Diagnostic accuracy of confocal microscopy imaging vs. punch biopsy for diagnosing and subtyping basal cell carcinoma

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Abstract

Background In vivo reflectance confocal microscopy (RCM) is a promising non-invasive skin imaging technique that could facilitate early diagnosis of basal cell carcinoma (BCC) instead of routine punch biopsies. However, the clinical value and utility of RCM vs. a punch biopsy in diagnosing and subtyping BCC is unknown.

Objective To assess diagnostic accuracy of RCM vs. punch biopsy for diagnosing and subtyping clinically suspected primary BCC.

Methods A prospective, consecutive cohort of 100 patients with clinically suspected BCC were included at two tertiary hospitals in Amsterdam, the Netherlands, between 3 February 2015 and 2 October 2015. Patients were randomized between two test-treatment pathways: diagnosing and subtyping using RCM imaging followed by direct surgical excision (RCM one-stop-shop) or planned excision based upon the histological diagnosis and subtype of punch biopsy (standard care). The primary outcome was the agreement between the index tests (RCM vs. punch biopsy) and reference standard (excision specimen) in correctly diagnosing BCC. The secondary outcome was the agreement between the index tests and reference standard in correctly identifying the most aggressive BCC subtype.

Results Sensitivity to detect BCC was similar for RCM and punch biopsy (100% vs. 93.94%), but a punch biopsy was more specific than RCM (79% vs. 38%). RCM expert evaluation for diagnosing BCC had a sensitivity of 100% and a specificity of 75%. The agreement between RCM and excision specimen in identifying the most aggressive BCC subtype ranged from 50% to 85% vs. 77% by a punch biopsy.

Conclusion Reflectance confocal microscopy and punch biopsy have comparable diagnostic accuracy to diagnose and subtype BCC depending on RCM experience. Although experienced RCM users could accurately diagnose BCC at a distance, we found an important difference in subtyping BCC. Future RCM studies need to focus on diagnostic accuracy, reliability and specific criteria to improve BCC subtype differentiation.

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Conflicts of interest

We declare that we have no conflict of interests.

Financial disclosure and products

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Introduction

Current international guidelines recommend a punch biopsy of clinically suspected basal cell carcinoma (BCC) to confirm clinical diagnosis and classify into histological subtypes (superficial, nodular and aggressive) to ensure optimal treatment selection.1,2 Although considered as the most reliable diagnostic technique, a
punch biopsy fails to diagnose an aggressive subtype in up to one of six BCCs. Other obvious disadvantages of a punch biopsy are pain and discomfort for the patient, scarring, doctor’s delay in the diagnostic process and costs for the healthcare system.

Ideally, BCC diagnosing and subtyping should be performed non-invasively by a painless procedure leading to an immediate diagnosis and treatment. This could be particularly relevant to the growing use of topical treatments as non-surgical first-line therapy for superficial BCC. As for nodular and aggressive BCC, surgical treatment with excision margins of, respectively, 3 or 5 mm remains the treatment of choice, reserving Mohs micrographic surgery for primary BCC on high-risk facial areas (depending on tumour size) and recurrent or previously incompletely excised BCC in all facial areas.

Recently, we performed a randomized controlled trial to assess the efficacy of a one-stop-shop concept with real-time in vivo reflectance confocal microscopy imaging (RCM) vs. standard care for surgical treatment of BCC. To determine the clinical value and utility of RCM vs. a punch biopsy in diagnosing and subtyping BCC, a diagnostic analysis is needed. Besides, previous diagnostic testing of RCM and punch biopsy for BCC was not done in accordance with the Standards of Reporting of Diagnostic Accuracy (STARD).

The aim of this study was to assess diagnostic accuracy of RCM vs. punch biopsy for diagnosing and subtyping clinically suspected primary BCC.

Methods

Study design

Diagnostic accuracy data were prospectively collected alongside the multicentre non-inferiority clinical trial in which patients with clinically suspected primary BCC were randomized between two test-treatment pathways: diagnosing and subtyping using RCM followed by direct surgical excision (RCM one-stop-shop) or planned excision based upon the histological diagnosis and subtype of a punch biopsy (standard care). Full details of the clinical trial are given elsewhere.

Participants

Between 3 February 2015 and 2 October 2015, 100 patients presented with clinically suspected BCC were consecutively included at the Department of Dermatology, Academic Medical Centre, University of Amsterdam (coordinating tertiary hospital), and the Department of Dermatology, the Netherlands Cancer Institute (participating tertiary hospital), in Amsterdam, the Netherlands. We included patients older than 18 years with previously untreated lesions, lesions suitable for conventional surgical excision and lesions present for at least 1 month. We excluded lesions on high-risk areas of the face (H-zone and ears), lesions larger than 20 mm, recurrent BCC, lesions not suitable for RCM (macroscopic ulceration or crust) and patients with basal cell nevus syndrome. Immunocompromised patients were not excluded.

Initial clinical assessment, most times including dermoscopy, was performed by experienced dermatologists. Patients with multiple clinically suspected new primary BCC were included for only one lesion being the most suitable for conventional surgical treatment according to the following order: (i) chest, (ii) extremities, and (iii) head and neck area.

Test methods

RCM (index test 1) Patients allocated to the RCM one-stop-shop group prospectively received RCM (VivaScope 1500; CaliberID, Henrietta, NY, USA; MAVIG GmbH, München, Germany) to diagnose and subtype BCC followed by direct surgical excision according to previously published protocol. DK performed RCM imaging including subsequent diagnosing of RCM cases at the Academic Medical Centre, and YE did the same at the Netherlands Cancer Institute. At the time of RCM, both assessors were masked to the results of surgical excision specimen but not to patients’ clinical history.

Prior to the study, DK and YE were trained in RCM and interpretation of the acquired images during a 1-week ‘Expert training in Confocal Laser Scanning Microscopy’ course organized by MAVIG GmbH (distributor of the VivaScope device) at the University of Modena in Italy. DK and YE had <1 year of RCM experience prior to the start of the study. After the trial was completed two independent international RCM experts (CI and MU) evaluated the RCM images for BCC presence and subtype through a secured online teleconsultation platform designed to share RCM cases (VivaNet, MAVIG, GmbH). The experts were masked to the results of surgical excision specimen as well as patients’ clinical history. Both experts had more than 10 years RCM experience.

Punch biopsy (index test 2) Patients allocated to the standard care group received planned excision after a punch biopsy was performed. The routine 3-mm punch biopsy was performed from the most elevated part of the lesion using infiltration anaesthesia (2% xylocaine/adrenaline 1 : 80 000). Biopsy specimens were subsequently analysed by an experienced pathologist within 2 weeks. Surgical excision of the lesion with adequate margins was performed within the following 4 weeks after receiving the report of the punch biopsy. At the time of the punch biopsy, the pathologists were masked to the results of surgical excision specimen but not to patients’ clinical history.

Surgical excision (reference standard) Histopathological confirmation of presence and subtype of BCC and inspection of resection margins with the use of haematoxylin and eosin...
stained sections taken from the excision specimen was defined as the reference standard.

An independent dermatologist or independent dermatology resident supervised by a dermatologist performed surgery under local anaesthetics (2% xylocaine/adrenaline 1 : 80 000) followed by primary wound closure in both treatment groups. Clinically suspected BCCs that were not confirmed by either RCM or punch biopsy were surgically treated with a 3-mm excision margin. To prevent bias DK and YE did not perform the subsequent surgical procedures.

After formalin fixation and treatment of resection borders with ink, standard vertical section processing of the surgical excision specimen was used. Reporting of histopathological findings was performed by an experienced pathologist within 2 weeks after surgery. During assessment of the reference standard, the pathologist was masked to the results of clinical assessment and RCM but not to the results of a punch biopsy and patients’ clinical history. In line with standard care, the pathologist re-evaluated the results of a punch biopsy during the assessment of the excision specimen in cases of doubt. Besides, a biopsy scar could be recognized in excision specimen.

Analysis
We recorded the following characteristics of participants and tumours at baseline and summarized them for each treatment group with descriptive statistics: age, gender, skin type, previous BCC, study site, immune status, tumour diameter and tumour localization. Rippey’s classification was used for classifying BCC subtypes. A distinction was made between superficial, nodular and aggressive (micronodular, infiltrating and basosquamous) growth patterns. In the case of mixed-type diagnosis, defined as two or more single growth patterns, the most aggressive component was used for analysis. Diagnoses of BCC and subtype by RCM vs. punch biopsy were separately compared to surgical excision for all tumours. The primary outcome was the agreement between the index tests (RCM vs. punch biopsy) and reference standard (excision specimen) in correctly diagnosing BCC. The secondary outcome was the agreement between the index tests and reference standard in correctly identifying the most aggressive BCC subtypes. We excluded from the analyses cases in which RCM or a punch biopsy was indeterminate and cases in which subsequent surgical excision was not performed. Reasons for not performing surgical excision were recorded.

The number of true and false positives as well as true and false negatives was recorded. We established the sensitivity, specificity, positive and negative likelihood ratios, and predictive values for diagnosing BCC. For BCC subtyping, concordant results were calculated as the proportion of tumours with the corresponding subtype diagnosis in RCM or punch biopsy compared to excision specimen. The statistical analysis was performed using SPSS version 21.0 (SPSS Inc., Chicago, IL, USA).

Results

Participants
A total of 50 patients were randomized to RCM one-stop-shop (48 received index test and treatment) and 50 to standard care (47 received index test and treatment) (Fig. 1). Five patients were excluded, two patients in the RCM one-stop-shop group who did not receive imaging and three patients in the standard care group who did not receive surgical treatment (Fig. 1).

Baseline characteristics are shown in Table 1. In the RCM on-stop-shop group, 40 BCCs were confirmed by surgical excision specimen compared to 33 BCCs in the standard care group. Most of the BCC had a superficial or nodular subtype. All patients in the RCM one-stop-shop group received surgical treatment directly after RCM at the same initial outpatient visit. The average time between the initial visit and surgical treatment in the standard care group was almost 10 weeks (66 days).

Test results
The RCM experts evaluated the images that were acquired at the Department of Dermatology, Academic Medical Centre, University of Amsterdam (coordinating tertiary hospital). CL evaluated 32/36 cases, and MU evaluated 36/36 cases.

RCM vs. punch biopsy for diagnosing BCC Table 2 shows the agreement between specimens in correctly diagnosing BCC. Sensitivity to detect BCC was similar for RCM and punch biopsy (100% [90.75–100] vs. 93.94% [79.77–99.26]), but a punch biopsy was more specific than RCM (79% [49.20–95.34] vs. 38% [8.52–75.51]).

The RCM expert evaluation for diagnosing BCC was the same for both readers with a sensitivity of 100% [85.75–100] and a specificity of 75% [34.91–96.81].

RCM vs. punch biopsy for subtyping BCC Table 3 shows the agreement between RCM vs. a punch biopsy compared to excision in correctly identifying the most aggressive BCC subtypes. The overall agreement was 68% for RCM (26/38 concordant RCM cases) vs. 77% for punch biopsy (24/31 discordant punch biopsy cases). The initial RCM assessment during the trial period led to overstaging of BCC subtype in 18% (7/38) vs. 10% (3/31) by punch biopsy. Understaging of BCC subtype was seen in 13% of both RCM and a punch biopsy (5/38 vs. 4/31).

The agreement between BCC subtype of the RCM experts and excision specimen ranged from 50% (12/24 discordant RCM cases diagnosed by CL) to 85% (23/27 discordant RCM cases diagnosed by MU) (Table 3). Overstaging of BCC subtype by RCM expert teleconsultation assessment after the trial period ranged from 11% (3/27) to 50% (12/24). Understaging of BCC subtype by RCM experts ranged from 0% to 4% (1/27).

There were no adverse events after performing RCM or punch biopsies. Adverse reactions after performing surgical excision
213 participants assessed for eligibility

Excluded, n = 113
- 68 lesions on high-risk localisation
- 14 declined to participate
- 13 treated or recurrent lesions
- 10 lesions not suitable for confocal imaging
- 3 lesions larger than 20mm
- 2 technical malfunction confocal imaging device
- 2 planned holidays on short term
- 1 was not able to understand study procedure

100 randomized

50 assigned to standard of care

50 underwent a punch biopsy (index test 2)

36 tested positive for BCC

2 did not receive surgery*
  - 1 due to leiomyosarcoma (protocol deviation)
  - 1 had received PDT treatment by mistake (protocol deviation)

34 received surgical excision (reference standard)

14 tested negative for BCC

2 tested in inconclusive

50 assigned to one-stop-shop

48 underwent RCM imaging (index test 1)

43 tested positive for BCC

1 did not receive surgery*
  - due to actinic keratosis (protocol deviation)

43 received surgical excision (reference standard)

3 tested negative for BCC

2 received surgical excision (reference standard)

11 tested negative for BCC

2 tested positive for BCC

38 tested positive for BCC

5 tested negative for BCC***

3 tested negative for BCC

2 tested positive for BCC***

31 tested positive for BCC

3 tested negative for BCC**

13 tested negative for BCC

2 tested positive for BCC

3 tested positive for BCC

2 tested positive for BCC****

*Three patient in the standard care group did not receive subsequent surgical excision after the punch biopsy. In one patient, the protocol was violated after histological assessment of punch biopsy specimen showed actinic keratosis with no visible signs of the biopsied lesion on the day of surgery. Another patient with a histologically confirmed superficial BCC was mistakenly treated with photodynamic therapy instead of surgery. The last patient with a histologically confirmed BCC developed a large leiomyosarcoma on the same localization. Surgical excision of the BCC was cancelled, and the patient was referred to an oncologic surgeon to treat the leiomyosarcoma. **In the standard care group, a punch biopsy identified three lesions as BCC while surgical excision specimen did not show (residual) histological signs of BCC. ***RCM incorrectly identified five lesions as BCC while surgical excision specimen diagnosed two non-malignant lesions, one actinic keratosis, one Bowen's disease and one squamous cell carcinoma. ****In the RCM one-stop-shop group, two histology proven BCC (excision specimen) cases were tested as inconclusive, one ulcerating lesion and one lesion with a superficial crust.

Figure 1 Flow chart. BCC, basal cell carcinoma; PDT, photodynamic therapy. Two patients in the RCM one-stop-shop group did not begin diagnosis and treatment. One refused directly after randomization, and the other one could not participate due to technical malfunction of the confocal imaging device.
including four patients of the RCM one-stop-shop group with postoperative wound infections. In all cases, the infection was successfully treated with oral antibiotics, without the need of hospitalization. One patient of the standard care group using anticoagulant medication developed an excessive postoperative bleeding requiring hospitalization for 3 days. She fully recovered. This was reported as the only serious adverse event.

**Discussion**

Our findings show that for experienced users, RCM can have a similar diagnostic accuracy to diagnose and subtype clinically suspected BCC compared to a punch biopsy.

This is the first study that prospectively compared RCM with a punch biopsy for diagnosing and subtyping BCC. Previous RCM studies for diagnosing BCC showed varying high sensitivity and specificity values ranging from 85% to 97% and from 89% to 99%, respectively. However as reported by Que et al., most of these studies involved RCM experts with prior experience in RCM interpretation. Our results confirm the high sensitivity of RCM for diagnosing BCC (100%), but the specificity ranged from 38% (RCM users with <1 year experience) to 75% (RCM experts with more than 10 years of experience). We caution that the range of specificities for confirming BCC diagnosis is large for both RCM (DK/YE 38% [8.52–75.51] vs. CL/MU 75% [34.91–96.81]) and a punch biopsy (79% [49.20–95.34]). A lower RCM specificity was previously reported by Rao et al. that studied RCM users with varying levels of experience. Furthermore, Farnetani et al. also recently emphasized on the importance of the RCM learning curve and confirmed that diagnostic accuracy of RCM increases with experience.

The agreement between histological subtype on a punch biopsy and surgical excision specimen in our study was 77% (24/31 concordant cases). This seems consistent with previous studies. Interestingly, we found that RCM proved to be almost as reliable for accurately subtyping BCC (68%, 26/38 concordant cases). However, we also found a large difference in subtyping BCC between RCM experts. This is an important finding that highlights the need for further training, guidelines and protocols for subtyping BCC using RCM.

With the growing number of patients suffering from BCC, new management strategies are needed. Non-invasive skin imaging could play a crucial role in improving BCC health care for both patients and clinicians. Previous diagnostic RCM studies have primarily focused on test accuracy (sensitivity and specificity for diagnosing BCC). However, other aspects such as time between diagnosis and treatment, direct health effects of testing, costs of testing and patients’ emotional and behavioural responses to testing should also be taken into consideration.

In our proposed RCM one-stop-shop, we have assessed the efficacy of such a test-treatment pathway. The main advantages of using RCM include an immediate diagnosis and treatment for patients suffering from BCC opposed to painful skin biopsies with a doctor delay in the diagnostic process. Moreover in selected cases of superficial BCC, patients could benefit from a totally non-invasive disease management.

Study strengths include adherence to the STARD guidelines. Furthermore, we prevented sampling error using final surgical excision specimen as our reference standard instead of a punch biopsy. We also prevented heterogeneity of our results using predefined RCM criteria and using the same VivaScope 1500 device at both participating centres. Although it may not be in line with daily practice to surgically treat superficial BCC, it was important in our study to histologically confirm all types of BCC and to prevent selection bias for specific BCC subtypes.

| Table 1 Tumour and patient characteristics separated by treatment group |
|-----------------------------------------------|
| **One-stop-shop** (n = 50) | **Standard of care** (n = 50) |
| **Age (years)** | 64 (39–88) | 68 (41–92) |
| **Sex** | | |
| Men | 31 (62%) | 25 (50%) |
| Women | 19 (38%) | 25 (50%) |
| **Fitzpatrick skin type** | | |
| I | 8 (16%) | 4 (8%) |
| II | 32 (64%) | 43 (86%) |
| III | 10 (20%) | 3 (6%) |
| **BCC in medical history** | | |
| Yes | 34 (68%) | 37 (74%) |
| No | 15 (30%) | 13 (26%) |
| **Study site** | | |
| Academic Medical Centre | 37 (74%) | 38 (76%) |
| Netherlands Cancer Institute | 13 (26%) | 12 (24%) |
| **Immunocompromised** | | |
| Yes | 4 (8%) | 4 (8%) |
| No | 46 (92%) | 46 (92%) |
| **Tumour diameter (mm)** | 8 (3–15) | 8 (3–20) |
| **Tumour location** | | |
| Head/neck | 9 (18%) | 12 (24%) |
| Trunk | 32 (64%) | 30 (60%) |
| Arm | 4 (8%) | 7 (14%) |
| Leg | 5 (10%) | 1 (2%) |
| **Number of BCC** | 40 (80%) | 33 (66%) |
| **BCC subtype distribution** | | |
| Superficial BCC | 17 (43%) | 14 (42%) |
| Nodular BCC | 17 (43%) | 17 (52%) |
| Aggressive BCC | 6 (14%) | 2 (6%) |

1 Patients who were taking immunosuppressive drugs such as oral steroids, methotrexate, ciclosporin for suppression of immunological disorder, or to prevent transplant rejection.

2 This number represents the histologically confirmed basal cell carcinoma based on surgical excision specimen. Basal cell carcinoma subtype distribution according to the most aggressive subtype found at histology of surgical excision.

Continuous variables are expressed as mean (range) and categorical variables as n (%).

BCC, basal cell carcinoma.
Limitations of our study include the limited sample size of aggressive BCC. Our randomized controlled trial was primarily designed and powered to assess non-inferiority of a RCM one-stop-shop in terms of tumour-free margins after surgical treatment of BCC compared to standard care. Another important limitation that needs to be considered when interpreting the results is the potential bias in favour of the punch biopsy diagnosis. Although in line with current practice, the pathologists were not blinded to the results of the punch biopsy during assessment of surgical excision specimen in the standard care group. Lastly, BCCs on high-risk areas of the face were excluded due to technical limitations of the VivaScope 1500/C226 device. This needs to be considered in terms of external validity. Nonetheless the potential value of RCM remains very high in the excluded patient population since the introduction of the VivaScope 3000/C226 flexible handheld version (VivaScope 3000/C226; CaliberID; MAVIG GmbH), that permits imaging of the more concave and convex high-risk facial areas.

Based on our findings, we believe that RCM could potentially replace a punch biopsy for diagnosing and subtyping selected BCC cases. Yet prior to doing so, it is mandatory to wait for the results of future and ongoing larger prospective clinical trials. In addition, a first report on a combined RCM/OCT skin modality for ex vivo BCC detection has been published. This approach could potentially be of significant interest for diagnosing and subtyping BCC in clinical practice as it combines the detailed features of RCM with the in-depth advantages of OCT.

Finally, we underline that both routine histology as non-invasive skin imaging modalities such as RCM and OCT remain morphology based and thus subject to interpretation bias.

Table 2 Diagnostic performance of RCM vs. a punch biopsy in diagnosing BCC compared to surgical excision

|                          | RCM (DK/YE) trial period | Surgical excision | Total |
|--------------------------|--------------------------|-------------------|-------|
|                          | BCC                      | No BCC            |       |
| BCC                      | 38                       | 5                 | 43    |
| No BCC                   | 0                        | 3                 | 3     |
| Total                    | 38                       | 8                 | 46    |

|                          | Punch biopsy trial period | Surgical excision | Total |
|--------------------------|--------------------------|-------------------|-------|
|                          | BCC                      | No BCC            |       |
| BCC                      | 31                       | 3                 | 34    |
| No BCC                   | 2                        | 11                | 13    |
| Total                    | 33                       | 14                | 47    |

|                          | RCM expert (MU) after trial period | Surgical excision | Total |
|--------------------------|------------------------------------|-------------------|-------|
|                          | BCC                                | No BCC            |       |
| BCC                      | 27                                 | 2                 | 29    |
| No BCC                   | 0                                  | 6                 | 6     |
| Total                    | 27                                 | 8                 | 35    |

|                          | RCM expert (CL) after trial period | Surgical excision | Total |
|--------------------------|------------------------------------|-------------------|-------|
|                          | BCC                                | No BCC            |       |
| BCC                      | 24                                 | 2                 | 26    |
| No BCC                   | 0                                  | 6                 | 6     |
| Total                    | 24                                 | 8                 | 32    |

| BCC vs. no BCC | Sensitivity % (n [95% CI]) | Specificity % (n [95% CI]) | Positive LR [95% CI] | Negative LR [95% CI] | PPV [95% CI] | NPV [95% CI] |
|----------------|-----------------------------|-----------------------------|----------------------|----------------------|-------------|-------------|
| RCM (DK/YE) trial period (n = 46) | 100% (38/38) [90.75–100] | 37.50% (3/8) [8.52–75.51] | 1.60 [0.94–2.74] | 0 | 88.37% [81.63–92.86] | 100% |
| Punch biopsy trial period (n = 47) | 93.94% (31/33) [79.77–99.26] | 78.57% (11/14) [49.20–95.34] | 4.38 [1.60–12.00] | 0.08 [0.02–0.30] | 91.18% [76.32–98.14] | 84.62% [54.55–98.08] |
| RCM expert (MU) after trial period (n = 35) | 100% (26/26) [87.23–100] | 75% (6/8) [34.91–96.81] | 4.00 [1.20–13.82] | 0 | 93.10% [80.26–97.82] | 100% |
| RCM expert (CL) after trial period (n = 32) | 100% (24/24) [85.75–100] | 75% (6/8) [34.91–96.81] | 4.00 [1.20–13.82] | 0 | 92.31% [78.32–97.55] | 100% |

Bold numbers indicate concordant cases. Values in brackets are 95% confidence intervals.

BCC, basal cell carcinoma; LR, likelihood ratio; NPV, negative predictive value; PPV, positive predictive value; RCM, reflectance confocal microscopy imaging.
Table 3 Diagnostic performance of RCM vs. a punch biopsy in subtyping BCC compared to surgical excision

| Diagnostic Method | Surgical excision | Total |
|-------------------|-------------------|-------|
| **RCM (DK/YE) trial period** |                |       |
| sBCC              | 14                | 17    |
| nBCC              | 1                 | 2     |
| aBCC              | 2                 | 4     |
| **Total**         | 17                | 38    |
| **Punch biopsy trial period** |                |       |
| sBCC              | 13                | 17    |
| nBCC              | 0                 | 9     |
| aBCC              | 0                 | 2     |
| **Total**         | 13                | 31    |
| **RCM expert (MU) after trial period** |                |       |
| sBCC              | 13                | 14    |
| nBCC              | 1                 | 10    |
| aBCC              | 1                 | 3     |
| **Total**         | 15                | 26    |
| **RCM expert (CL) after trial period** |                |       |
| sBCC              | 3                 | 18    |
| nBCC              | 1                 | 10    |
| aBCC              | 9                 | 6     |
| **Total**         | 13                | 24    |

Bold numbers indicate concordant cases.

sBCC, superficial basal cell carcinoma; nBCC, nodular basal cell carcinoma; aBCC, aggressive basal cell carcinoma; BCC, basal cell carcinoma.

In conclusion, RCM and punch biopsy have comparable diagnostic accuracy to diagnose and subtype BCC depending on RCM experience. Although experienced RCM users could accurately diagnose BCC at a distance, we found an important difference in subtyping BCC. Future RCM studies need to focus on diagnostic accuracy, reliability and specific criteria to improve BCC subtype differentiation.

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