INTRODUCTION

Carbapenem-resistant Enterobacteriaceae (CRE) represent a serious growing threat to public health in Korea. Serious Infections with CRE are difficult to treat or non-treatable, and thus are associated with high mortality rates. Carbapenemase-mediated CRE is unique because of its capacity for horizontal transmission via a plasmid, by which carbapenem-resistance increases rapidly and crosses the species barrier [1]. In a previous study of five Korean university hospitals, ertapenem susceptibilities were 99.5% among Escherichia coli and 94.0% among Klebsiella pneumoniae in 2015 [2]. CRE has been designated as a legally notifiable disease since October.
2010, and Korea Centers for Disease Control and Prevention (KCDC) implemented a laboratory-based monitoring system in representative regional hospitals to collect and genotype CRE and CPE (Carbapenemase-producing Enterobacteriaceae) nationwide [1]. CRE has been considered as a group III notifiable disease since June 2017, and all CRE cases and patients harboring CRE should be reported to the KCDC [1].

CRE surveillance culture is an essential element in the control and prevention of CRE infection [3]. Because CRE infection rates have increased since the first four cases of NDM-producing K. pneumoniae were found at our hospital in 2010 [4], CRE surveillance culture has become a routine practice in many clinical microbiology laboratories in Korea. To accelerate the turn-around time of CRE surveillance culture, matrix assisted laser desorption ionization-time of flight mass spectrometry (MALDI-TOF MS, MALDI-TOF) was recently introduced for direct species identification of colonies on selective agar [5-7]. However, species identification of CRE grown on MacConkey agar containing imipenem (IMP-Mac) failed biochemical identification tests in this hospital.

This study was performed to evaluate the impact of the media type on species identification using MALDI-TOF in CRE surveillance cultures. The influence of colony aging on species identification was also measured for incubation times of 1 and 2 days.

MATERIALS AND METHODS

1. Specimens

For 2 months, March and April 2017, four types of media were evaluated: blood agar (BA; Komed, Seongnam, Korea), Mueller Hinton agar agar (MH, Komed), MacConkey agar (Mac, Komed), MacConkey agar containing imipenem (IMP-Mac). The samples subjected to CRE surveillance culture were inoculated on IMP-Mac and pink colonies were considered as CRE-likely colonies after overnight incubation at 36°C±1°C. Because routine practices in the laboratories use extended culture at room temperature to detect colonies in the case of holiday, the incubation was extended for one additional day at room temperature. All pink colonies on IMP-Mac were consecutively collected and subcultured on BA and Mac plates. Disk diffusion tests were conducted on MH agar. The colonies on each media type were tested for species identification using a MALDI biotyper (Bruker Daltonics, Bremen, Germany) and MicroScan NC72 panel (Beckman Coulter, Brea, CA, USA).

2. MALDI-TOF Identification

The colonies on each agar plate were directly prepared for MALDI-TOF identification. Briefly, a single colony was picked using a wooden toothpick and gently smeared onto a polished steel target plate. The spots were then overlaid with 1 μL of IVD matrix solution (Bruker Daltonics) consisting of 47.5% deionized water, 50% acetonitrile (Sigma-Aldrich, St. Louis, MO, USA) and 2.5% trifluoroacetic acid (Sigma-Aldrich) and allowed to dry. The MALDI biotyper operation was conducted according to the manufacturer’s instructions and species identification was based on Bruker Taxonomy with Mass Spectrometry Profiling. According to the manufacturer’s instructions, the identification results with score values ≥1.7 were considered reliable at the genus level. The correct identification was considered as showing consistent with results with MicroScan NC72 or Klebsiella oxytoca in MicroScan NC72 for Raoultella ornithinolytica identified by Biotyper. The successful identification rates (SIRs) and score values of the identification results were analyzed for each media type and incubation time.

3. Statistical Analysis

The SIRs of each media type were compared by chi-square tests. The score values of each specimen resulting in correct identifications were compared by paired t-test. Analysis was conducted using PASW SPSS ver. 18.0 software (SPSS Inc., Chicago, IL, USA). P-value of <0.05 was considered statistically significant.

RESULTS

A total of 117 CRE were detected, including 84 K. pneumoniae, 12 Escherichia coli, 9 Enterobacter cloaceae,
<table>
<thead>
<tr>
<th>Species</th>
<th>No. of isolate</th>
<th>Incubation time (day)</th>
<th>No. of successful identification (%)</th>
<th>Blood agar</th>
<th>Muller Hinton Agar</th>
<th>MacConkey Agar</th>
<th>IMP-MacConkey agar</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Klebsiella pneumoniae</strong></td>
<td>84</td>
<td>1</td>
<td>82/82 (97.6)/2 (2.4), 2.209±0.16 (82)</td>
<td>82/82 (97.6)/2 (2.4), 2.258±0.14 (82)</td>
<td>68/68 (81.0)/12 (14.3), 2.006±0.23 (72)</td>
<td>59/59 (70.2)/18 (21.4), 1.967±0.24 (66)</td>
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<td></td>
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<td>2</td>
<td>81/81 (96.4)/3 (3.6), 2.216±0.14 (81)</td>
<td>81/81 (96.4)/3 (3.6), 2.255±0.12 (81)</td>
<td>53/53 (63.1)/27 (32.1), 2.033±0.24 (57)</td>
<td>49/49 (58.3)/27 (32.1), 1.964±0.27 (57)</td>
<td></td>
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<tr>
<td><strong>Escherichia coli</strong></td>
<td>12</td>
<td>1</td>
<td>12/8 (91.7)/1 (8.3), 2.268±0.09 (11)</td>
<td>11/8 (91.7)/1 (8.3), 2.065±0.19 (11)</td>
<td>9/7 (75.0)/2 (16.7), 1.923±0.31 (10)</td>
<td>0.06/0.06 (0.06)</td>
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<td></td>
<td></td>
<td>2</td>
<td>12/12 (100)/0 (0), 2.294±0.09 (12)</td>
<td>11/11 (90.9)/1 (9.1), 2.199±0.13 (11)</td>
<td>8/7 (66.7)/4 (33.3), 2.023±0.19 (8)</td>
<td>8/8 (66.7)/4 (33.3), 1.989±0.18 (8)</td>
<td></td>
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<tr>
<td><strong>Entrobacter cloacae</strong></td>
<td>9</td>
<td>1</td>
<td>9/9 (100)/0 (0), 2.061±0.09 (9)</td>
<td>9/9 (100)/0 (0), 2.084±0.12 (9)</td>
<td>8/8 (88.8)/1 (11.2), 1.945±0.15 (8)</td>
<td>7/7 (77.8)/1 (22.2), 1.885±0.13 (8)</td>
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<tr>
<td></td>
<td></td>
<td>2</td>
<td>9/9 (100)/0 (0), 2.106±0.09 (9)</td>
<td>8/8 (88.8)/1 (11.2), 2.032±0.17 (9)</td>
<td>4/4 (44.4)/2 (55.6), 1.748±0.34 (7)</td>
<td>5/5 (55.5)/2 (44.4), 1.807±0.14 (7)</td>
<td></td>
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<tr>
<td><strong>Klebsiella oxytoca</strong></td>
<td>5</td>
<td>1</td>
<td>5/5 (100)/0 (0), 2.268±0.11 (5)</td>
<td>5/5 (100)/0 (0), 2.271±0.09 (5)</td>
<td>3/3 (60.0)/2 (40.0), 2.048±0.26 (4)</td>
<td>3/3 (60.0)/2 (40.0), 2.187±0.03 (3)</td>
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<td></td>
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<td>2</td>
<td>5/5 (100)/0 (0), 2.168±0.03 (5)</td>
<td>5/5 (100)/0 (0), 2.230±0.10 (5)</td>
<td>2/2 (100)/0 (0), 1.906±0.40 (3)</td>
<td>1/1 (100)/0 (0), 1.762±0.25 (2)</td>
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<tr>
<td><strong>Klebsiella aerogenes</strong></td>
<td>4</td>
<td>1</td>
<td>4/4 (100)/0 (0), 2.180±0.24 (4)</td>
<td>4/4 (100)/0 (0), 2.281±0.13 (4)</td>
<td>3/3 (75.0)/1 (25.0), 2.277±0.10 (3)</td>
<td>4/4 (100)/0 (0), 2.231±0.10 (4)</td>
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<tr>
<td></td>
<td></td>
<td>2</td>
<td>3/3 (75.0)/1 (25.0), 2.340±0.07 (3)</td>
<td>4/4 (100)/0 (0), 2.278±0.17 (4)</td>
<td>2/2 (50.0)/2 (50.0), 2.056±0.22 (2)</td>
<td>3/3 (75.0)/1 (25.0), 2.196±0.05 (3)</td>
<td></td>
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<tr>
<td><strong>Raoultella ornithinolytica</strong></td>
<td>2</td>
<td>1</td>
<td>2/2 (100)/0 (0), 2.515±0.01 (2)</td>
<td>2/2 (100)/0 (0), 2.509±0.04 (2)</td>
<td>2/2 (100)/0 (0), 1.962±0.08 (2)</td>
<td>1/1 (50.0)/1 (50.0), 2.371±0.00 (1)</td>
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<tr>
<td></td>
<td></td>
<td>2</td>
<td>2/2 (100)/0 (0), 2.482±0.05 (2)</td>
<td>2/2 (100)/0 (0), 2.470±0.07 (2)</td>
<td>2/2 (100)/0 (0), 2.461±0.01 (2)</td>
<td>2/2 (100)/0 (0), 2.155±0.04 (2)</td>
<td></td>
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<tr>
<td><strong>Citrobacter freundii</strong></td>
<td>1</td>
<td>1</td>
<td>1/1 (100)/0 (0), 2.200 (1)</td>
<td>1/1 (100)/0 (0), 2.438 (1)</td>
<td>1/1 (100)/0 (0), 2.073 (1)</td>
<td>1/1 (100)/0 (0), 1.977 (1)</td>
<td></td>
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<td></td>
<td></td>
<td>2</td>
<td>1/1 (100)/0 (0), 2.301 (1)</td>
<td>1/1 (100)/0 (0), 2.311 (1)</td>
<td>1/1 (100)/0 (0), 2.116 (1)</td>
<td>1/1 (100)/0 (0), 2.261 (1)</td>
<td></td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>117</td>
<td>1</td>
<td>115/115 (98.3)/2 (1.7), 2.213±0.16 (115)</td>
<td>114/114 (97.4)/3 (2.6), 2.300±0.14 (114)</td>
<td>96/96 (82.1)/16 (13.7), 2.017±0.22 (101)</td>
<td>83/83 (70.9)/24 (20.5), 1.978±0.24 (83)</td>
<td>P&lt;0.001/P&lt;0.001</td>
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<td></td>
<td></td>
<td>2</td>
<td>113/113 (96.6)/4 (3.4), 2.223±0.14 (113)</td>
<td>113/113 (96.6)/4 (3.4), 2.236±0.14 (113)</td>
<td>72/72 (61.5)/37 (31.6), 2.015±0.27 (80)</td>
<td>68/68 (51.3)/37 (31.6), 1.965±0.25 (80)</td>
<td>P&lt;0.001/P&lt;0.001</td>
</tr>
</tbody>
</table>

Abbreviations: SD, standard deviation; IMP-MacConkey agar, MacConkey agar containing 1 µg/mL imipenem.

*aRaoultella ornithinolytica* was identified by MALDI-TOF, *P*-values of each results of 1 and 2 days were calculated by comparison of successful identification rates/score value of each media type to those of BAP on the same day of incubation, and *P*-values of 1 vs. 2 days results were calculated by comparison of successful identification rates/score values of 2 days results to those of 1 day on each media.
5 K. oxytoca, 4 Klebsiella aerogenes, 2 R. ornithinolytica, and 1 Citrobacter freundii. The MALDI-TOF identification results are shown in Table 1. At overnight incubation, 98.3% of isolates were correctly identified from the colonies on BA, 97.4% on MH (P=0.9), 82.1% (P<0.001) on MAC, and 70.9% (P<0.001) on IMP-MAC. After 2 days of incubation, the SIRs were decreased to 61.5% (P<0.001) on MAC and 58.1% (P<0.001) on IMP-MAC, but not on BA and on MH which showed values of 96.6% and 96.6%, respectively. Unsuccessful identification mainly resulted from the lack of a peak, and the number of samples without peaks was increased after 2 days of incubation: 2 to 4 in BA, 3 to 4 in MH, 16 to 37 in Mac, and 24 to 37 in IMP-Mac (Table 1). The average score values for correctly identified specimens were significantly lower for Mac (2.017±0.22, P<0.001) and IMP-Mac (1.978±0.24, P<0.001), while there was no significant difference between those for MH (2.300±0.14) and BA (2.213±0.16). After 2-day incubation, the score values were slightly lower for each media type than those after day 1, but the differences were not significant among the specimens identified correctly except for E. coli on Mac and E. cloacae on IMP-Mac (Table 1). Among species for which 4 or more isolates were examined in this study, identification performance of the Biotyper was significantly lower when colonies grown on Mac or IMP-Mac were used, except for E. cloacae. The SIRs of colonies grown on Mac were further decreased by colony aging, except for E. coli, while those grown on BA and MH were not (Table 1).

DISCUSSION

In this study, SIRs were significantly lower for colonies grown on Mac compared to those grown on BA and MH. The identification rate was further decreased when colonies on IMP-Mac were used. Because microbial identification by MALDI-TOF is based on unique protein signatures, certain media types and supplements may interfere with identification [6,8,9]. In a previous study, the type of media was found to affect the identification performance of MALDI-TOF MS for enteric gram-negative rods (EGNR), while growth on Mac showed similar performance as growth on BA, even when using the direct colony method [10]. Pseudomonas species are affected by growth on Mac more significantly when using the direct method described in the previous study. The reason for the difference in SIRs between media is unclear. The type of media is thought to affect the cell wall condition, making the colonies more mucoid in Pseudomonas species and affecting identification performance using the direct method [11]. This explanation may also apply to EGNR such as K. pneumoniae, which shows the greatest mucoid character and is the most prevalent CRE. Pigmented media such as Mac may affect protein spectral profiles, as has been observed for fungal pigments [12]. Additionally, bile salts in Mac may block ionization effects during examination [11]. In conclusion, species identification of EGNR by MALDI-TOF MS using the direct method was affected by the type of media used: growth on BA and MH is preferred to growth on Mac and IMP-Mac.

Colony aging also affected the success rate of identification when the colonies were grown on Mac. Colony aging from 5 to 48 hours did not impact identification performance [11]. Thus, very short-term incubation of 4 hours on solid media is sufficient to identify gram-negative rods, and further extending the incubation time does not improve SIR using MALDI-TOF [13]. Although the score value of correctly identified specimens did not change significantly with colony aging, the number of isolates showing no peak was significantly increased (Table 1). The effects of colony aging on BA and MH were not significant regardless of species, while results from Mac or IMP-Mac yielded lower confidence scores after 2 days of incubation compared to 1 day of incubation among E. cloacae and K. oxytoca. The bile salts present in Mac may interfere with the ionizing process because Hektoen Enteric agar or Salmonella–Shigella agar containing more bile salts showed lower confidence scores than MAC or xylose lysine deoxycholate agar [11]. The effect of aging on E. coli was lower on MAC than on K. pneumoniae when using the direct method [11]. This suggests that interference is species-dependent. Aging on Mac may cause more interference when using MALDI-TOF for identification than that when using the direct method. Considering that CRE surveillance
culture is not considered as an urgent condition in most clinical laboratories, colony reading is often delayed during weekends or holidays. The aging effect on selective agar may be a significant limitation in the application of MALDI-TOF to identify colonies on MAC or IMP-MAC.

A significant limitation of this study was the small sample size, which did not include all common species of CRE. However, *K. pneumoniae* is the most prevalent species among CRE in Korea and showed a significant difference by media type and colony aging. The results demonstrated that growth on IMP-Mac certainly significantly reduces the ability of MALDI-TOF to identify CRE. In conclusion, for reliable species identification of CRE grown on IMP-Mac, a disk diffusion set on MH should be applied when using the direct method with MALDI-TOF. Further evaluation of larger sample sizes and a greater number of species is required to generalize these conclusions.

REFERENCES

배지종류와 배양시간에 따른 카바페넴내성 장내세균의 MALDI Biotyper 동정능 평가
조영은 • 최병후 • 장정현 • 성홍섭 • 김미나
울산대학교 의과대학 서울아산병원 진단검사의학과

배경: T카바페넴내성 장내세균(carbapenem-resistant Enterobacteriaceae, CRE)의 감시배양배지에서 자란 집락을 직접 matrix assisted laser desorption ionization–time of flight mass spectrometry (MALDI–TOF)를 이용하여 동정할 때 배지종류와 배양시간에 따른 동정능을 평가하였다.

방법: 2017년 3월부터 2달간 의뢰된 CRE 감시배양에서 imipenem이 1 μg/mL 첨가된 MacConkey agar (IMP–Mac)에서 양성으로 선별된 집락들을 MALDI Biotyper (Bruker Daltonics, Germany)를 이용하여 직접 동정하였다. 또한 혈액한천배지(blood agar, BA), Mueller Hinton agar (MH), MacConkey agar (Mac)에 계대배양하여 각각의 배지에 하룻밤 배양한 것과 이틀째까지 배양한 것을 비교평가하였다. 속수준까지 동정에 성공한 경우 동정률과 동정한 스코어 값을 분석하였다.

결과: 총 117개 검체에서, Klebsiella pneumoniae 84개, Escherichia coli 12개, Enterobacter cloacae 9개, Klebsiella oxytoca 5개, Enterobacter aerogenes 4개, Raoultella ornithinolytica 2개가 동정되었다. Mac와 IMP–Mac의 동정률이 82.1% (P < 0.001)와 70.9% (P < 0.001)였던 반면, BA와 MH의 동정률은 각각 98.3%, 97.4% (P = 0.9)였었다. 배양 이틀째 BA, MH, Mac과 IMP–Mac 배지별 동정률은 96.6%, 96.6% (P = 1.0), 61.5% (P < 0.001), 58.1% (P < 0.001)로 Mac과 IMP–Mac에서 유의하게 감소하였다. MALDI 스코어 평균값은 Mac (2.017±0.22)와 IMP–Mac (1.978±0.24)이 BA (2.213±0.16) (P < 0.001)에 비해 유의하게 낮았다.

결론: Mac 또는 IMP–Mac에서 직접 MALDI Biotyper로 동정하는 경우 동정률이 낮았다. 이는 MALDI–TOF 동정 전 BA 또는 MH에서 하룻밤 계대배양함으로써 동정 정확도를 높일 수 있다는 것을 나타낸다.
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