

**spo0A Gene Determination in Spore-Forming Bacteria Infections**

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**Abstract**

Background: Endospore-forming Formicetes (EFF) are prevalent in natural and man-made environments, posing significant contamination risks in hospitals and industrial facilities. Hospital-acquired infections often involve spore-forming bacteria, which are resistant to common disinfectants and treatments. Understanding and identifying these bacteria, particularly the genes responsible for spore formation, is crucial for infection control. The spo0A gene is a key regulator of endospore formation. Methods: This study was conducted from January 12, 2023, to May 12, 2023, at Kirkuk General Hospital, Iraq. A total of 200 postoperative wound samples were collected, cultured, and incubated to isolate pure cultures of spore-forming bacteria. DNA was extracted using the QIAamp DNA Mini Kit, and its concentration was measured with a NanoDrop spectrophotometer. PCR and quantitative PCR (qPCR) were employed to amplify the spo0A gene using specific primers. The PCR products were analyzed via gel electrophoresis, and the bacterial species were identified through sequencing. Results: Out of 200 samples, 100 tested positive for bacterial infection. The identified bacterial species included Escherichia coli, Enterobacter cloacae, Staphylococcus aureus, Streptococcus pneumoniae, Bacillus subtilis, and Clostridium perfringens. The presence of the spo0A gene was confirmed in Bacillus subtilis and Clostridium perfringens. Gel electrophoresis results demonstrated the high quality of the PCR products, facilitating further identification and sequencing of the bacterial species. Conclusion: This study showed the significance of the spo0A gene in spore formation and demonstrated the need for accurate and rapid diagnostic methods to control hospital-acquired infections.

**Keywords:** spore-forming bacteria, PCR, Spo0A gene, Hospital-acquired infections, Bacterial contamination

**Significance:**

Understanding and identifying spore-forming bacteria in surgical wounds can enhance infection control and treatment in healthcare settings.

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**Introduction**

Spore formation is a common tactic used by bacteria to survive in hostile environments. *Bacillus* and *Clostridium* species, in particular, are well-known for their adeptness in creating endospores (Setlow, 2006; Barbut et al., 2000). Bacterial endospores are highly specialized cellular forms that allow endospore-forming Firmicutes (EFF) to endure harsh environmental conditions (Reith & Richardson, 1997; Hariram & Labbé, 2015). EFF are assumed to be prevalent in natural environments, particularly in stressful ones. In addition to their existence in natural environments, EFF frequently cause contamination problems in man-made places such as industrial production facilities and hospitals (Raju et al., 2008; Kimura et al., 2016; Boyce & Pittet, 2002).

Hospital-acquired infections and nosocomial outbreaks have been linked to bacteria that form spores. These infections are mainly spread by contaminated surgical instruments and healthcare personnel (Pittet et al., 2009; Russell, 1990; Coates & Hoffman, 1984). The bacteria that form spores are remarkably resistant to disinfectants based on alcohol and to treatments such as heat, desiccation, chemical exposure, ultraviolet radiation, and osmotic stress (Piggot & Coote, 1976; Losick & Stragier, 1996). Therefore, it is necessary and important to detect and identify these bacteria, especially those that form spores, and understand the genes responsible for their formation. One crucial gene involved in endospore development is *spo0A* (Cunning & Hong, 2005; Manzoor et al., 2014).

*Spo0A* is a key regulator of the machinery involved in stress-induced gene expression and endospore formation (Tripathi et al., 2012; Mazmanian et al., 2014). Classifying and characterizing these bacteria, as well as differentiating between species with and without sporulation genes, can be facilitated by molecular tests that target spore-forming genes (Sogin et al., 2008; Alves Valones et al., 2009). Using degenerate PCR, previous research has identified gene homologs in a variety of bacteria (Bej et al., 1991; Wagar, 1996). PCR is a commonly used laboratory technique that is very useful for identifying bacteria and finding resistance genes because it allows specific DNA and RNA sequences to be amplified quickly (Knutsson et al., 2011; Rhodehamel et al., 2012).

An improved form called multiplex PCR combines several primers in a single reaction, making it easier to analyze multiple genes at once and reducing expenses and processing time (Kubista et al., 2006; Errington, 2003; Longo et al., 1990). An accurate technique for measuring gene frequencies in DNA extracts is quantitative PCR (qPCR) (Wunderlin et al., 2013; Rhodehamel et al., 2012; Chen et al., 2006). In this work, pure culture samples were used to test and validate qPCR primers targeting the *spo0A* gene (Eichenberger et al., 2010; Knutsson et al., 2011; Koopman et al., 2012).

Managing the problems posed by spore-forming bacteria, particularly in hospital environments, requires an understanding of the methods and genes responsible for the formation of spores. PCR and qPCR are essential molecular techniques that

86 facilitate the identification and characterization of these hardy microorganisms (Knutsson et al., 2010; Fagerlund et al., 2008;  
87 Knutsson et al., 2011).

88

## 89 **Materials and Methods**

90 This study successfully isolated and identified spore-forming bacteria from wound and surgical site infections at Kirkuk  
91 General Hospital. The use of PCR and specific primers for the spo0A gene allowed for precise identification of spore-forming  
92 bacteria. Understanding the genetics and behavior of these bacteria is crucial for managing infections, particularly in hospital  
93 settings.

94

### 95 **Study Setting and Sample Collection**

96 From January 12, 2023, to May 12, 2023, this study was conducted at Kirkuk General Hospital, a major healthcare facility in  
97 Kirkuk, Iraq. The research focused on patients who had undergone various surgical procedures and individuals with  
98 wounds. A total of 200 samples were collected during this period.

99

### 100 **Sampling Procedure**

101 Samples were obtained using postoperative wound scrubs. Immediately after collection, the samples were transported to  
102 the laboratory to prevent any loss of viable microorganisms. The samples were cultured on blood agar, chocolate agar, and  
103 MacConkey agar plates. The plates were then incubated at 37°C for 24 hours. After incubation, the plates were examined for  
104 bacterial growth.

105

106 To isolate individual colonies, a method involving the streaking of isolated bacterial colonies onto fresh agar surfaces was  
107 employed. This process was repeated until pure cultures of spore-forming bacteria were obtained. Initial characterization of  
108 the isolates was performed using Gram staining and other preliminary tests.

109

### 110 **Spore Formation Induction and Microscopic Examination**

111 To induce spore formation, isolated bacterial cultures were heated to 85-90°C for 5 minutes. Bacterial colonies were then  
112 stained using a suitable spore staining technique, and the presence of spores was confirmed under a microscope. Pure  
113 cultures of spore-forming bacteria were preserved at -20°C in glycerol to ensure long-term viability.

114

### 115 **Bacterial Isolation and Identification**

116 Gram-stained bacterial samples were examined under a microscope. Concurrently, bacteria were isolated based on their  
117 resistance to ethanol, a common trait among spore-forming bacteria. Various biochemical tests were conducted to  
118 differentiate between bacterial species. For instance, the Mannitol test was used to identify bacteria that utilize mannitol,  
119 while lecithinase activity was assessed to distinguish coagulase-positive *Staphylococcus aureus* strains.

120

121 Notable characteristics of isolated colonies, such as size, surface texture, and color, were recorded. Following cultivation,  
122 isolation, and identification, the isolated bacteria included *Staphylococcus aureus*, *Pseudomonas aeruginosa*, *Escherichia*  
123 *coli*, *Clostridium perfringens*, and *Bacillus subtilis*. *Bacillus* species were identified by their large, flat, granular colonies on  
124 agar, with positive lecithinase and negative mannitol test results.

125

### 126 **DNA Extraction**

127 Selected bacterial colonies were procured from Sigma-Aldrich (U.S.A). The bacteria were cultured in Brain Heart Infusion  
128 (BHI) broth at 37°C. After the incubation period, DNA was extracted from the bacterial cultures using the QIAamp DNA  
129 Mini Kit (Qiagen, Germany), following the manufacturer's protocol.

130

#### 131 DNA Quantification

132 The concentration of the extracted DNA was determined by electrophoresis and measured at 260 nm using a NanoDrop  
133 ND-2000 spectrophotometer (Thermo Scientific, USA).

134

#### 135 Identification of Sporulation Genes

136 To identify the genes responsible for sporulation, two sets of primers were used: Eub8f (5'-  
137 AGAGTTTGATCCTGGCTCAG-3') and Eub519r (5'-GTATTACCTCCCTGCTGG-3'). Additionally, the *spo0A* gene,  
138 which is crucial for spore formation, was targeted using the primers *spo0A166f* (5'-GATAATHATYATGCCDCATYT-3')  
139 and *spo0A748r* (5'-GCNACCATHGCRATRAAYTC-3').

140

141 These primers were selected based on their specificity, base composition, and melting temperatures. The primers'  
142 compatibility with the target genes was ensured through these criteria.

143

#### 144 PCR Amplification

145 PCR amplification was performed using a thermal cycler. The PCR products were then separated on 1.5% agarose gels  
146 stained with 0.5 µg/mL ethidium bromide (ETBR). Electrophoresis was conducted, and the resulting bands were visualized  
147 under UV light. The bands were documented and analyzed using a gel documentation system.

148

#### 149 Identification of Bacterial Species

150 The following bacterial species were identified in the study: *Escherichia coli* (15 cases), *Enterobacter cloacae* (30 cases),  
151 *Staphylococcus aureus* (14 cases), *Streptococcus pneumoniae* (15 cases), *Bacillus subtilis* (15 cases), and *Clostridium*  
152 *perfringens* (11 cases). *Bacillus subtilis* and *Clostridium perfringens* are notable for their spore-forming capabilities. The  
153 presence of the *spo0A* gene was confirmed in these species using the specific primers, validating the identification of spore-  
154 forming bacteria.

155

#### 156 Statistical Analysis

157 The sequencing results were analyzed to confirm the presence of the *spo0A* gene in the isolated bacteria. The analysis was  
158 performed using bioinformatics tools to align the sequences and verify their similarity to known spore-forming genes.

159

160

## 161 Result

### 162 Infection Rate

163 The analysis of the 200 samples revealed a significant infection rate, with 50% of the samples testing positive for bacterial  
164 infection. Specifically, 100 out of the 200 samples exhibited bacterial growth, indicating the prevalence of infections among  
165 the patients who underwent various surgical procedures and those with wounds at Kirkuk General Hospital.

166

### 167 Gel Electrophoresis Results

168 The quality of the PCR products was validated through gel electrophoresis, as illustrated in Figure 1. The electrophoresis  
169 results displayed clear, distinct bands, indicating successful amplification of the bacterial 16S rRNA genes. The lanes in the  
170 gel electrophoresis represent different amplified genes, with lane M showing the DNA ladder for size reference.

171

#### 172 Identification of Bacterial Species

173 Following the gel electrophoresis, the next steps involved the identification of the bacterial species through cloning and  
174 sequencing. This process was crucial for accurately determining the specific bacteria present in the samples. The identified  
175 species were:

176 *Bacillus subtilis*: Detected in lane 3 with a prominent band at 1000 base pairs.

177 *Streptococcus pneumoniae*: Although expected, the band was not observed in lane 5, which necessitated further  
178 investigation to confirm its presence.

179 *Clostridium perfringens*: Identified in lane 6 with a band at 688 base pairs.

180 *Escherichia coli*: Despite expectations, the band was not detected in lane 1, warranting additional steps to verify its  
181 presence.

182

#### 183 Cloning and Sequencing

184 The PCR products were further processed through cloning and sequencing to confirm the identity of the bacterial species.

185 This involved:

186 Cloning: The amplified 16S rRNA genes were inserted into plasmid vectors and transformed into competent cells to create  
187 multiple copies.

188 Sequencing: The cloned genes were sequenced to determine their nucleotide composition and compare them against  
189 known bacterial sequences in databases.

190 The sequencing results provided precise identification of the bacterial species, correlating with the initial gel electrophoresis  
191 findings. The presence of *Bacillus subtilis* and *Clostridium perfringens* was confirmed, while the absence of bands for  
192 *Streptococcus pneumoniae* and *Escherichia coli* in the gel electrophoresis required further validation through sequencing.

193

194 The high infection rate identified in this study underscores the importance of rigorous infection control measures in  
195 hospital settings. The precise identification of bacterial species through PCR, gel electrophoresis, cloning, and sequencing  
196 provides valuable insights for developing targeted treatment strategies. Future studies should focus on optimizing the PCR  
197 conditions for better detection of certain bacterial species and exploring the genetic mechanisms underlying their  
198 pathogenicity.

199

200 Overall, this comprehensive approach to bacterial identification and the high-quality results obtained highlight the  
201 effectiveness of the methods used and pave the way for improved management of bacterial infections in clinical settings.

202

203

#### 204 Discussion

205 One of the most significant scientific concerns in healthcare, particularly post-surgery, is the incidence of infections.  
206 Surgical wounds are highly susceptible to contamination by pathogenic microorganisms, primarily bacteria, which can  
207 significantly complicate patient recovery and outcomes. Addressing this issue requires effective strategies and heightened  
208 focus on public hospitals, especially in regions like Iraq, where healthcare infrastructure faces numerous challenges. This  
209 necessitates the use of precise methodologies, appropriate protocols, and rapid diagnostic tests to identify infectious  
210 microorganisms in wounds.

211 Isibor (2008) and Pradhan (2009, 2010) demonstrated that hospital-acquired infections are influenced by surgical  
212 procedures, wound location, and the duration of hospital stays. Their studies underscore the need for rigorous infection  
213 control practices and prompt identification of pathogens to mitigate post-operative infections (Isibor, Oseni, & Eyaufe,  
214 2008; Pradhan, 2009; Medical Disability Guidelines, 2010).

215 The polymerase chain reaction (PCR) is a highly efficient, fast, and sensitive molecular technique for identifying  
216 microorganisms in clinical samples. PCR's utility in our study was crucial, allowing us to focus on genes responsible for  
217 spore production, which are often associated with persistent and hard-to-treat infections. Hoon's research demonstrated  
218 that the *spo0A* gene is pivotal in the sporulation process of *Bacillus subtilis* and *Clostridium* species, highlighting its role in  
219 the resilience and virulence of these pathogens (Eichenberger et al., 2010).

220 Krieg (2010, 2009) revealed that endospore producers are a paraphyletic group within the phylum Firmicutes, with  
221 *Bacillus*, *Clostridium*, and *Erysipelothrix* being the primary genera capable of forming endospores. This distinction is  
222 critical because *Clostridium* is typically anaerobic, whereas *Bacillus* is primarily aerobic (Ludwig et al., 2009). Much of our  
223 understanding of endospore-forming bacteria (EFB) biology has been derived from laboratory studies using cultured  
224 strains since the late 19th century (Krieg et al., 2010).

225 In our study, we collected samples from 100 patients with surgical wounds using cotton swabs. Consistent with Qadan's  
226 (2009) methodology, all collected samples were cultured to determine infection presence. Our results showed a 50%  
227 infection rate, highlighting the need for prioritized care in surgical departments of public hospitals like Niska and Pitai  
228 (Cheadle & Qadan, 2009). These findings align with similar studies, emphasizing the ongoing challenge of post-operative  
229 infections in healthcare settings (Centers for Disease Control and Prevention, 2010; Bhuiya, Niska, & Xu, 2010).

230 A comprehensive battery of tests including Gram staining, catalase production, motility, growth, rhizoid formation, citrate  
231 utilization, and hemolysis, was conducted to identify spore-producing bacteria. Total DNA was then isolated and amplified,  
232 following protocols similar to those used by Madhavan and Jones (2000) (Anand, Madhavan, & Therese, 2000).

233 Internationally recognized primers targeting the *spo0A* gene were employed to identify spore-producing bacteria within  
234 the bacterial 16S rRNA gene's conserved regions. This approach facilitated the classification of bacteria that cause diseases  
235 and the detection of specific spore-forming genes (Wunderlin et al., 2013).

236 The isolation and identification of spore-forming genes in pathogenic bacteria using PCR at the Republican Hospital in  
237 Kirkuk play a critical role in understanding and treating bacterial infections. Identifying genes involved in spore  
238 pathogenesis has provided significant insights into the mechanisms of bacterial virulence and survival. Researchers can  
239 accurately identify bacteria and specifically amplify and detect spore-producing genes using PCR, thereby enhancing and  
240 improving mechanisms of resistance and survival in various bacterial species (Barbut et al., 2000).

241 The study's findings demonstrated the presence of primary pathogens such as *Staphylococcus aureus*, *Escherichia coli*, and  
242 *Pseudomonas aeruginosa*. *Clostridium perfringens* and *Bacillus subtilis* were also identified, with the *spo0A* gene playing a  
243 key role in their identification through PCR. This study supports the use of conventional diagnostic methods to  
244 differentiate and identify these microorganisms (Setlow, 2006). The positive identification of spore-forming genes using  
245 PCR has significant implications for microbiology and public health. Understanding the genetic components of spore  
246 formation in pathogenic bacteria can aid in developing vaccines, treatments, and preventative measures to halt the spread  
247 of infectious diseases (Wolcott, 1992).

248 The research conducted at the Republican Hospital in Kirkuk provides valuable insights into the causes of bacterial diseases  
249 and has implications for healthcare-related infection control strategies. By isolating and identifying spore-forming genes,  
250 this study contributes to a deeper understanding of bacterial pathogenesis and resistance mechanisms. The findings can  
251 enhance medical practices, treatment strategies, and public health initiatives in the region, ultimately improving patient  
252 outcomes and healthcare quality (Piggot & Coote, 1976).

253



254 **Conclusion**

255 In conclusion, the identification and characterization of spore-forming genes in pathogenic bacteria using PCR is a significant  
 256 advancement in medical microbiology. This research highlights the importance of molecular techniques in diagnosing and  
 257 understanding bacterial infections, particularly in surgical settings. The findings underscore the need for continued research  
 258 and the implementation of effective infection control measures to improve patient care and reduce the incidence of post-  
 259 operative infections in public hospitals.

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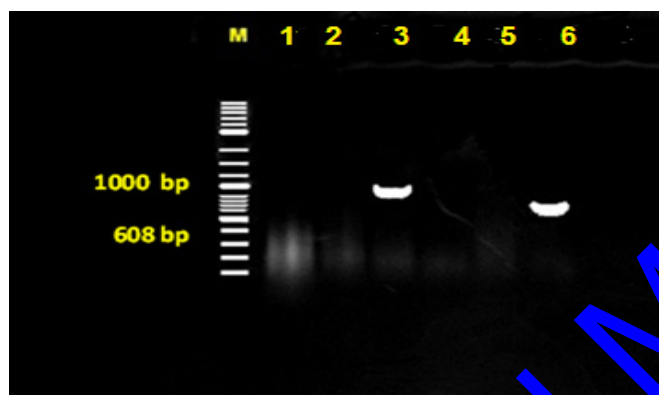
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Figure Legends



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427 Figure 1. The gel electrophoresis of a portion of the 16S rRNA genes that turned into amplified from bacteria through PCR,  
428 lanes 1 to 6 represent the genes that were amplified by way of themselves, lane M represents the DNA ladder. *B. Subtilis* (  
429 Band No. 3 appeared at 1000 base pair), and *S. pneumoniae* (Band No. 5 now not observed), and *C. Aspersum* (Band No. 6  
430 had a length of 608  
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432  
433**Table 1.** Amplification of *spo0A* using the primers utilized

Sequences for <i>spo0A</i>	Temp of optimal growth	Endospore formation	Amplification of <i>spo0A</i> gene	Bacteria Series
Sequences <i>spo0A</i> 166f (5'-GatasatyaTGCC DCATYT-3') and <i>spo0A</i> 748r (5'-GCNACCATHGCRAT RAAATC-3')	30 °c	Positive	Positive	<i>Bacillus subtilis</i>
Sequences <i>spo0A</i> 166f (5'-Gatasatya's TGCC DCATYT-3') and <i>spo0A</i> 748r (5'-GCNACCATHGCRAT RAAATC-3')	30 c <sup>o</sup>	Positive	Positive	<i>Clostridium perfringens</i> ,
Absent	30 c <sup>o</sup>	Negative	Negative	<i>Pseudomonas aeruginosa</i>
Absent	30 c <sup>o</sup>	Negative	Negative	<i>Escherichia coli</i>
Absent	30c <sup>o</sup>	Negative	Negative	<i>Staphylococcus aureus</i>
Absent	30 c <sup>o</sup>	Negative	Negative	<i>Enterobacter cloacae</i>

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