Chromogenic Agar Medium : A Versatile Tool for the Diagnosis of UTI

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Abstract

Objective: The present study was done on Chromogenic agar media to identify uropathogens more efficiently by its characteristic colony colour for each of the organism.

Methodology: A total 300 sample were collected from Rajshahi Medical College Hospital, Bangladesh. Urine samples of the suspected UTI cases, showing pus cells >5/HPF on microscopic examination were included for urine culture simultaneously onto Chromogenic agar media, Blood agar and MacConkey agar media.

Results: Culture yielded 139 (46.33%) bacterial growth among them, 133 (44.33%) showed single organism and remaining 06 (2.00%) showed mixed growth of two organisms in different combinations. It is evident from the present study that both Chromogenic agar media and Blood agar (BA) media supported growth of all 145 bacteria, while MacConkey (MAC) agar yielded 133(91.72%) bacterial growths. The rate of presumptive identification of the isolates was found significantly higher (97.24%) on Chromogenic agar media when compared with the MacConkey agar (80.68%) and Blood agar (27.58%) media. Out of 91 E. coli isolated, 88(96.70%) could be identified differentially on Chromogenic agar media in contrast to 85(93.40%) on MacConkey agar and only 06(06.59%) on Blood agar. Again, all 06 (100%) of the isolate-pairs of mixed growth were identified distinctly on Chromogenic agar media, whereas both Blood agar and MacConkey agar media could revealed only 01(16.66%) of the polymicrobial growth.

Conclusion: Chromogenic agar media has been documented for its very high yielding rate, rapid presumptive identification of both single and polymicrobial growths with greater precision and avoidance of biochemical tests for further identification of uropathogens. Thus it can be recommended as primary urine culture medium to be used by the clinical microbiology laboratories.

Key Words: Diagnosis of UTI, Chromogenic agar media.

Introduction

UTI refer to invasion of urinary tract including urinary bladder, prostate, collecting system or kidney by different microorganisms¹. Isolation and identification of the causative organism from urine through culture is the gold standard for the diagnosis of UTI. Predominant causative agents of UTI are E. coli, Klebsiella spp., Pseudomonas spp., Proteus spp. and Enterococci spp., Staph. saprophyticus ² ³. Urinary tract infections result from the interaction between uropathogens and the host. Non-uropathogenic strains can induce acute infection in the presence of urological abnormalities or when the host's defense mechanisms are impaired⁴.

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The aim of the clinical microbiology laboratory in the management of UTI is to reduce the morbidity and mortality through accurate and timely identification of the aetiological agent with selection of appropriate antimicrobials through in-vitro sensitivity testing.

The media must support the growth of all urinary pathogens, inhibit possible contaminants and distinguish between pathogenic lactose and non-lactose fermenters. Routinely used conventional media like Blood agar (BA), MacConkey agar (MAC) have some drawbacks as all uropathogens cannot be cultured and differentiated in a single medium. On CLED agar medium the presence of Enterococci is frequently masked by larger colonies of gram negative bacteria. Therefore it cannot be used alone as a primary isolation medium.

The problem of urine culture can be addressed by using chromogenic agar (CA) medium. CA medium is increasingly being used as a versatile tool in early differentiation and identification of bacterial isolates from clinical specimens. This single medium supports not only the growth of all uropathogens but mixed infection can also be diagnosed more easily. Since Chromogenic agar medium facilitates direct identification of the organism on the basis of distinct colour production and characteristic colonial morphology thus it reduces the burden of biochemical characterization of the bacterial species in most instances.

The principle of chromogenic agar medium is based on the fact that bacteria posses many enzymes for their physiological function that help them to utilize substrates. In such media specific chromogenic substrates are broken down by the enzymes liberated by the particular bacteria thereby imparting a distinct colour to the growing bacterial colonies that can be visually observed.

Over the last few years, several chromogenic urine culture media have been developed and commercialized, allowing more specific and direct differentiation of microorganism on the primary plate itself. Chromogenic medium designed to isolate and identify all uropathogens. E. coli appears as pink-red colonies because of β-galactosidase production, thus allows a definite identification of E. coli without the need for further biochemical tests. Strain that produces β-glucosidase, such as Enterococci and the Klebsiella-Enterobacter-Serratia group, form colonies that generate a blue coloration as a result of hydrolysis of glucoside chromogenic substrate. Tryptophan is also present in the medium to detect members of the Proteus group, which generates a diffuse brown coloration as a result of tryptophan deaminase production. Pseudomonas spp. produce colourless colony whereas Staph. saprophyticus white coloured. An important aspect of HiCrome UTI agar is that it allows an easy differentiation of various species from mixed cultures due to specific colony colour. Another advantage of this chromogenic agar medium is that it can be used to perform antibiotic sensitivity testing without subculture onto another basic medium.

Thus chromogenic agar medium is advocated as an attractive and easy to use primary screening medium that hopefully will considerably reduce the daily workload and limits the use of further identification tests.

**Methodology**

**Patients**

Study cases are 300 clinically suspected patients of UTI of different age and sex attending either at the outpatient department (OPD) or admitted in the Rajshahi Medical College Hospital from July, 2007 to June, 2008.

Clinically suspected patients of UTI having pus cells >5/ HPF detected on microscopy of centrifuged deposit of urine were included as study cases.

**Culture of urine**

Urine samples for culture were selected on the basis of presence of pus cells >5/ HPF on microscopy. Inoculation was done aseptically with urine samples using a calibrated wire loop of 28G with an internal diameter of 3.26 mm holding 0.004 ml of urine into all three selected media plate. The plates were incubated at 37°C aerobically and after overnight incubation, they
were checked for bacterial growth. Colony count was done with bacterial growth to calculate the number of colony forming unit (CFU) or viable bacteria per ml of urine\textsuperscript{18}.

**Presumptive identification**
Bacterial growth was identified presumptively on HiCrome UTI agar, MAC agar and Blood agar, following colony characteristics against each of the uropathogens (Table-I).

The isolates were further identified using standard identification protocol such as Gram’s staining, motility test, catalase test, coagulase test, oxidase test and other relevant biochemical tests as appropriate for the isolates\textsuperscript{18}.

**Table I**: Colony characteristics of uropathogens on three urine culture media

| Organisms          | Chromogenic agar                                      | Blood agar                                                                                               | MacConkey agar                                                                                      |
|--------------------|-------------------------------------------------------|----------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------|
| *E. coli*          | Pink to Red colony due to β-D-galactosidase           | Large, grey coloured colonies, Some strains haemolytic                                                     | Smooth, Pink colonies, lactose fermenting, Some strains are late or non-lactose fermenting           |
| *Klebsiella* spp.  | Blue colour due to β-glucosidase. Large, mucoid colony| Yellow to cream or occasionally white, large colonies, Some strains beta-haemolytic                        | Mucoid, Pink colonies, lactose fermenting                                                           |
| *Enterococcus* spp.| Blue colour due to β – glucosidase, Small colony      | Non-haemolytic colonies                                                                                   | some strains lactose fermenter/non-lactose fermenter, Pale/Pink coloured colonies                  |
| *Pseudomonas* spp. | Colourless : not explained by the manufacturer        | Large flat, spreading greenish-blue coloured colonies                                                       | Pale coloured colonies, non-lactose fermenting, grape like odour                                   |
| *Staph. Aureus*    | Golden Yellow: not explained by the manufacturer      | Yellow to cream or occasionally white, large colonies Some strains beta-haemolytic                         | Smaller colonies and some strains lactose fermenting                                                |
| *Proteus* spp.     | Light brown colour due to tryptophan deaminase        | Characteristic fishy odour, and swarming growth                                                            | Pale (non-lactose fermenting) colonies                                                               |
| *Enterobacter* spp.| Blue colour due to β – glucosidase. Mucoid colony     | Produces large and mucoid colonies                                                                          | Produces large, mucoid, pink coloured colonies, lactose fermenting                                 |
| *Staph. saprophyticus* | White, small colony: not explained by the manufacturer | Produce white or yellow and non-haemolytic colonies.                                                      | Growth does not occur                                                                                 |
HiCrome UTI Agar, Modified (M1418)
1. Escherichia coli
2. Pseudomonas aeruginosa
3. Enterococcus faecalis
4. Proteus mirabilis
5. K. pneumoniae

distinguishes between Escherichia coli and Enterobacter, and also between Proteus mirabilis and other species. Coliforms produce purple coloured colonies due to cleavage of both the chromogenic substrates.

Peptic digest of animal tissue, beef extract and casein enzymic hydrolysate provides nitrogenous, carbonaceous compounds and other essential growth nutrients.

**Quality Control:**

**Appearance of Powder:**
Light yellow coloured, homogeneous, free flowing powder.

**Gelling:**
Firm, comparable with 1.5% Agar gel.

**Colour and Clarity:**
Light amber coloured, clear to slightly opalescent gel forms in petri plates.

**Reaction:**
Reaction of 5.54% w/v aqueous solution is pH 7.2 ± 0.2 at 25°C.

**Cultural Response:**
Cultural characteristics after 24 hours at 35-37°C.

| Organisms (ATCC) | Growth | Colour of colony | TDA | DMSAC |
|------------------|--------|------------------|-----|-------|
| Escherichia coli (25922) | luxuriant | pink to red | — | + |
| Proteus mirabilis (10875) | luxuriant | light brown | + | — |
| K. pneumoniae (13883) | luxuriant | blue to purple, mucoid | — | — |
| Pseudomonas aeruginosa (27653) | luxuriant | colourless | — | — |
| Staphylococcus aureus (25923) | luxuriant | golden yellow | — | — |
| Enterococcus faecalis (29212) | luxuriant | blue (small) | — | — |

Key: + = positive reaction, — = negative reaction
Results
A total of 300 patients of different age and sex suffering from UTI were included in this study. Out of 300 samples of urine cultured, a total of 139(46.33%) samples yielded significant bacterial growth and 161(53.67%) samples yielded no growth. Culture-positive cases included 133(44.33%) growth of single organism and 06(02.00%) mixed growth of two organisms each.

Table II: Pattern of bacteria isolated from urine culture (N=145)

| Bacteria         | Number | Percentage |
|------------------|--------|------------|
| E. coli          | 91     | 62.75      |
| Klebsiella spp.  | 18     | 12.41      |
| Enterococcus spp.| 16     | 11.03      |
| Pseudomonas spp. | 09     | 06.28      |
| Staph. saprophyticus | 05 | 03.44      |
| Enterobacter spp.| 04     | 02.75      |
| Proteus spp.     | 02     | 01.37      |
| Total            | 145    | 100        |

Patterns of bacterial isolates from urine culture include: 139 culture positive samples of urine yielded 145 bacterial isolates including both single (133) and polymicrobial growths (06) of two bacteria each. Out of 145 isolates, E. coli was the leading bacteria 91(62.75%) followed by Klebsiella spp. 18(12.41%), Enterococcus spp. 16(11.03%), Pseudomonas spp. 09(06.28%), Staph. saprophyticus 05(03.44%), Enterobacter spp. 04(02.75%) and Proteus spp. 02(01.37%).

Table III: Comparison of three culture media for the rate of isolation of uropathogens (N=145)

| Bacteria         | Total isolates | HiCrome UTI agar | MAC agar | Blood agar |
|------------------|----------------|------------------|----------|------------|
| E. coli          | 91             | 91(100)          | 91(100)  | 91(100)    |
| Klebsiella spp.  | 18             | 18(100)          | 18(100)  | 18(100)    |
| Enterococcus spp.| 16             | 16(100)          | 09(56.25)| 16(100)    |
| Pseudomonas spp. | 09             | 09(100)          | 09(100)  | 09(100)    |
| Staph. Saprophyticus | 05  | 05(100)          | 00       | 05(100)    |
| Enterobacter spp.| 04             | 04(100)          | 04(100)  | 04(100)    |
| Proteus spp.     | 02             | 02(100)          | 02(100)  | 02(100)    |
| Total            | 145            | 145(100)         | 133(91.72)| 145(100)   |

Regarding presumptive identification of bacterial isolates by defined colony characteristics on primary culture plates, it was found that out of total 145 bacterial isolates, 141(97.24%) could be differentially identified on chromogenic agar media and 117(80.68%) on MAC agar media whereas only 40(27.58%) on BA. The rate of presumptive identification of the isolates was found significantly higher on chromogenic agar media in comparison to MAC agar and BA (Table IV).

Table IV: Comparison of rate of presumptive identification on different media

| Bacterial strains | HiCrome UTI agar | MAC agar | Blood agar |
|------------------|------------------|----------|------------|
| E. coli (n=91)   | 88 (96.70)       | 85 (93.40)| 06(06.59)  |
| Klebsiella spp.  | 18 (100)         | 15 (83.33)| 14(77.77)  |
| Enterococcus spp.| 16 (100)         | 05 (31.25)| 10(62.50)  |
| Pseudomonas spp. | 08 (88.88)       | 07 (77.77)| 03(33.33)  |
| Staph. Saprophyticus | 05  | 00       | 05 (100)   |
| Enterobacter spp.| 04 (100)         | 03 (75.00)| 00         |
| Proteus spp.     | 02 (100)         | 02 (100)  | 02 (100)   |
| Total (N=145)    | 141 (97.24)      | 117(80.68) | 40(27.58)  |
Discussion
Urinary tract infections continue to be an important clinical problem with significant morbidity. UTI accounts for approximately 23% of all hospital acquired infections19.

For the isolation and identification of uropathogens, routinely used culture media like MacConkey agar, Blood agar, and Cystine Lactose Electrolyte Deficient (CLED) agar media are used together conventionally for long time. Each of these media has several limitations of their own for identification of the common uropathogens.

In order to overcome disadvantages of traditional urine culture media, a range of chromogenic media has become commercially available in recent years. Chromogenic agar medium supports not only the growth of all uropathogens but mixed infections can also be diagnosed properly. Since Chromogenic agar medium facilitates prompt isolation and presumptive identification of the organisms on the basis of distinct colour production and colonial morphology, thus it reduces the burden of biochemical characterization of the bacterial species in most instances.

Out of 300 urine samples, a total of 139(46.33%) samples yielded bacterial growths and 161(53.67%) had no growth. Culture-positive cases included 133(44.33%) with significant growth of single organism and remaining 06(02.00%) yielded mixed growth of two organisms.

The growths of all 145(100%) isolates were supported by chromogenic agar media and Blood agar whereas MacConkey agar yielded 133(91.72%) bacterial growths (Table-III). Blood agar is an enriched medium and chromogenic agar media contains all essential nutrients to support the growth of possible uropathogens that is why all isolates have grown on to these two media. On the other hand, the slightly lower yielding rate on MAC agar can be explained by its limitations of not supporting all organisms involved in UTI like Staph. saprophyticus and Enterococcus spp., because it is a selective medium to support the growth of members of Enterobacteriaceae.

Out of 145 isolates of uropathogens, 141(97.24%) had been presumptively identified on chromogenic agar media by matching with standard colours as stated and supplied by the manufacturer. In contrast, 117(80.68%) isolates were identified on MAC agar and only 40 (27.58%) on Blood agar media (Table-IV). The rate of presumptive identification of the uropathogens was found significantly higher on chromogenic agar media in comparison to Blood agar and MAC agar. This high rate of identification could be correlated with the ease of distinct colour production by each of the bacterial isolate on Chromogenic agar medium. Moreover, chromogenic agar media offered the advantage of limiting the spread of some isolates such as Proteus spp., Klebsiella spp. and E. coli mucoid strains thus increased the ability of the medium to detect urinary tract pathogens when mixed organisms were present12.

As far as the rate of presumptive identification of individual bacterial isolates on different culture plates is concerned, it has been seen that except E. coli (96.70%) and Pseudomonas spp. (88.88%), all bacteria could be differentially identified onto the chromogenic agar media. While, MAC agar provided 93.40% identification for E. coli, followed by 83.33% for Klebsiella spp., 31.25% for Enterococcus spp., 77.77% for Pseudomonas spp., 75% for Enterobacter spp., 100% for Proteus spp. and 00% for Staph. saprophyticus. Blood agar also showed variable rate of identification ranging from 00 to 77.77% for all bacterial isolates except Proteus spp. and Staph. saprophyticus, which were identified 100% on BA plates (Table-V).

The reason behind the highest rate of identification of E. coli, Klebsiella spp., Enterococci spp. and Enterobacter spp. on chromogenic agar media in comparison to other two conventional media used is that each of these organisms produces characteristic, distinct and easily perceivable colour on chromogenic agar media. In fact, this differential colour production by individual bacterial isolates is among the exciting features of chromogenic agar for which it has been advocated to be used as primary culture medium. The chromogenic media also provided added
advantage on identification of a few non-lactose variety of *E. coli*, which might be the reason of decreased rate of identification on MAC agar.

However, chromogenic agar media failed to produce expected colony colours for 03(03.33%) of the *E. coli* and 01(11.12%) *Pseudomonas* spp. This failure can be correlated with inadequate production of enzymes at that point of time or absence of enzymes in those strains.

The chromogenic agar media also reigned over the conventional media by providing specific identifying characteristics of the organisms isolated in mixed growth. As it was found in this study, all 06(100%) of the isolate-pairs of mixed growth (*E. coli* and *Enterococci* spp., *E. coli* and *Klebsiella* spp., *E. coli* and *Proteus* spp., *E. coli* and *Staph. saprophyticus*) were all identified distinctly on HiCrome UTI agar, whereas only 01(16.66%) in combination of *E. coli* and *Proteus* spp. was identified on both BA and MAC agar media (Table-VI). The rate of identification of mixed culture on the BA and MAC agar was poor due to difficulty in differentiating the colonies.

It is obvious from the results of the present and similar studies that chromogenic media can replace CLED and MAC agar in order to identify uropathogens more efficiently by its characteristic colony colour for each of the organism in both community acquired and nosocomial UTI cases. Moreover, chromogenic media also provide an added advantage of requiring less time in mastering the skill in the identification of the uropathogens. This will be an attractive easy to use primary screening medium that considerably reduce the daily workload and thus minimize the use of troublesome biochemical tests.

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