Klebsiella species in hospital food and kitchens: a source of organisms in the bowel of patients

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SUMMARY

Hospital food was examined for the presence of Klebsiella species. Salads and cold meat were most frequently contaminated, often containing more than $10^3$ organisms per g. Klebsiella species were also widely distributed in the kitchen environment which was considered, at least in part, to be the source of the organisms in food.

By the use of serological and bacteriocin typing, intestinal carriage of strains ingested in food could be demonstrated.

INTRODUCTION

The epidemiology of klebsiella infections in hospitals is not well defined (Lancet, 1971). Selden et al. (1971) have demonstrated that an important source of klebsiella strains causing infections may be the patient’s own bowel. In addition to being a potential source of autoinfection, the acquisition of a strain in the bowel during hospitalization provides a possible source for transmission of the organism.

If intestinal carriage of klebsiella is an important source of infection, then the sources of the strains colonizing the patient’s bowel are of interest. Hospital food has been shown to contain klebsiellas in numbers great enough to suggest that colonization of the bowel might occur (Montgomerie et al. 1970; Shooter et al. 1971) and infant feeds have also been shown to be contaminated (Ayliffe, Collins & Pettit, 1970).

The three surveys reported in this paper were designed to examine the extent of klebsiella contamination of food prepared in a hospital kitchen, to define in greater detail the sites at which food became contaminated with klebsiella by examining the kitchen environment and food both before and after preparation, and to determine whether ingestion of klebsiellas in food could lead to intestinal colonization.

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MATERIALS AND METHODS

Survey 1

Food prepared in the hospital kitchen was sampled daily for 16 weeks and examined for klebsiellas.

Survey 2

During a 7-week period work surfaces, floors, utensils and sinks in a hospital kitchen were examined twice weekly for klebsiellas. Food at different stages of preparation was also sampled.

Survey 3

Every portion of food and drink and any medicaments taken in 1 week by five patients in a female orthopaedic ward was sampled. During the week faecal specimens were obtained from each patient and examined for klebsiellas. Faecal specimens were also sampled during the week before the food sampling and during the week after food sampling had ended. Patients’ hands were examined for klebsiellas on one occasion. The patient’s locker and articles on it were sampled weekly.

Bacteriology

In Survey 1 a volume of sterile water equal to the weight of the food sample in grams was added to each sample and suspensions of food were made using either a wrist-action shaker or a stomacher. Each suspension was inoculated on MacConkey agar (Difco), Endo agar (Difco) and deoxycholate agar (Difco) and plates were incubated at 37 °C for 48 h. The number of klebsiella colonies was counted and klebsiellas were identified biochemically as previously (Cooke et al. 1979).

In Surveys 2 and 3 solid food samples were mixed with 1/4 strength Ringer’s solution in the proportion of 4 ml of Ringer’s solution to 1 g of food, and were homogenized in a stomacher. Volumes of 0.1 ml of suspension were spread on MacConkey agar, citrate agar (Difco) and MacConkey-inositol-carbenicillin (MIC) agar (Cooke et al. 1979) and were also inoculated into citrate broth. Liquid samples were diluted tenfold in 1/4 strength Ringer’s solution and plated out as for solid food samples.

When examining environmental sites hard surfaces were sampled by swabbing with a cotton wool swab moistened in nutrient broth, which was then immersed in a tube containing 1 ml of 1/4 strength Ringer’s solution. The tube was agitated on a vortex mixer and the resulting suspensions were inoculated to the same media as for food.

All plates were incubated at 37 °C for 24 h and up to ten klebsiella colonies were selected from each food and environmental sample. Where plate cultures were negative after 24 h, the citrate broth was subcultured on MacConkey, citrate and MIC agar plates.

Faeces were examined as previously (Cooke et al. 1979).

Patients’ hands were sampled by agitation in 1/4 strength Ringer’s solution, which was filtered through 0.45 μm membrane filters; the filters were then incubated on MacConkey agar at 37 °C for 24 h.
Table 1. Isolation of Klebsiella from food

<table>
<thead>
<tr>
<th>Type of food</th>
<th>No. of samples</th>
<th>No. of samples containing Klebsiella (percentage in parentheses)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Salads</td>
<td>211</td>
<td>49 (23.2)</td>
</tr>
<tr>
<td>Cold meat</td>
<td>54</td>
<td>14 (25.9)</td>
</tr>
<tr>
<td>Other cold food</td>
<td>443</td>
<td>13 (2.9)</td>
</tr>
<tr>
<td>Hot food</td>
<td>637</td>
<td>18 (2.8)</td>
</tr>
</tbody>
</table>

Table 2. Isolation of Klebsiella from the kitchen environment

<table>
<thead>
<tr>
<th>Source of sample</th>
<th>No. of samples</th>
<th>No. of samples containing Klebsiella (percentage in parentheses)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surfaces</td>
<td>53</td>
<td>23 (43.4)</td>
</tr>
<tr>
<td>Utensils</td>
<td>87</td>
<td>34 (39.1)</td>
</tr>
<tr>
<td>Sinks</td>
<td>57</td>
<td>37 (65.0)</td>
</tr>
<tr>
<td>Washing-up water</td>
<td>16</td>
<td>8 (50.0)</td>
</tr>
<tr>
<td>Water used for soaking</td>
<td>12</td>
<td>8 (66.7)</td>
</tr>
</tbody>
</table>

Typing methods

All klebsiella strains were serotyped by the quellung method (Kauffmann, 1949), using 77 capsular antisera produced in this laboratory. Strains which were thought to be related were also bacteriocin typed by the method of Edmondson & Cooke (1979).

RESULTS

The results from the three surveys of the isolation of Klebsiella from food prepared in the hospital kitchen are given in Table 1. The number of klebsiellas in different types of food varied from 10 to > 10⁵/g, salads being the most heavily contaminated.

Food was also sampled as it entered the kitchen. Of 136 salad constituents sampled before preparation, 14 (10.3%) contained Klebsiella while 6 of 13 (46%) raw meat samples contained Klebsiella.

The contamination of the kitchen environment with Klebsiella is shown in Table 2. Estimations of the numbers of klebsiella organisms in the kitchen environment could not be made in many cases as there was frequently confluent growth on the isolation media. Where estimations could be made there were 10³–10⁴ organisms/ml of suspension prepared from an environmental sample.

Forty-six different serotypes were isolated from food and 50 types were isolated from the kitchen environment. Thirty-four serotypes were found in both situations.

Table 3 shows the serotypes of klebsiella strains isolated from food, faeces and the immediate environment (i.e. the locker and any articles on it) of four patients studied in Survey 3. No klebsiellas were isolated from any source from the remaining patient examined in this survey.

There was no evidence of acquisition of klebsiellas from food in patients A and B, for the non-typable strains isolated from the food and faeces of the two patients had distinct bacteriocin typing patterns. Similarly the two strains of serotype 35
isolated from patient B had distinct bacteriocin typing patterns and had differing antibiotic sensitivity patterns and biochemical reactions. Evidence of acquisition was obtained for patient C, as this patient ingested klebsiellas of serotype 24 on three occasions, and the same serotype was isolated from the faeces of the patient on four occasions. The first isolation of type 24 from faeces was made on the day following the first ingestion of this type, and the final isolation was made three days after the examination of food had ended. The bacteriocin typing patterns, biochemical reactions and antibiotic sensitivity patterns of these seven cultures of serotype 24 suggested that they were identical. Patient D ingested klebsiellas of type G23 on two occasions and klebsiellas of the same serotype and bacteriocin typing pattern were isolated from the faeces of this patient on two later occasions.

There was no correlation between the serotypes found in the environment of the patients and those isolated from the food and faeces of the patients. No klebsiellas were isolated from the patients’ hands, nor from any medicaments.

**DISCUSSION**

The results of the three surveys described in this paper show that all types of hospital food may be contaminated with large numbers of klebsiella organisms. Klebsiellas were most frequently isolated from salads, as has been reported in other surveys (Shooter et al. 1971; Wright, Kominos & Yee, 1976) but were also very common in cold meat. There was a considerable difference between the number of klebsiella isolations from salads which had been prepared in the hospital kitchen and salads before preparation since 23.2% of prepared salads contained klebsiella while only 10.3% of salads sampled as they entered the kitchen were contaminated with klebsiella. This suggests that salads became contaminated during preparation. The similarity of the serotypes of klebsiella strains isolated from food and the kitchen environment also suggests that contamination of food from the environment may have occurred.

A total of 62 different serotypes were found in food and the kitchen environment from the three surveys, and no one serotype predominated. A similar range of
Klebsiella spp. in hospital food

Serotypes has been isolated from infections in the hospital, again with no particularly common single serotype.

The acquisition in the gastro-intestinal tract of klebsiella serotypes ingested in food has previously been reported in renal-transplant patients (Montgomerie et al. 1970) and in intensive-care patients (Casewell & Phillips, 1978). In this paper we demonstrate intestinal colonization by klebsiella strains ingested in food in two patients using two sensitive epidemiological typing methods, capsular serotyping and bacteriocin typing. The use of more than one typing method is valuable in studies of this kind as it improves discrimination between strains.

The results obtained here confirm those of other workers showing that hospital food may be contaminated with klebsiellas, and indicate that the source of the organisms may, at least in part, be the kitchen environment.

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REFERENCES


