A simple and novel method for retrieval of Pasteurellaceae from swab samples collected in the field

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Abstract
Traditionally it has been difficult or impossible to collect and preserve bacterial samples of especially fastidious bacteria in mixed primary cultures, unless the samples could be transported to a laboratory within approximately 24 h. Therefore, a simple novel method for preserving swab samples until bacterial isolation can be completed in the laboratory was developed and evaluated. Pasteurellaceae bacteria were used as a representative for fastidious bacteria. A 7.5% glucose serum medium was used as freeze medium. Swab samples were soaked in the medium a maximum of 2 h after collection and stored at –20°C.

Introduction
Collection and preservation of bacterial samples in the field can be very challenging, especially due to the distance to proper laboratory facilities. When dealing with fastidious bacteria that survive poorly outside the host, quick transportation of the samples to the laboratory can be an impossible obstacle to overcome. Commercially available culture swabs may alleviate this problem to some extent, but when the transport time exceeds 24 h, the probability of isolating host-dependent bacteria, like Pasteurellaceae, decreases rapidly (Schwarz 2008).

The Pasteurellaceae family is a large and diverse family of obligate parasites most of which are closely related to a single vertebrate host (Christensen and Bisgaard 2008). Pasteurellaceae typically colonize the upper respiratory tract, the reproductive tract, and perhaps also parts of the intestinal tract (Olsen et al. 2005; Christensen and Bisgaard 2008). Pasteurellaceae bacteria are classified as
fastidious bacteria, and often require specific growth factors outside the host (Schwarz 2008).

The aim of this study was to develop and test a system to preserve swab samples until bacterial isolation could be completed in a microbiological laboratory. *Pasteurellaceae* bacteria were used as representatives for fastidious bacteria.

**Materials and Methods**

The study consisted of two parts, a control study and a field study, respectively. BBL culture swabs (BD Biosciences, Le Point de Clai, France) were used for swabbing the canine teeth gingival/dental fossa in all included animals.

The control study was conducted using samples from a single captive brown bear (*Ursus arctos*) (Table S1), collected while it was under anesthesia for veterinary procedures. Samples for the field study were collected from eight wild polar bears (*Ursus maritimus*) (Table S1), shot by local subsistence hunters near Scoresbysound, Central East Greenland (69°00′N and 74°00′N) in March–February 2011 as part of the legal hunting quota. The polar bear samples were stored at −20°C for 240 to 259 days prior to and during sea transport to Denmark.

Swabs were kept in Stuart semi liquid medium after sampling at ambient temperatures (four to 20°C) for up to 2 h and then soaked for 3 sec in 0.25 mL of a custom made 7.5% glucose serum medium (Lapage and Redway 1974) (15 mL sterile 50% glucose solution [SAD, 2100 København Ø, Denmark] was mixed with 100 mL sterile calf serum [In Vitro, 3480 Fredensborg, Denmark] and stored at −20°C). The swabs were then stored at −20°C until further processing. Following storage, swabs were thawed for 30 min at 20°C and plated on 5% bovine blood agar (BA) (Blood agar base, CM55; Oxid, Roskilde, Denmark). BA plates were incubated aerobically in sealed plastic bags for 24 h at 37°C.

In the control study, a total of 15 samples were retrieved from a brown bear kept in captivity (Table S1). Three control swabs were not dipped in the glucose serum medium prior to plating on BA, where one was plated 1 h after collection. The two others were stored at −20°C for 7 and 30 days, respectively, prior to thawing and plating on BA. The remaining 12 brown bear swabs were preserved using the glucose serum medium and stored at −20°C for a variable time period. After 7 days of storage, and then once every 30 days during 330 days, a swab was thawed and plated. The flora was described based on colony morphology and the total number of colonies on the plates were counted.

The polar bear samples from the field study were plated 240 to 259 days after collection (Table S1). Colonies typi-
Table 1. Total colony plate count for brown bear samples with and without freeze medium.

| Time after sampling | Total colony plate count with medium | Total colony plate count without medium |
|---------------------|-------------------------------------|----------------------------------------|
| 1 h (no freezing)   | –                                   | 3744                                   |
| 7 days              | 3468                                | 23                                     |
| 30 days             | 3876                                | 0                                      |
| 60 days             | 4092                                | –                                      |
| 90 days             | 3528                                | –                                      |
| 120 days            | 4116                                | –                                      |
| 150 days            | 3480                                | –                                      |
| 180 days            | 3504                                | –                                      |
| 210 days            | 3732                                | –                                      |
| 240 days            | 3432                                | –                                      |
| 270 days            | 3984                                | –                                      |
| 300 days            | 3624                                | –                                      |
| 330 days            | 3876                                | –                                      |

–20°C. In contrast, unpreserved swab samples yielded no Pasteurellaceae growth following just 1 week of storage.

Also, it is noteworthy that the total colony plate count variation was independent of storage time and that all samples showed a similar nonspecific mixed flora.

Besides Pasteurellaceae, Neisseriaceae, which are also classified as fastidious bacteria, were also preserved with the method, thus underlining the ability of this method to preserve a mixed bacterial flora including an array of fastidious bacteria.

The polar bear samples were taken in the field in Greenland and were transported to Denmark in a freezer by boat, which proves that the method is very usable for remote sampling and consecutive long lasting storage and transport.

In summary, the method has demonstrated that even fastidious bacteria in a mixed primary culture can survive for many months and allow relevant microbiological procedures on field samples.

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Conflict of Interest

None declared.

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Supporting Information

Additional Supporting Information may be found in the online version of this article:

Table S1. Animals included in the study.