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To cite this version:

Lucie Bardet, Stephanie Le Page, Thongpan Leangapichart, Jean-Marc Rolain. LBJMR medium: a new polyvalent culture medium for isolating and selecting vancomycin and colistin-resistant bacteria. BMC Microbiology, BioMed Central, 2017, 17 (1), pp.220. <10.1186/s12866-017-1128-x>. <hal-01729759>

HAL Id: hal-01729759

https://hal.archives-ouvertes.fr/hal-01729759

Submitted on 23 Apr 2018

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LBJMR medium: a new polyvalent culture medium for isolating and selecting vancomycin and colistin-resistant bacteria

Lucie Bardet†, Stéphanie Le Page†, Thongpan Leangapichart and Jean-Marc Rolain*

Abstract

Background: Multi-drug resistant bacteria are a phenomenon which is on the increase around the world, particularly with the emergence of colistin-resistant Enterobacteriaceae and vancomycin-resistant enterococci strains. The recent discovery of a plasmid-mediated colistin resistance with the description of the transferable mcr-1 gene raised concerns about the need for an efficient detection method for these pathogens, to isolate infected patients as early as possible. The LBJMR medium was developed to screen for all polymyxin-resistant Gram-negative bacteria, including mcr-1 positive isolates, and vancomycin-resistant Gram-positive bacteria.

Results: The LBJMR medium was developed by adding colistin sulfate salt at a low concentration (4 μg/mL) and vancomycin (50 μg/mL), with glucose (7.5 g/L) as a fermentative substrate, to a Purple Agar Base (31 g/L). A total of 143 bacterial strains were used to evaluate this universal culture medium, and the sensitivity and specificity of detection were 100% for the growth of resistant strains. 68 stool samples were cultured on LBJMR, and both colistin-resistant Gram-negative and vancomycin-resistant Gram-positive strains were specifically detected.

Conclusions: The LBJMR medium is a multipurpose selective medium which makes it possible to identify bacteria of interest from clinical samples and to isolate contaminated patients in hospital settings. This is a simple medium that could be easily used for screening in clinical microbiology laboratories.

Keywords: Colistin resistance, Culture media, Mcr-1, mcr genes, Vancomycin-resistant enterococci, Enterobacteriaceae, Multi-drug resistant, Detection method, Screening method

Background

The worldwide emergence of multidrug-resistant (MDR) bacteria represents a major public health issue. Controlling the spread of these bacteria relies upon both reducing the prescription of antibiotics and preventing transmission from carrier patients to others [1]. More specifically, this prevention targets the emerging carbapenemase-producing Enterobacteriaceae and vancomycin-resistant enterococci (VRE) strains, with the development of specific detection methods, mostly based on chromogenic and selective culture media [2].

The increase in infections due to carbapenemase-producing Enterobacteriaceae has led to the revival of colistin as a last-resort treatment [3]. Its widespread use, particularly in the livestock food in many countries [4], has inevitably led to the emergence of colistin-resistant strains over the past ten years [5, 6]. Until recently, all colistin resistance mechanisms which had been described were attributed to chromosomal mutations [7]. In China in 2015, Liu and colleagues were the first to report the plasmid-mediated colistin-resistance gene in animals and humans, which they named mcr-1 [8]. This was followed by descriptions of the variants mcr-1.2 and mcr-1.3 [9, 10], mcr-2, mcr-3, mcr-4 and mcr-5 genes [11–14]. A significant number of human cases (in the majority of cases asymptomatic carriers) was described [15, 16]. Particularly, the worldwide analysis of colistin-resistant strains with an unknown mechanism allowed the detection of a consequent number of mcr-1-positive bacteria [17–19]. This mobile colistin resistance gene presents low resistance levels, with minimal inhibitory concentrations (MIC)
of colistin around 4 µg/ml, which is close to the clinical breakpoint of colistin resistance (> 2 µg/mL), according to the European Committee on Antimicrobial Susceptibility Testing (EUCAST) [20]. Those data raised concerns about the detection and isolation of these pathogens, and the screening of the mcr-1 gene into carbapenemase-producing bacteria has been added to recommendations for clinical microbiology laboratories in France [21]. While carbapenemase screening is well-defined with phenotypic and molecular techniques [22], there is a need for colistin-resistance screening tools [2]. Given the diversity of mechanisms of colistin resistance [23], phenotypic methods, such as chromogenic culture media [2, 24] are preferred for a rapid detection.

In addition to colistin-resistant Enterobacteriaceae, the detection of vancomycin-resistant enterococci isolates is of clinical concern. The prevalence of VRE strains is increasing in Europe, especially Enterococcus faecium [25] and has led to nosocomial infections in the United States [26], and their dissemination is associated with high mortality rates [27]. The development of an effective screening tool for the early detection of those multi-drug resistant pathogens has become a priority, in order to adapt treatment and isolate patients who are either infected or carriers.

For such purposes, we developed a polyvalent selective culture medium which can isolate both colistin-resistant Gram-negative bacteria, including Enterobacteriaceae strains harboring the mcr-1 gene, and vancomycin-resistant Gram-positive bacteria. In this study, we evaluated the performances of this medium on a representative set of bacterial strains and fecal samples. This universal culture medium was named LBJMR, standing for Lucie Bardet-Jean-Marc Rolain, and has recently been patented [28].

**Methods**

Bacterial strains and samples cultured on LBJMR

A total of 143 bacterial strains were used in this study, of which 101 were Enterobacteriaceae: 75 with an acquired mechanism of resistance to colistin (including 32 strains harboring the mcr-1 gene, and vancomycin-resistant Gram-positive bacteria. In this study, we evaluated the performances of this medium on a representative set of bacterial strains and fecal samples. This universal culture medium was named LBJMR, standing for Lucie Bardet-Jean-Marc Rolain, and has recently been patented [28].

The colistin-resistance genes which were potentially involved were previously screened by RT-PCR for Enterobacteriaceae [6, 44], as were the vancomycin resistance genes vanA and vanB for the VRE strains [45]. MICs of colistin and vancomycin were determined using E-test® strips (BioMérieux, Marcy l’Étoile, France) for Gram-negative and Gram-positive strains, respectively, and MICs of daptomycin were determined for the four reference strains of Enterococcus faecium (DSM 17050, 25,698, 25,697 and 13,590), and were evaluated according to the EUCAST guidelines [20].

In addition, 68 samples were cultivated on LBJMR medium, including 66 stool samples (56 from humans and ten from chickens) that were previously screened for mcr-1 by RT-PCR (56 positive and ten negative) and two clinical rectal swabs obtained from patients in the La Timone hospital, from which a vancomycin-resistant E. faecium VRE strain was isolated. Colonies with different morphologies isolated by culture of samples on LBJMR were isolated by subculture on Trypticase Soy Agar (TSA, Becton-Dickinson, Heidelberg, Germany), and the species were identified using Matrix-Assisted Laser Desorption-Ionization-Time Of Flight Mass Spectrometry (MALDI-TOF MS), as described previously [46].

**LBJMR development**

In order to choose the most suitable medium for selecting colistin-resistant Enterobacteriaceae, including strains harboring the mcr-1 gene, five commercial differential media usually used for the detection of Enterobacteriaceae were supplemented with a range of concentration of colistin sulfate salt (MP Biomedicals, Illkirch, France) from 0 to 32 µg/mL: Drigalski Lactose Agar (Biokar diagnostics, France), BBL™ Eosin Methylene Blue Agar (EMB; Becton-Dickinson, Heidelberg, Germany), Difco™ Violet Red Bile Agar (Becton-Dickinson, Heidelberg, Germany), BBL™ MacConkey Agar II (Becton-Dickinson, Heidelberg, Germany), and Difco™ Purple Agar Base (Becton-Dickinson, Heidelberg, Germany) supplemented with 7.5 g/L of glucose (MP Biomedicals, Illkirch, France). Four Klebsiella pneumoniae, including two colistin-resistant, and four Escherichia coli, including two which were colistin-resistant with the mcr-1 gene, were inoculated at 10^6 CFU on each prepared agar [19].

The Purple Agar Base supplemented with glucose (7.5 g/L), colistin sulfate (0, 4 and 8 µg/mL) and vancomycin (0 and 50 µg/mL) (Sandoz, Levallois-Perret, France) was evaluated with 24 representative strains, including 20 Enterobacteriaceae (five intrinsic genera, eight with acquired resistance and seven colistin-susceptible strains as control) and four Gram-positive strains susceptible to vancomycin, which were inoculated at 10^6 CFU. This experiment was also conducted by replacing glucose with lactose (Laboratoires Humeau, La Chapelle-sur-
Erdre, France). The optimum concentration of vancomycin for the medium composition was then determined using 20 Gram-positive strains representative of vancomycin resistance (three intrinsically-resistant genera and 17 enterococci, including seven VRE) and ten chicken stool samples from Algeria, five of which were positive for mcr-1 by RT-PCR, and five of which were negative [19, 44].

LBJMR evaluation
The sensitivity and specificity of the LBJMR medium to detect all colistin-resistant Gram-negative bacteria and all vancomycin-resistant Gram-positive bacteria were evaluated with all the strains listed in Additional file 1: Table S1 and samples listed in Additional file 1: Table S2, which were incubated on LBJMR in an aerobic atmosphere at 37 °C for between 24 and 48 h.

Inocula of the strains were prepared at 0.5 McFarland (corresponding to 1.5 × 10^9 CFU/mL according to the EUCAST expert system) and serial 10-fold dilutions were then performed in Phosphate Buffer Saline. 10 μL of these suspensions were deposited on agars and simultaneously on TSA medium to determine the detection limits for each bacterium by assessing the viable concentration by colony count. Growth or inhibition of these microorganisms was observed after overnight incubation at 37 °C in an aerobic atmosphere. Those experiments were performed in duplicate.

Finally, as the LBJMR medium was developed so as to avoid Proteus swarming, Proteus mirabilis and Proteus vulgaris strains were inoculated alone or mixed with E. coli P10 (mcr-1) at the same concentration to evaluate the isolation ability of the LBJMR medium. Cultures were incubated at 37 °C for 72 h.

Mcr-1 screening on stool samples
A total of 1052 stool samples were used in this study, 899 from humans from different regions of the world: Marseille (n = 212) and Angers (n = 128) in France, Thailand (n = 212), Laos (n = 189), Nigeria (n = 143), and Senegal (n = 15). 153 samples were from animals, including chickens from Algeria (n = 10), rodents from Cambodia (n = 21), and rodents (n = 60), pigs (n = 16) and goats (n = 46) from Laos.

The presence of the mcr-1 gene in those stools was first screened using RT-PCR as previously described [44]. Briefly, the DNA of the stools was extracted, after pretreatment by proteinase K and incubation at 95 °C, using Biorobot EZ1 Advanced XL (Qiagen, Hilden, Germany). Positive DNA was confirmed using standard PCR. Samples with positive DNA were then cultured on LBJMR, after a prior enrichment step performed by inoculating approximately 1 g of the sample in tryptic soy broth (TSB) medium (BioMérieux, Marcy l’Étoile, France) and incubating for 24 h at 37 °C. 100 μL of these liquid media were then plated on agars that were also incubated at 37 °C for 24 h.

For each Gram-negative isolate identified by MALDI-TOF MS, colistin MIC was determined with a colistin E-test® strip. For each colistin-resistant isolate, the presence of mcr-1 was investigated by RT-PCR as previously described [44]. Susceptibility testing to 26 antibiotics was performed on mcr-1-positive strains using the disk diffusion method, and interpreted according to the EUCAST guidelines [20].

Comparison of the LBJMR medium with other polymyxin-containing media
The sensitivity and specificity of the LBJMR medium was first compared to the BD™ Cepacia Medium (Cepacia, Becton-Dickinson, Heidelberg, Germany) medium with all the colistin-resistant Enterobacteriaceae strains listed in Additional file 1: Table S1. The LBJMR medium was then compared to the SuperPolymyxin medium with 103 bacterial strains, including 61 of the Enterobacteriaceae, 17 non-fermentative Gram-negative, and 25 Gram-positive strains (Additional file 1: Table S1). Other polymyxin-containing media were also compared concomitantly to the LBJMR medium, using EMB Agar (Sigma-Aldrich, Illkirch, Germany) as a control, with some of these strains as described below. Inocula were prepared following the same protocol as for the LBJMR evaluation. Columbia Colistin Nalidixic Acid Agar +5% sheep blood (CNA) (BioMérieux, Marcy l’Étoile, France) and three mixed culture media: EMB with colistin sulfate (4 μg/mL) and vancomycin (50 μg/mL), LBJMR with Daptomycin (10 μg/mL, Novartis, Horsham, United Kingdom) in place of vancomycin and LBJMR with Ampotericin B (5 μg/mL, Bristol-Myers Squibb, Rueil-Malmaison, France) were concomitantly tested with 45 Enterobacteriaceae strains representative of colistin resistance (30 mcr-1, ten susceptible and five intrinsically resistant genera). LBJMR with Ampotericin B was also evaluated with the eight VRE strains. Finally, the LBJMR medium was compared to the SuperPolymyxin medium with 12 samples, including the two clinical rectal swabs from the La Timone hospital and ten human stools from Thailand, five of which had been identified as positive by culture for the detection of an isolate harboring the mcr-1 gene, and five controls that were negative for mcr-1 using RT-PCR.

Statistical analysis
The data were analyzed using a Chi-test to compare the Cepacia and LBJMR media and a Student-t test for a pairwise comparison of SuperPolymyxin and LBJMR selective media for the detection of non-fermentative Gram-negative colistin-resistant strains. Significance was assessed at p < 0.05.
Results

LBJMR development

Because colistin is a cationic molecule, we looked for a medium which was deprived of any ions in its original composition and which was also deprived of any electrolytes to avoid Proteus spp. swarming. The optimal conditions for LBJMR medium (growth of colistin- and vancomycin-resistant strains and inhibition of colistin- or vancomycin susceptible strains) were obtained on Purple Agar Base with glucose (Additional file 1: Table S3), 4 μg/mL of colistin sulfate and 50 μg/mL of vancomycin (Additional file 1: Table S4). The culture of ten chicken stools from Algeria enabled the detection of E. coli strains harboring the mcr-1 gene from three of the five samples which were positive for mcr-1 by RT-PCR, including strain 235 which had previously been isolated on Cepacia medium [19] and no Gram-negative strains were detected from the five negative samples (Additional file 1: Table S2). A correlation was observed between the Ct-values that reflect the DNA concentration and the results of culture (Additional file 1: Table S2).

The final preparation of the LBJMR medium was as follows: 31 g/L of Purple Agar Base, 7.5 g/L of glucose, 4 μg/mL of colistin sulfate and 50 μg/mL of vancomycin. 7.5 g/L of lactose was used in place of glucose for the 24 strains initially tested and the same results were obtained. Glucose was selected because, combined with bromocresol purple, a pH indicator, it revealed the polyvalent capacity of the LBJMR medium with an easy and fast visualization of the species of interest in the clinical routine: both Enterobacteriaceae and enterococci gave yellow colonies, contrasting on the purple agar and exhibiting different sizes (2–3 mm and 0.1–1 mm, respectively).

LBJMR evaluation

All the polymyxin-resistant Enterobacteriaceae strains (n = 81) tested were able to grow on the LBJMR medium, with the lowest detection limit (10⁴ CFU), including those with the mcr-1 gene, as well as all the colistin-resistant non-fermentative Gram-negative (n = 10) and vancomycin-resistant Gram-positive strains (n = 11), with detection limits dependent on their MIC, as shown in Table 1. Meanwhile, all the bacteria susceptible to colistin or vancomycin were inhibited (not detected at 10⁵ CFU), giving rise to 100% sensitivity and specificity for the growth of Gram-negative colistin-resistant and Gram-positive vancomycin-resistant strains (Table 1).

The swarming of Proteus sp. strains was fully inhibited on LBJMR, even after 48 h of incubation.

Of the 1052 stools screened for mcr-1 by RT-PCR, 66 were cultured on LBJMR, including 56 that were positive after 32 cycles of RT-PCR, as well as ten negative stool samples as controls. 16 cultures were positive on LBJMR medium, enabling the detection of 17 colistin-resistant Enterobacteriaceae strains harboring the mcr-1 gene (colistin MICs ranging from 2 to 8 μg/mL): 15 were identified as E. coli and two as K. pneumoniae by MALDI-TOF MS (Additional file 1: Table S2). Indeed, two bacterial species, E. coli and K. pneumoniae, were detected from the culture of the same sample (FHM128) (Additional file 1: Table S2). Five isolates were already known from a previous study on Cepacia medium [6, 19]. All the E. coli isolates from France were Extended-Spectrum Beta-Lactamase-producing strains (data not shown). In conclusion, the LBJMR medium enabled the isolation of new colistin-resistant strains harboring the mcr-1 gene. The weak culture results compared to PCR could be explained by the length of storage of the tested samples (up to four years) at −80 °C, possibly inducing the death of bacteria while their DNA was still detectable [47]. However, it should be noted that, for some samples, we isolated another kind of strain, mostly Gram-positive bacteria which have an intrinsic resistance to vancomycin, largely Pediococcus pentosaceus and Weissella cibaria, as shown in Additional file 1: Table S2.

Finally, the culture of two clinical samples from the hospital enabled the isolation of two vancomycin-resistant E. faecium isolates (VRE1 and VRE2). In addition, two colistin-resistant Enterobacteriaceae isolates were also detected in the VRE2 sample, identified as K. pneumoniae LB3 (MIC = 64 μg/mL) and E. coli LB4 (MIC = 16 μg/mL), both negative for the mcr-1 gene (Fig. 1).

Comparison between LBJMR and other selective polymyxin-containing media

The LBJMR medium was more sensitive than the Cepacia and CNA media for detecting the strains exhibiting the mcr-1 gene (Additional file 1: Table S5). Indeed, only four of those strains could grow on the CNA medium, because it also contains nalidixic acid and fewer than half of them grew on the Cepacia medium. Comparison of the sensitivities of the LBJMR (100%) and Cepacia (47%) media using a Chi-square statistical test gave a significant p-value (<10⁻⁴).

LBJMR and SuperPolymyxin media presented the same sensitivity for Enterobacteriaceae, with the growth of all colistin-resistant strains (detection limit of 10⁴ CFU), including the 30 mcr-1 strains tested, and the inhibition of all susceptible strains (Table 1). In addition, the culture of ten human stools gave the same results: five were positive, with the detection of an E. coli positive for mcr-1, and the five controls remained negative (Additional file 1: Table S2). In contrast, the LBJMR medium showed a significantly higher sensitivity than the SuperPolymyxin medium for the detection of colistin-resistant non-fermentative Gram-negative strains, with a significant p-value of 0.019 (<0.01) with a pairwise comparison using a Student-t test (Table 1). More
| Bacterial strains             | Detection limits on culture media (CFU): |
|------------------------------|------------------------------------------|
|                              | CT | MIC | LBJMR | SuperPolymyxin |
| **Colistin-resistant Enterobacteriaceae** |    |     |       |                |
| *E. coli* SE65               | 4  | 10^1 | 10^1  |                |
| *E. coli* 117R               | 4  | 10^1 | 10^1  |                |
| *E. coli* 1R                 | 4  | 10^1 | 10^1  |                |
| *E. coli* 1R 2104            | 4  | 10^1 | 10^1  |                |
| *E. coli* 44A                | 4  | 10^1 | 10^1  |                |
| *E. coli* 6R                 | 4  | 10^1 | 10^1  |                |
| *E. coli* 85R                | 4  | 10^1 | 10^1  |                |
| *E. coli* 95R                | 4  | 10^1 | 10^1  |                |
| *E. coli* 96R                | 4  | 10^1 | 10^1  |                |
| *E. coli* 134R               | 3  | 10^1 | 10^1  |                |
| *E. coli* 143R               | 3  | 10^1 | 10^1  |                |
| *E. coli* LH121              | 16 | 10^1 | 10^1  |                |
| *E. coli* LH140^1            | 12 | 10^1 | 10^1  |                |
| *E. coli* LH257              | 12 | 10^1 | 10^1  |                |
| *E. coli* LH57^1             | 8  | 10^1 | 10^1  |                |
| *E. coli* LH1                | 6  | 10^1 | 10^1  |                |
| *E. coli* LH30               | 6  | 10^1 | 10^1  |                |
| *E. coli* TH214              | 6  | 10^1 | 10^1  |                |
| *E. coli* TH99               | 4  | 10^1 | 10^1  |                |
| *E. coli* 235                | 4  | 10^1 | 10^1  |                |
| *E. coli* P6                 | 6  | 10^1 | 10^1  |                |
| *E. coli* P10                | 4  | 10^1 | 10^1  |                |
| *E. coli* P17                | 4  | 10^1 | 10^1  |                |
| *E. coli* FHA102^2           | 12 | 10^1 | 10^1  |                |
| *E. coli* FHA113^4           | 12 | 10^1 | 10^1  |                |
| *E. coli* NH94^5             | 12 | 10^1 | 10^1  |                |
| *E. coli* TH176              | 6  | 10^1 | 10^1  |                |
| *K. pneumoniae* FHA60        | 8  | 10^1 | 10^1  |                |
| *K. pneumoniae* FHA128       | 4  | 10^1 | 10^1  |                |
| *K. pneumoniae* 119R         | 3  | 10^1 | 10^1  |                |
| *K. pneumoniae* LH131^4       | 32 | 10^1 | 10^1  |                |
| *K. pneumoniae* LH61^7        | 24 | 10^1 | 10^1  |                |
| *K. pneumoniae* LH17         | 12 | 10^1 | 10^1  |                |
| *K. pneumoniae* LH92         | 12 | 10^1 | 10^1  |                |
| *K. pneumoniae* LB1^6         | 32 | 10^1 | 10^1  |                |
| *P. mirabilis* FH112          | >256 | 10^1 | 10^1  |                |
| *P. vulgaris* PV148          | >256 | 10^1 | 10^1  |                |
| *P. alcalifaciens* TH44      | >256 | 10^1 | 10^1  |                |
| *M. morganii* FM102          | >256 | 10^1 | 10^1  |                |
| *S. marcescens* E13         | 128 | 10^1 | 10^1  |                |
| *Others* colistin-resistant  |    |     |       |                |
| *B. cepacia* FHM-BC1         | 64 | 10^5 | 10^5  |                |
| *B. cepacia* FHM-BC2         | >256 | 10^1 | 10^1  |                |
| Bacterial strains | Detection limits on culture media (CFU): |
|-------------------|-----------------------------------------------|
| **A. xylosoxidans FHM-AX** | 3 | $10^1$ | $>10^5$ |
| **S. maltophilia FHM-SM** | 12 | $10^4$ | $>10^5$ |
| **I. limosus FHM-IL** | $>256$ | $10^4$ | $10^4$ |
| **P. pulmoncola FHM-PP** | $>256$ | $10^1$ | $10^3$ |
| **S. putrefaciens FHM-SP** | NC | $10^4$ | $>10^5$ |
| **O. anthropi FHM-OA** | NC | $10^2$ | $>10^5$ |
| **P. aeruginosa FHM-PACOLR1** | $>256$ | $10^1$ | $10^1$ |
| **A. baumannii ABIsac_ColiR** | 8 | $10^5$ | $10^1$ |

**Colistin-susceptible**

| **E. asburiae P113** | 0.19 | $>10^5$ | $10^4$ |
| **E. cloacae NH151** | 0.50 | $>10^5$ | $10^4$ |
| **E. cloacae NH74** | 0.38 | $>10^5$ | $10^4$ |
| **K. pneumoniae CIP 82.91** | 0.125 | $>10^5$ | $10^5$ |
| **K. pneumoniae LB2** | 0.128 | $>10^5$ | $10^5$ |
| **K. pneumoniae TH205** | 0.125 | $>10^5$ | $10^5$ |
| **K. pneumoniae TH285** | 0.125 | $>10^5$ | $10^5$ |
| **Proteus vulgaris P100** | 0.94 | $>10^5$ | $10^5$ |
| **S. enterica 108R** | 1 | $>10^5$ | $10^5$ |
| **S. enterica 122R** | 0.5 | $>10^5$ | $10^5$ |
| **E. coli ATCC 25922 CIP 76.24** | 0.094 | $>10^5$ | $10^5$ |
| **E. coli 161** | 0.094 | $>10^5$ | $10^5$ |
| **E. coli 169** | 0.094 | $>10^5$ | $10^5$ |
| **E. coli FHM88S** | 0.125 | $>10^5$ | $10^5$ |
| **E. coli TH134S** | 0.094 | $>10^5$ | $10^5$ |
| **E. coli LH53S** | 0.094 | $>10^5$ | $10^5$ |
| **E. coli LH165S** | 0.074 | $>10^5$ | $10^5$ |
| **E. coli TH775S** | 0.064 | $>10^5$ | $10^5$ |
| **E. coli 2825** | 0.074 | $>10^5$ | $10^5$ |
| **E. coli FHM19S** | 0.125 | $>10^5$ | $10^5$ |
| **P. aeruginosa FHM-PA4** | 2 | $>10^5$ | $10^5$ |
| **P. aeruginosa FHM-PA5** | 0.50 | $>10^5$ | $10^5$ |
| **P. aeruginosa FHM-PA6** | 0.38 | $>10^5$ | $10^5$ |
| **S. xianemensis 111P** | 0.125 | $>10^5$ | $10^5$ |
| **S. xianemensis 111A** | 0.094 | $>10^5$ | $10^5$ |
| **A. nosocomialis ABG135** | 0.064 | $>10^5$ | $10^5$ |
| **A. pitti MK** | 0.125 | $>10^5$ | $10^5$ |

**Gram-positives**

| **E. faecium DSM17050** | $>256$ | $10^1$ | $10^5$ |
| **E. faecium DSM13590** | $>256$ | $10^1$ | $10^5$ |
| **E. faecium DSM25698** | $>256$ | $10^1$ | $10^5$ |
| **E. faecium DSM25699** | $>256$ | $10^1$ | $10^5$ |
| **E. faecium FHMVRE1** | $>256$ | $10^1$ | $10^5$ |
| **E. faecium FHMVRE2** | $>256$ | $10^1$ | $10^5$ |
| **E. faecium VRE** | 24 | $10^3$ | $10^5$ |
| **E. faecalis JH2–2; Tn 1549** | 64 | $10^2$ | $10^5$ |
specifically, the LBJMR medium was able to detect those screened in the cystic fibrosis samples, such as Burkholderia cepacia, even at low concentrations, and could replace specific media such as the Cepacia medium for their isolation. The addition of amphotericin B or the replacement of vancomycin by daptomycin in the LBJMR medium did not affect the growth of Enterobacteriaceae strains (Additional file 1: Table S5). Finally, all the VRE strains were inhibited on the SuperPolymyxin medium because of the presence of daptomycin, while they were detected on the LBJMR medium (Table 1).

These results were confirmed by the culture of the two clinical samples from the hospital: vancomycin-resistant E. faecium could only grow on the LBJMR medium and not on the SuperPolymyxin medium for both samples, and the two Enterobacteriaceae from the VRE2 sample found on the LBJMR medium were also detected on the Super-Polymyxin medium.

Finally, the concomitant addition of colistin and vancomycin to EMB agar showed the systematic inhibition of E. coli strains harboring the mcr-1 gene, corresponding to the tested strains with the lowest MIC.

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**Table 1** Detection limits of targeted bacteria on LBJMR and SuperPolymyxin culture media (Continued)

| Bacterial strains | Detection limits on culture media (CFU): |
|-------------------|--------------------------------------|
| W. cibaria P18A   | >256 10^7 >10^5 |
| W. cibaria P18B   | >256 10^7 >10^5 |
| L. lactis P18C    | >256 10^7 >10^5 |
| S. aureus CF_Marseille | 1 >10^5 >10^5 |
| E. faecium TH26   | 0,75 >10^5 >10^5 |
| E. faecium 349    | 0,75 >10^5 >10^5 |
| E. faecium TH12   | 2 >10^5 >10^5 |
| E. faecium LH165  | 0,75 >10^5 >10^5 |
| E. faecium AI7    | 0,75 >10^5 >10^5 |
| E. faecium TH43   | 0,75 >10^5 >10^5 |
| E. faecium TH95   | 0,75 >10^5 >10^5 |
| E. faecium 282    | 0,75 >10^5 >10^5 |
| E. faecalis JH2-2-C2 | 1,5 >10^5 >10^5 |
| E. faecalis JH2-2S | 1,5 >10^5 >10^5 |
| E. gallinarum SE3 | 1,5 >10^5 >10^5 |
| E. casseliflavus SE3 | 1,5 >10^5 >10^5 |
| E. hirae LH111    | 1 >10^5 >10^5 |

MIC of colistin (CT) and vancomycin (VCN) are given in μg/mL. Enterobacteriaceae with mcr-1 gene are in bold font. COLISTIN-resistant strains with a mutation on another gene are indicated as follows: 1 phoP E375K, 2 pmrB A159V, 3 pmrB P7-Q12(del 6aa), 4 pmrB T156 K, 5 pmrB I92 insertion, 6 mgrB Stop, 7 mgrB Sub, 8 pmrB T157P. Enterococci strains with vancomycin-resistant genes are indicated as follows: a vanA and b vanB.

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**Fig. 1** Aspect of the different types of colonies on LBJMR after culture of a clinical sample. Enterobacteriaceae exhibited yellow colonies that are larger (2–3 mm) than enterococci (0.1–1 mm).
Discussion
In accordance with recommendations aiming to isolate patients carrying multi-drug resistant bacteria, it is becoming essential to detect colistin-resistant Gram-negative pathogens, particularly with the recent description of plasmid-mediated colistin-resistant genes among Enterobacteriaceae strains [8–12]. We do not currently know the potential consequences of patient colonization by those resistant bacteria, as it has already been demonstrated that acquisition of resistance is associated with a decrease in virulence [48–50], but it is necessary to implement efficient tools to prevent and monitor as closely as possible any epidemic outbreak and develop a rapid therapeutic strategy.

Currently, the only available method for detecting colistin resistance, according to joint EUCAST and Clinical Laboratory Standard Institute (CLSI) expert systems, is antibiotic susceptibility testing using broth microdilution, which is not suitable for daily screening in clinical microbiology laboratories. Current polymyxin-containing culture media are also not appropriate because of their high concentration of polymyxin. Indeed, they were not developed to isolate Enterobacteriaceae but to specifically detect bacteria that are intrinsically resistant to polymyxins, avoiding contaminants such as Enterobacteriaceae species. Thus, the establishment of an effective protocol to detect colistin resistance with the development of a reliable culture medium was necessary in clinical microbiology laboratory.

The significant number of strains tested on our selective medium showed that the LBJMR medium allows all colistin-resistant Enterobacteriaceae strains to grow, even those with a very low MIC for colistin and harboring the mcr-1 gene, when all the susceptible strains were inhibited. Furthermore, the screening conducted on stool samples that were positive for mcr-1 by PCR allowed the isolation of new mcr-1 strains (Additional file 1: Table S2). The LBJMR medium could also detect colistin-resistant non-fermentative Gram-negative bacteria, including pathogens which are often found in patients suffering from cystic fibrosis (Table 1). We thereby created a universal culture medium, able to replace specific polymyxin-containing culture media usually used in routine diagnosis. All the vancomycin-resistant bacteria tested were detected at low concentrations on the LBJMR medium, including the VRE strains, when susceptible were inhibited. To attest this capacity, the LBJMR medium should be tested on more VRE strains and comparatively to VRE selective culture media.

The performances of the LBJMR medium are summarized in Table 2: samples can be cultured directly on the LBJMR selective medium, without a previous decontamination step, and colonies can be directly analyzed from the primary culture, without subcultures. Bacteria of interest are easily recognizable, as shown in Fig. 1: both colistin-resistant Enterobacteriaceae and vancomycin-resistant enterococci exhibit yellow colonies (2–3 mm and 0.1–1 mm respectively), on a purple agar base. Because LBJMR also permits the growth of intrinsic resistant bacteria, screening should be completed with a rapid identification using MALDI-TOF MS [51], directly from the LBJMR medium. Antibiotic susceptibility testing and PCR screening can be performed in the same time. Nordmann et al. developed a different selective medium called SuperPolymyxin, based on EMB agar and containing 3.5 μg/mL of colistin sulfate, 10 μg/mL of daptomycin and 5 μg/mL of amphotericin B [52], associated with a complementary phenotypic test, the rapid polymyxin NP [53], to screen for colistin-resistant Gram-negative bacteria [54]. These two media are differential and are able to recognize bacterial species based on color. Sensitivities for colistin-resistant Enterobacteriaceae were the same for the SuperPolymyxin and LBJMR media, but only the LBJMR medium could detect VRE strains (Table 1). Indeed, the SuperPolymyxin medium is composed of daptomycin and amphotericin B, which prevents the growth of contaminants including bacteria of interest such as VRE strains, because they are mostly susceptible to daptomycin. In our study, we tested stool samples and obtained some contaminants, including Gram-positive bacteria which were intrinsically resistant to vancomycin (Additional file 1: Table S2) or some yeasts.

More recently, the culture medium ChromAgar COL-APSE was developed and compared to the SuperPolymyxin medium [55]. The comparison of the LBJMR and the CHROMAgar COL-APSE performances have to be assessed. All those media should be also evaluated on

| Table 2 Performance of LBJMR medium |
|-------------------------------------|
| Criteria                         | LBJMR medium |
|-----------------------------------|--------------|
| Isolates screened                 | Colistin-resistant Gram-negatives: |
| - Enterobacteriaceae, including those harboring the mcr-1 gene |
| - Non-fermentative Gram-negative colistin-resistant strains, including those involves in cystic fibrosis samples |
| Vancomycin-resistant Gram positives, including Enterococci |
| Aspect of colonies                | Yellow on purple agar: 2–3 mm for Enterobacteriaceae, 0.1–1 mm for Enterococci |
| Incubation                        | Aerobic atmosphere, 37 °C, 24H (sterile at 48H) |
| Culture of samples                | Direct on LBJMR, no previous decontamination |
| Isolates analysis                 | Colonies can be picked directly from primary cultures on LBJMR for analysis: |
| - MALDI-TOF identification         | |
| - Antibiotic Susceptibility testing| |
| - PCR screening for resistance genes. | |
| Avoid contamination               | - Inhibition of Proteus swarming |
| - Inhibition of yeast possible by adding amphotericin B |

Further reading

[8] Bardet et al. BMC Microbiology (2017) 17:220
heteroresistant strains, as they are difficult to detect and their frequency is widely underestimated [56].

The advantage of the LBJMR medium in terms of isolating vancomycin resistant enterococci can be also a disadvantage in a few cases, because Gram-positive bacteria which are intrinsically resistant to vancomycin bacteria, including *Pediococcus*, can also be isolated. In the LBJMR medium, Gram-positive bacteria appear with small colonies and a simple identification by MALDI-TOF can identify them. For the elimination of yeast in samples, amphotericin B can be added in our medium.

**Conclusion**
The LBJMR medium is an adequate screening tool for all colistin-resistant isolates from clinical samples, independently of their resistance level or mechanism. LBJMR could be used in routine laboratory work to detect colistin-resistant and vancomycin-resistant bacteria, allowing for the direct analysis of colonies and, thus, the early isolation of contaminated patients in hospital settings. This medium is currently being investigated as a routine medium in our institute to determine its usefulness in detecting such bacteria.

**Additional file**

**Additional file 1: Table S1.** List of the studied strains. The MIC of colistin (CT) or vancomycin (VCN) are indicated in μg/mL. Table S2. List of stools used and Gram-negative isolates detected on LBJMR. Positive isolates for *mcr-1* and their corresponding samples are in bold font. New isolates harboring *mcr-1* are underlined. Their MICs of colistin are indicated in μg/mL. – (−) Negative (ND) refers to Not Done. Table S3. Comparison of different agar bases supplemented with different concentrations of colistin. Genes involved in colistin resistance are indicated in bold and colistin MICs are indicated in μg/mL. Table S4. Development of the selective medium for the detection of colistin-resistant strains. Table S5. Comparison of different polymyxin-containing media. Numbers indicate the total number of strains that grew on each culture media. (DOCX 99 kb)

**Abbreviations**

CLSI: Clinical Laboratory Standard Institute; CNA: Colistin Nalidixic Acid; EMB: Eosin-Methylene Blue; EUCAST: European Committee on Antimicrobial Susceptibility Testing; MALDI-TOF MS: Matrix-Assisted Laser Desorption-Ionization-Time Of Flight Mass Spectrometry; MIC: Minimum Inhibitory Concentration; TSA: Trypticase Soy Agar; VRE: Vancomycin-Resistant Enterococci

**Acknowledgements**

We thank to Marielle Bedotto-Buffet for technical assistance and TradOnline for English corrections.

**Funding**

IHU Méditerranée Infection.

**Availability of data and materials**

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**Authors’ contributions**

LB designed the study, performed the experiments and analyzed and interpreted the data. SLP was analyzed, interpreted the data and involved in revising the manuscript. TL contributed to the development and analysis of data. JMR was responsible for the conception and design of the study, and the analysis of data. All authors read and approved the final manuscript.

**Ethics approval and consent to participate**

Not applicable

**Consent for publication**

Not applicable

**Competing interests**

The authors declare that they have no competing interests

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Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Received: 25 August 2017 Accepted: 15 November 2017

Published online: 23 November 2017
39. Rolain J-M, Diene SM, Kempf M, Gimenez G, Robert C, Raoult D. Real-time most recent history of travel. J Antimicrob Chemother. 2017;62:2942–4.
40. Vila-Farré S, Ferrer-Navarro M, Callaria AE, et al. Loss of LPS is involved in the virulence and resistance to colistin of colistin-resistant Acinetobacter nosocomialis mutants selected in vitro. J Antimicrob Chemother. 2015;70: 2981–6.