c1 16S rRNA gene, | 1068 | 1068 | 100 | New Seq 10 |
| 11.   | Staphylococcus aureus strain S11 16S rRNA gene, | 1068 | 1068 | 100 | New Seq 11 |
| 12.   | S. aureus gene for 16S rRNA | 1068 | 1068 | 100 | New Seq 12 |
| 13.   | Uncultured bacterium clone nck261c02c1 16S rRNA gene, | 1062 | 1062 | 99.83 | New Seq 13 |
| 14.   | Uncultured bacterium nck159c1 1c1 16S rRNA gene, | 1062 | 1062 | 99.83 | New Seq 14 |
| 15.   | Uncultured bacterium clone nck138b11c1 16S rRNA gene, | 1062 | 1062 | 99.83 | New Seq 15 |
| 16.   | Uncultured bacterium clone nck121d02c2 16S rRNA gene, | 1062 | 1062 | 99.83 | New Seq 16 |
| 17.   | Uncultured bacterium clone ncd2387a04c1 16S rRNA gene | 1062 | 1062 | 99.83 | New Seq 17 |
| 18.   | Uncultured bacterium clone ncd2370e04c2 16S rRNA gene | 1062 | 1062 | 99.83 | New Seq 18 |
| 19.   | Uncultured bacterium clone ncd2357c11c1 16S rRNA gene | 1062 | 1062 | 99.83 | New Seq 19 |
| 20.   | Uncultured bacterium clone ncd2304e08c1 16S rRNA gene | 1062 | 1062 | 99.83 | New Seq 20 |
| 21.   | Uncultured bacterium clone ncd1985a06c1 16S rRNA gene | 1062 | 1062 | 99.83 | New Seq 21 |
| 22.   | Uncultured bacterium clone ncd1925e04c1 16S rRNA gene | 1062 | 1062 | 99.83 | New Seq 22 |
| 23.   | Uncultured bacterium clone ncd1368b07c1 16S rRNA gene | 1062 | 1062 | 99.83 | New Seq 23 |
| 24.   | Uncultured bacterium clone ncd1367a09c1 16S rRNA gene | 1062 | 1062 | 99.83 | New Seq 24 |
| 25.   | Uncultured Staphylococcus sp. clone VA21_59 16S rRNA | 1062 | 1062 | 99.83 | New Seq 25 |
| 26.   | Uncultured bacterium clone ncd943g01c1 16S rRNA gene | 1062 | 1062 | 99.83 | New Seq 26 |
| 27.   | Uncultured bacterium clone ncd886e03c1 16S rRNA gene | 1062 | 1062 | 99.83 | New Seq 27 |
| 28.   | Uncultured bacterium clone ncd858b10c1 16S rRNA gene | 1062 | 1062 | 99.83 | New Seq 28 |
| 29.   | Uncultured bacterium clone ncd702c03c1 16S rRNA gene | 1062 | 1062 | 99.83 | New Seq 29 |
| 30.   | Uncultured bacterium clone ncd672d02c1 16S rRNA gene | 1062 | 1062 | 99.83 | New Seq 30 |
| 31.   | Uncultured bacterium clone ncd739g04c1 16S rRNA gene | 1062 | 1062 | 99.83 | New Seq 31 |
| 32.   | Uncultured bacterium clone ncd610b11c1 16S rRNA gene | 1062 | 1062 | 99.83 | New Seq 32 |
| 33.   | Uncultured bacterium clone ncd597e08c1 16S rRNA gene | 1062 | 1062 | 99.83 | New Seq 33 |
| 34.   | Uncultured bacterium clone ncd586e07c1 16S rRNA gene | 1062 | 1062 | 99.83 | New Seq 34 |
| 35.   | Uncultured bacterium clone ncd585f03c1 16S rRNA gene | 1062 | 1062 | 99.83 | New Seq 35 |
| 36.   | Uncultured bacterium clone ncd578e03c1 16S rRNA gene | 1062 | 1062 | 99.83 | New Seq 36 |
| 37.   | Uncultured bacterium clone ncd567b11c1 16S rRNA gene | 1062 | 1062 | 99.83 | New Seq 37 |
| 38.   | Uncultured bacterium clone ncd1480e09c1 16S rRNA gene | 1062 | 1062 | 99.83 | New Seq 38 |
| 39.   | Uncultured bacterium clone ncd32e11c1 16S rRNA gene | 1062 | 1062 | 99.83 | New Seq 39 |
| 40.   | Staphylococcus aureus strain SAW1, complete genome | 1057 | 1057 | 99.65 | CP047651.1 |
| 41.   | Staphylococcus aureus strain UP_338 chromosome, genome | 1057 | 1057 | 99.65 | CP047651.1 |
| 42.   | Staphylococcus aureus strain UP_1442, genome | 1057 | 1057 | 99.65 | CP047802.1 |
| 43.   | Staphylococcus aureus strain UP_1106, genome | 1057 | 1057 | 99.65 | CP047802.1 |
| 44.   | Staphylococcus aureus strain UP_1322, genome | 1057 | 1057 | 99.65 | CP047802.1 |
| 45.   | Staphylococcus aureus strain UP_1525, genome | 1057 | 1057 | 99.65 | CP047802.1 |
| 46.   | Staphylococcus aureus strain UP_322, genome | 1057 | 1057 | 99.65 | CP047802.1 |
| 47.   | Staphylococcus aureus strain UP_1097, genome | 1057 | 1057 | 99.65 | CP047802.1 |
| 48.   | Staphylococcus aureus strain UP_1405, genome | 1057 | 1057 | 99.65 | CP047802.1 |
| 49.   | Staphylococcus aureus strain UP_296, genome | 1057 | 1057 | 99.65 | CP047802.1 |

It is clear from the BLAST results that newly sequenced Staphylococcus aureus 16S rRNA gene have shown 100% identity with different Staphylococcus aureus 16S rRNA genes submitted at NCBI GenBank such as, Staphylococcus aureus strain sam2 16S ribosomal RNA gene (MN540925.1), Staphylococcus aureus strain OS 16S ribosomal RNA gene (MN508958.1), Staphylococcus aureus strain S1 16S ribosomal RNA gene (MK881023.1), Staphylococcus aureus
strain RB3 16S ribosomal RNA gene (MK271755.1), Staphylococcus aureus strain ATCC 12600 16S ribosomal RNA (NR_118997.2), Staphylococcus aureus strain FDS17 16S ribosomal RNA gene (FJ434470.1) and Staphylococcus aureus strain S11 16S ribosomal RNA gene (FJ434470.1). However, other uncultured 16S rRNA gene sequences with accession number KF099686.1, KF092179.1, KF090696.1, KF09615.1, JF208017.1, JF207012.1, JF199790.1, JF197863.1, JF172707.1, JF165386.1, JF119960.1, JF119857.1, HM077151.1, HM330051.1, HM307769.1, HM297895.1, HM291954.1, HM289952.1, HM289008.1, HM285212.1, HM284435.1, HM280583.1, HM280506.1, HM280017.1, HM279200.1, HM260038.1, HM247340.1, CP045682.1, CP047851.1, CP047802.1, CP047863.1, CP047861.1, CP047854.1, CP047799.1, CP047803.1, CP047860.1 and CP047865.1 shown 99.65 to 99.83 homology with newly sequenced 16S rRNA of Staphylococcus aureus.

Phylogenetic analysis

The phylogenetic analysis included the newly sequenced Staphylococcus aureus 16S RNA and retrieved 16S RNA sequences of different isolates from NCBI database. Staphylococcus aureus 16S rRNA gene sequences alignments were generated using MEGA7 (ver 7.0.26) tool. Individual dendrograms were generated using different methods, namely the maximum likelihood methods. Phylogenetic groups and subgroups were defined by the length and branching order of the concatenated gene tree. The resulting groups were supported by high bootstrap values.

In Phylogenetic analysis, alignment of nucleotide sequences is a major consideration, particularly in studies of genes from divergent taxa. It seems obvious to state that the phylogenetic analysis of sequences begins with the appropriate alignment of the data themselves, yet alignment remains one of the most difficult and poorly understood facets of molecular data analysis. Alignments of the genomic sequences are required to analyze the phylogenetic tree. Phylogenetic analysis often includes the search for evidence of directional selection in molecular evolution. Evolution of the 16S RNA was studied in different isolates of Staphylococcus aureus and adaptive changes were in the sequences. The phylogenetic analysis of the Staphylococcus aureus 16S rRNA gene dataset resulted in a tree consistent with modern systematic understanding of the relatedness among different species of Staphylococcus genus, mainly based on DNA sequences homology (Table 5).

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In order to determine the genus of the bacterial isolates collected from the different food samples, we have amplified and sequenced the 16S RNA gene from bacterial group. The obtained sequences were BLAST against NCBI's 16S rRNA GenBank. The evolutionary history was inferred by using the Maximum Likelihood method based on the Tamura-Nei model. The phylogenetic analysis performed using 39 sequences of 16S rRNA gene from newly and retrieved 16S rRNA sequences, including Staphylococcus aureus. The consensus tree inferred from 10 most parsimonious trees is shown. Branches corresponding to partitions reproduced in less than 50% trees are collapsed. The consistency index is 1.000000 (1.000000), the retention index is 1.000000 (1.000000), and the composite index is 1.000000 (1.000000) for all sites and parsimony-informative sites (in parentheses). The MP tree was obtained using the Subtree-Pruning-Regrafting (SPR) algorithm with search level 0 in which the initial trees were obtained by the random addition of sequences (10 replicates). The tree is drawn to scale with branch lengths calculated using the average pathway method and are in the units of the number of changes over the whole sequence. The analysis involved 39 nucleotide sequences. Codon positions included were 1st+2nd+3rd+Noncoding. All positions containing gaps and missing data were eliminated. There were a total of 578 positions in the final dataset. Evolutionary analyses were conducted in MEGA7.

Figure 4. Molecular Phylogenetic analysis of 16S rRNA gene using Maximum Likelihood method. The evolutionary history was inferred using the Maximum Parsimony method and evolutionary analyses were conducted in MEGA7.
The phylogenetic trees were constructed using Maximum likelihood method for the sequence of newly isolated bacteria from the food specimens. Maximum likelihood method is most suitable model to understand the evolutionary history of an organism. The bootstrap consensus trees inferred from 1000 replicates were taken to represent the evolutionary history of the taxa analyzed. The Maximum likelihood trees were obtained using the Nearest Neighbor-Interchange heuristic algorithm. All positions containing gaps and missing data were eliminated from the dataset (Complete Deletion option). Phylogenetic analyses were conducted in MEGA7 and obtained three major cluster (depicted in Figure 4), classified as Clade A (Red color), Clade B (Green Color) and Clade C (Blue Color). As depicted in figure 4, newly sequenced 16S rRNA gene sequences were grouped with other isolates of Staphylococcus aureus in Clade A, with 100% homology obtained similar as in local alignment analysis. Clade B and Clade C shown 99% homology with Clade A. however, both clades include different isolates of uncultured Staphylococcus species. The observations on phylogenetic analysis of 16S RNA gene of Staphylococcus aureus using Maximum Likelihood method revealed the relationships and percent similarity of 16S RNA gene within different bacterial isolates, including Staphylococcus species. Observations based on molecular techniques verified the major presence of Staphylococcus aureus in collected food specimens.

CONCLUSION:
Phylogenetic analysis of the Staphylococcus species, including new isolates from food specimens revealed that they are the same strain and are affiliated to Staphylococcus aureus. In recent years, Next Generation of Sequencing technologies boosted the genome databases and a remarkable increase in the number of sequenced genomes, drafts or complete, are available, but the correct assignment of the sequenced strains to the corresponding species with the accepted taxonomic tools is important before comparative analyses with other genomes can be performed. The need for the whole genome sequences of all the type strains, which are the only species references that are publicly available in culture collections, is evident. In the present study, we have identified and characterized Staphylococcus aureus from the clinical specimens, using molecular biology techniques. New 16sRNA sequences of Staphylococcus aureus isolated were aligned with Staphylococcus species and constructed phylogeny tree to determine the molecular evolution and population structure of Staphylococcus species using bioinformatics tools. The phylogenetic affiliations of the different species of the genus Staphylococcus aureus were shown by the Maximum likelihood based phylogenetic analyses using the 16S rRNA sequences. Our study demonstrated that positive selection of 16S RNA gene during the divergence of different isolate of Staphylococcus species during evolution. These evolutionary acquisitions have made necessary changes in the genetic control of ontogeny, and this, in turn, might have caused adaptive changes in the 16S RNA gene.

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Conflicts of Interest Statement
The authors declare no conflict of interests.
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