Two new techniques for obtaining consistent results when growing *Renibacterium salmoninarum* on KDM2 culture medium

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ABSTRACT: We recently showed that the inconsistent performance of the culture medium KDM2 in supporting growth of the fastidious kidney disease agent (*Renibacterium salmoninarum* = Rs) was due to a lot-to-lot variation in the composition of a commercially available peptone used as a medium ingredient. We now report on 2 procedures for obtaining consistent growth of the pathogen on KDM2, regardless of the lot of peptone used in preparing the medium. The first procedure involves the recently described nurse culture technique that has the added advantages of accelerating growth of Rs and of increasing the sensitivity with which the pathogen can be detected. The second procedure is to supplement KDM2 with a small amount of KDM2 broth that has previously been used to grow Rs (= spent broth). A metabolite (or metabolites) produced by the pathogen during its growth in the broth appears to substitute effectively for the nurse culture by providing the Rs growth initiating factor(s) that, apparently, is in short supply in certain lots of peptone. These techniques remove the need to pre-test peptone lots to ensure that they will perform satisfactorily when used in KDM2 for growing Rs.

INTRODUCTION

*Renibacterium salmoninarum* (Rs), the causative agent of bacterial kidney disease (BKD) in salmonids, is a fastidious and slow-growing organism. Recently, Evelyn et al. (1989) described a technique, the nurse culture technique, that significantly accelerates the growth of the pathogen when the organism is present in low numbers. The technique takes advantage of the fact that low numbers of Rs cells always show accelerated growth when they are plated on an agar medium (KDM2) adjacent to a nurse culture consisting of a heavy inoculum of Rs cells. Until recently, it was thought that growth acceleration of Rs and an increased sensitivity at detecting the pathogen were the only benefits of the technique. The experiments outlined below, however, highlight another advantage of the nurse culture technique. We have reported frustrating instances in which the normally satisfactory Rs culture medium, KDM2, failed to support growth of the pathogen (Evelyn & Prosperi-Porta 1989). An investigation of these occurrences revealed that they were due to the particular production lot of peptone used in preparing the culture medium. Production lots of peptone varied enormously in their growth promoting properties for Rs even when they were produced by the same manufacturer. The purpose of this paper is to show that growth failures of Rs on KDM2 due to the inadvertent use of faulty lots of peptone can be avoided by (1) using the nurse culture technique or (2) supplementing the growth medium with an unidentified metabolite(s) produced by the nurse culture.

MATERIALS AND METHODS

General. Two experiments were performed. The first examined the effect of the nurse culture technique on the growth of Rs resulting on batches of KDM2 prepared with production lots of peptone that had previously been shown to be satisfactory or unsatisfactory at promoting growth of Rs (Evelyn & Prosperi-Porta 1989). In the second experiment, Rs was grown from a heavy inoculum in KDM2 broth and the spent broth was tested to determine whether it contained Rs metabolites that would substitute for the nurse culture in KDM2 prepared with an unsatisfactory lot of peptone. The 3 Rs isolates used in the experiments (Rs 384, Rs...
8894, Rs 9902) were obtained from different stocks of cultured chinook salmon *Oncorhynchus tshawytscha*, affected with BKD. Each isolate was typical of Rs and 2 of them have been described elsewhere (Evelyn et al. 1989). The Rs cultures were 8 to 16 d old when used in the experiments. The 4 lots of peptone used for preparing the various batches of KDM2 had earlier been shown to be satisfactory (Lot 4) or unsatisfactory (Lots 2, 3, and 7) for promoting the growth of Rs (Evelyn & Prosperi-Porta 1988, Evelyn et al. unpubl.).

In the first experiment, 3 different batches of KDM2 (Evelyn & Prosperi-Porta 1989) were prepared, each with a different production lot of peptone (Lot 2, 3, or 4). Petri plates, poured with a particular batch of KDM2, were then drop-inoculated (Miles & Misra 1938) peripherally with 25 μl aliquots of Rs cells suspended in peptone (0.1%) saline (0.85% NaCl) (= PS). The inocula consisted of cells of Rs 8894 or Rs 9902 that were suspended in PS to a turbidity of 1.25 OD at 540 nm and then diluted to 10⁻³, 10⁻⁴, and 10⁻⁵. A single dilution of a single Rs isolate was inoculated onto any given plate. Those plates inoculated peripherally with Rs 8894 were simultaneously inoculated centrally with a 25 μl drop of nurse culture (Rs 9902 cells in PS at 1.25 OD at 540 nm) or with a 25 μl drop of sterile PS (controls). Plates inoculated peripherally with Rs 9902 were simultaneously inoculated centrally with a 25 μl drop of nurse culture (Rs 384 cells in PS at 1.25 OD at 540 nm) or with a 25 μl drop of sterile PS (controls). Following imbibition of the drops of inoculum, plates were sealed in plastic bags to prevent desiccation, and incubated inverted, at 15°C for 20 or 21 d. At this stage, counts of colonies developing from each 25 μl drop of 10⁻³-diluted suspension were made using a stereomicroscope at a magnification of 120. In addition, the turbidity yielded by the growth from the drops of the variously diluted suspensions was measured at 540 nm, using the procedure outlined earlier (Evelyn et al. 1989).

In Expt 2, 2 batches of KDM2 were prepared, one with a satisfactory lot of peptone (Lot 4), the other with an unsatisfactory lot of peptone (Lot 7). Petri plates, poured with these production lots of peptone (1 batch of KDM2 plate⁻¹), were drop-inoculated peripherally with Rs 384 cells (25 μl drops of cells suspended in PS to a turbidity of 1.25 OD at 540 nm and diluted to 10⁻³, 10⁻⁴, and 10⁻⁵). These plates were not inoculated with nurse cultures. Instead, they were 'inoculated' centrally with spent, sterile, 10× concentrated KDM2 broth, the preparation of which is described below, or with fresh (unspent), sterile, 10× concentrated KDM2 broth (controls). Spent KDM2 broth was prepared as follows. Several loopfuls of Rs 384 cells from a KDM2 plate were inoculated into a flask containing 100 ml of KDM2 broth (prepared with peptone Lot 4) and the culture was incubated at 15°C with occasional shaking until a turbidity of 0.83 OD at 540 nm was reached (18 d). At this point, the culture was centrifuged to remove the Rs cells and other particulate matter (12 000 × g for 30 min at 4°C). The clarified supernatant was then freeze-dried and redissolved in 1/10th of its original volume using distilled water. The concentrated supernatant was then filter-sterilized directly, using a 0.22 μm pore diameter filter (Millipore), or following its passage through a molecular sieve (nominal molecular weight limit of 10 000 Daltons, Millipore). In addition, sieved material was also sterilized by autoclaving (121°C for 15 min). Samples (25 μl) of the variously treated spent broth concentrate were applied centrally (in place of the nurse culture) on test plates of KDM2 (1 sample plate⁻¹). Control plates were 'inoculated' centrally with 25 μl of sterile, 10× concentrated, unspent KDM2 broth. Plates were then incubated, as described above, for 19 d, at which time the resulting Rs colonies were recorded turbidimetrically (10⁻³ sample) and photographically (10⁻⁴ sample), or counted (10⁻⁵ sample).

**RESULTS AND DISCUSSION**

Results indicate that when the nurse culture technique was used, growth of Rs was consistently obtained on all batches of KDM2, irrespective of the lot number of the peptone used to prepare the medium. In contrast, when nurse cultures were not used, only that batch of KDM2 prepared with a peptone lot (Lot 4), previously shown to promote satisfactory growth of Rs, supported growth of Rs. These results held true for each of the 2 Rs isolates tested (Table 1), indicating that the nurse culture phenomenon is not specific to a particular isolate. Clearly, reproducible results in growing Rs depend on using batches of KDM2 prepared with previously proven peptone lots or on using the nurse culture technique. The latter approach, however, provides 2 additional advantages: faster Rs growth and increased sensitivity in detecting Rs cells (Evelyn et al. 1989). These advantages were again obvious with the 2 Rs isolates used in the present experiment when their short term (20 or 21 d) growth, obtained from the more highly diluted samples (the 10⁻³ and 10⁻⁴ diluted samples), was considered. In all cases, there was no detectable growth on any of the batches of KDM2 media — not even on the batch of KDM2 prepared with the satisfactory lot (Lot 4) of peptone — when the nurse culture technique was not used. In contrast, when nursed, both isolates grew well on all batches of KDM2 even at these dilutions of inoculum (turbidities on the various batches of medium inoculated with the highest dilution of inoculum averaged, for example, 0.100 OD for Rs 8894 and 0.096 OD for Rs 9902).
Table 1. Use of nurse culture of *Renibacterium salmoninarum* (Rs) results in consistent growth of RS isolates on KDM2 culture medium, irrespective of the lot of peptone used in preparing the medium.

<table>
<thead>
<tr>
<th>Peptone lot</th>
<th>Classification of peptone lot</th>
<th>Growthb (OD 540 nm) of:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Rs 384 Nursed with Rs 9902</td>
</tr>
<tr>
<td>2</td>
<td>Unsatisfactory</td>
<td>0.450 ± 0.061</td>
</tr>
<tr>
<td>3</td>
<td>Unsatisfactory</td>
<td>0.447 ± 0.051</td>
</tr>
<tr>
<td>4</td>
<td>Satisfactory</td>
<td>0.484 ± 0.062</td>
</tr>
</tbody>
</table>

a These lots of peptone had earlier been shown to promote satisfactory or unsatisfactory growth of Rs (Evelyn & Prosperi-Porta 1989)

b Growth at 20 d (Rs 384) or 21 d (Rs 9902) from inocula containing $3.1 \times 10^4$ Rs 384 cells or $4.5 \times 10^4$ Rs 9902 cells per 25 µl drop. Turbidity values (± SD) were produced by the growth resulting from the 25 µl drop and are averages of 6 replicate drops.

Results obtained in Expt 2 indicate that Rs 384 cells produce a metabolite that stimulates growth of Rs cells and that can be successfully substituted for the nurse culture (Fig. 1, Table 2). The growth stimulating effect of the Rs 384 metabolite was not Rs isolate specific because it also stimulated the growth of the 2 other Rs isolates tested (Rs 8894, Rs 9902; data not shown). We conclude that the metabolite is in short supply in those lots of peptone (e.g. Lots 2, 3, 7) that fail to promote satisfactory growth of Rs because when it is provided, those lots of peptone yield satisfactory growth of the pathogen.

The identity of the metabolite is not known, but it is a small, heat-stable molecule because it passed through a molecular sieve of 10 000 MW exclusion limit and it resisted autoclaving (121°C for 15 min). Thus, when sieved or sieved and autoclaved spent broth was tested in place of filter-sterilized spent broth, growth stimulation similar to that illustrated in Fig. 1 for the filter-sterilized spent broth was obtained. The identity of the factor is being investigated because it would greatly simplify growing Rs if the medium could be formulated using known, commercially available ingredients. Until, however, the active factor can be identified, 2

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Fig. 1. Use of spent KDM2 broth (previously used for growing *Renibacterium salmoninarum* = RS) ensures growth of Rs on a batch of KDM2 prepared with an unsatisfactory lot of peptone (Lot 3). The Rs isolate (Rs 384), inoculated peripherally on the 2 plates, shows no signs of growth on this particular batch of KDM2 at 19 d of incubation when the supplement (white central spot) was fresh, 10× concentrated KDM2 broth (A); but shows good growth when the supplement (white central spot) was spent, 10× concentrated KDM2 broth (B). The white central spots are the result of the dried, precipitated, bovine serum proteins present in the KDM2 broth used as the supplement. Note: stimulation of growth similar to that shown in B was obtained when molecular sieved or molecular sieved and autoclaved spent broth concentrate was substituted for the filter-sterilized preparation used in B.
Table 2. Use of a spent broth supplement* results in consistent growth of *Renibacterium salmoninarum* (Rs 384) on KDM2 culture medium, irrespective of the lot of peptone used in preparing the medium.

<table>
<thead>
<tr>
<th>Peptone Lot</th>
<th>Classification of peptone lotb</th>
<th>Supplement</th>
<th>No supplement</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>Satisfactory</td>
<td>0.791 ± 0.050</td>
<td>0.595 ± 0.070</td>
</tr>
<tr>
<td>7</td>
<td>Unsatisfactory</td>
<td>0.562 ± 0.070</td>
<td>0.000</td>
</tr>
</tbody>
</table>

*Spent broth supplement = KDM2 broth that had been used to grow Rs, was freed of the Rs cells, and then used in place of a nurse culture following filter sterilization.

*bThese lots of peptone had earlier been shown to be satisfactory (Evelyn et al. 1989b) or unsatisfactory (Evelyn et al. unpubl.) at promoting growth of Rs.

**Growth at 19 d from an inoculum containing 4.5 x 10^4 Rs cells per 25 μl drop. Turbidity values (± standard deviation) were produced by the growth resulting from the 25 μl drop and are averages of 7 replicate drops.

Procedures are available for reproducibly growing Rs: the nurse culture technique and a method based on supplementation of KDM2 with small amounts of spent KDM2 broth, produced as outlined above. It is not necessary to add the supplement to the center of an already poured agar plate as was done in Expt 2. Addition of the spent broth supplement may be conveniently done when adding the serum supplement to the KDM2 basal medium. (If supplement sterilized by autoclaving is used, a cloudy medium will result due to the presence of heat-precipitated proteins). Recent results, not reported on here, indicate that supplementation of KDM2 with quantities of unconcentrated spent broth of only 1.5% (v/v) will suffice to ensure reproducible Rs growth on KDM2, regardless of the peptone lot used in preparing the medium. In addition, we also found that the satisfactory spent broth supplements can be obtained with batches of broth prepared with faulty peptone lots (e.g. Lot 3). With such batches of broth, however, care must be taken to use a heavy enough inoculum of Rs to ensure that it grows to produce the required metabolite(s). Finally, although the supplement can be safely stored at 4°C for 2 wk, we advise storing the supplement in the frozen state until more is known about its stability.

**LITERATURE CITED**


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