New host associations and a novel species for the gall-inducing acacia rust genus *Ravenelia* in South Africa

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Abstract

Trees in the genus *Vachellia* (previously *Acacia*) are commonly infected by the gall-inducing rusts *Ravenelia macowaniana* and *R. evansii*. Rust galls bearing aecial infections and relating uredinial and telial infections on the leaves of nine *Vachellia* species not previously recorded to be infected by *Ravenelia* spp. have recently been collected in South Africa. The rust fungi causing these infections were characterised using molecular phylogenetic analyses of DNA sequence data of the LSU and ITS rDNA regions as well as morphological examinations. The host range of *R. macowaniana* and *R. evansii* was thus re-assessed and extended from four to nine species and from one to three species, respectively. Application of Principal Component Analyses (PCA) of telial morphological characters provided evidence of an effect of the host species on the teliospore morphology in *R. evansii*, but only minor effects in *R. macowaniana*. A novel gall-inducing *Ravenelia* sp. closely related to *R. macowaniana*, was found on *Vachellia xanthophloea* and it is described here as *R. xanthophloeae*.

Keywords

*Ravenelia xanthophloeae* sp. nov., *Vachellia xanthophloea*, novel host record, aecial galls, teliospore morphology, intraspecific variability, Principal Component Analysis

Introduction

Trees in the genus *Vachellia* (formerly *Acacia* subg. *Acacia*) and referred to here as acacias make up one of the most prominent floral elements of the Southern African...
landscape. Acacias can be found in all South African biomes. Here they play important ecological roles by providing food for insects, birds and game, as well as improving soil fertility through nitrogen fixation by their associated rhizobia (Ross 1979, Coe and Coe 1987, Coates Palgrave 2005, Smit 2008, Grellier et al. 2012).

In South Africa, acacias are commonly infected by rust fungi (Pucciniales) of the genus Ravenelia (Doidge 1950). Ravenelia includes more than 200 described species and is amongst the most species-rich genera of rust fungi (Cummins and Hiratsuka 2003). These fungi are obligate parasites of various genera of the legumes (i.e. in sub-families Mimosoideae, Faboideae, Caesalpinioideae) and they are globally distributed in the tropics and sub-tropics (Dietel 1906, Cummins and Hiratsuka 2003). While the aecial stage of several macrocyclic species is known to cause hypertrophied tissues such as galls and witches brooms within host organs (Dietel 1894, Hernandez and Hennen 2003), the multicellular teliospores of Ravenelia are amongst the most complex spore forms found in the rusts.

In South Africa, 20 species of Ravenelia have been described, the majority of which infect trees of the acacia genera Senegalia and Vachellia (Doidge 1939, 1950, van Reenen 1995). Four of these including R. natalensis Syd., P. Syd. & Pole-Evans, R. deformans (Maublanc) Dietel, R. macowaniana Pazschke and R. evansii Syd. & P. Syd. induce galls on their hosts. Based on the number of deposited specimens in the National Collection of Fungi, South Africa (PREM) and our own field observations, the macrocyclic and gall-inducing R. macowaniana and R. evansii are likely the most abundant Ravenelia species in South Africa. Ravenelia macowaniana has been reported from V. karroo only, which in turn is the most frequently occurring acacia in this region. Ravenelia evansii has been reported from four acacia species including V. gerrardii (Benth.) P.J.H. Hurter, V. rehmanniana (Schinz) Kyal. & Boatwr. V. robusta ssp. robusta (Burch.) Kyal. & Boatwr. and V. sieberiana var. woodii (Burtt Davy) Kyal. and Boatwr. (Doidge 1939, 1950). In contrast, R. deformans and R. natalensis, both reported from V. karroo (Hayne) Banfi and Galasso, have only rarely been collected (Doidge 1939, Farr and Rossman 2017).

During several field surveys focused on re-assessing the diversity of Ravenelia species in South Africa, we sampled rust infections associated with galls on eight Vachellia species. The aim of this study was to identify the collected rust specimens using morphological and phylogenetic analyses. In addition, new host associations and a new species were reported and the influence of the host on teliospore characters was analysed.

**Material and methods**

**Specimens examined**

Infected V. borleae, V. davyi, V. exuvialis, V. hebeclada, V. natalitia, V. permixta, V. swazica and V. xanthophloea trees were sampled during several field surveys in South Africa between 2004 and 2015. Leaves bearing uredinal and telial rust sori and short branch-
es having aecial galls were collected and subsequently dried between paper sheets in a plant press. In total, 49 specimens were studied based on morphology and 31 of these could be used for phylogenetic analyses based on DNA-sequence data. Ten of the 49 specimens were either type or voucher specimens collected in the late 19th and early 20th century and used by Doidge for her studies of the southern African Ravenelia spp. (Doidge 1927, 1939). These herbarium specimens were used only for morphological comparisons with the newly collected material (Table 1). Three specimens were deposited in KR, all others in PREM.

**DNA extraction and PCR**

Spores from individual sori were collected separately using sterile insect needles. Genomic DNA extractions were made using the INNUPrep Plant DNA Kit (Analytik Jena, Germany) following the manufacturer’s protocols with the following modifications: Spores were crushed using a Retsch mixer mill MM2000 (Retsch, Haan, Germany) by shaking them together with 2 steel beads of 2.5 mm diameter in a 2.0 ml Eppendorf tube. This process was repeated in three consecutive cycles. In the first step, the closed tubes were cooled in liquid nitrogen and immediately shaken for 2 min at 100 Hz. Thereafter 10–40 μl of lysis buffer was added to the tube to loosen spore remnants from the inner side of the Eppendorf tube lid using a vortex mixer followed by a centrifugation step. Samples were again cooled in liquid nitrogen and shaken for an additional 2 min at 100 Hz followed by centrifugation for 1 min at 6000 rcf. The last two steps were repeated once.

For PCR of the ribosomal nrITS and LSU rDNA gene regions, the Taq-DNA-Polymerase Mix (PeqLab, Erlangen, Germany) was used with the primers ITS1F (Gardes and Bruns 1993) and RustITS1F (Toome and Aime 2014), respectively and ITS4BR (Vialle et al. 2009). PCR of the LSU rDNA region was performed using the primer pairs LR0R and LR6 (Vilgalys and Heester 1990). Two additional primers (5.8SrustF: 5’-CGA TGA AGA ACA CAG TGA AAT GTG; D1D2RustR: 5’-CTY TGC TAT CCT GAG GGA) were designed with improved specificity for Ravenelia and that reduced amplification of ascomycetous non-target organisms. The thermal cycling conditions for primers ITS1F/ITS4BR and RustITS1F/ITS4BR were as follows: 2 min at 96 °C followed by 40 cycles of 20 sec at 96 °C, 40 sec at 50 °C and 50 sec at 72 °C, final extension was for 5 min at 72 °C; for primers LR0R/LR6: 3 min at 96 °C followed by 40 cycles of 30 sec at 95 °C, 40 sec at 49 °C and 1 min at 72 °C, final extension was for 7 min at 72 °C; for primers 5.8SrustF/D1D2rustR: 3 min at 96 °C followed by 40 cycles of 30 sec at 96 °C, 45 sec at 54 °C and 1 min 20 sec at 72 °C, final extension was for 7 min at 72 °C.

PCR products were purified using Sephadex G-50 columns (Sigma-Aldrich, Steinheim, Germany). Where PCR products showed only weak bands on agarose gels, purification was undertaken using the Zymo Research DNA Clean & Concentrator™-5 Kit (Zymo Research GmbH, Freiburg, Germany) following the manufacturer’s pro-
Table 1. List of specimens included in the present study, including host information, collection data and GenBank accession numbers of rDNA sequences.

| Voucher | Host | Species name | Origin | Date | Collector | GenBank accession-Nos. |
| --- | --- | --- | --- | --- | --- | --- |
| PREM61208 | Ravenelia evansii | Vachellia evansii | South Africa, North-West Province, Groen Marico, River Still Guest Farm | 15 Apr 2009 | W. Maier | MG945960, MG945992 |
| PREM21209 | | | | 18 Apr 2010 | M. Ebinghaus | MG945959, MG945991 |
| PREM71105 | Vachellia robusta, Vachellia termitina | | | 6 Jul 1913 | I. B. Pole Evans | – |
| PREM61105 | Vachellia robusta, Vachellia termitina | | | 3 Jul 1913 | M. Ebinghaus | MG945956, MG945990 |
| PREM61224 | Vachellia sieberiana var. woodii | | South Africa, Mpumalanga, R40 north of Nelspruit | 22 June 2005 | W. Maier | – |
| PREM61005 | Vachellia sieberiana var. woodii | | | 11 Apr 2013 | M. Ebinghaus | MG945966, MG945998 |
| PREM61222 | Vachellia hebeclada | | South Africa, North-West Province, Leeuwfontein Farm | 30 Dec 2006 | A. E. van Wyk | MG945969, MG945997 |
| PREM61228 | Vachellia exuvialis | | South Africa, Mpumalanga; Marloth Park; 25°20'44.4"S; 31°46'26.1"E | 9 Apr 2013 | M. Ebinghaus | MG945964, MG945996 |
| PREM61846 | Vachellia luederitzii var. retinens | | South Africa, North-West Province, Jonimo; 27°30'57.5"S; 31°11'52.0"E | 17 Feb 2015 | M. Ebinghaus | MG945985, MG945995 |
| PREM61876 | Vachellia meridionalis | | South Africa, Mpumalanga, Belfast; 24°56'08.7"S; 31°47'52.4"E | 17 Feb 2015 | M. Ebinghaus | MG945986, MG945994 |
| Voucher | Species name | Host | Origin | Date       | Collector | GenBank accession-No. |
|---------|-------------|------|--------|------------|-----------|----------------------|
| ME384   | Vachellia borleae | South Africa, KwaZulu-Natal; 20 km north of Empangeni; 28°41'30.1"S; 31°43'16.9"E | 9 Feb 2015 | M. Ebinghaus | MG945971 MG946003 |
| PREM61869 | Vachellia borleae | South Africa, Mpumalanga; Masibekela; 25°51'36.2"S; 31°49'51.8"E | 16 Feb 2015 | M. Ebinghaus | MG945970 MG946002 |
| PREM61222 | *Ravenelia macowaniana* | Vachellia karroo | South Africa, Limpopo, Sekhukhune Land, Winterveld Mine | 23 June 2005 | W. Maier | MG945975 MG946007 |
| PREM61221 | Vachellia karroo | South Africa, North-West Province, Hartbeespoort Dam | June / Jul 2005 | W. Maier | MG945973 MG946004 |
| PREM61210 | Vachellia karroo | South Africa, Eastern Cape, Haga Haga | Dec 2005 | W. Maier | MG945972 MG946004 |
| PREM61220 | *Ravenelia macowaniana* | South Africa, unknown | 15 May 2006 | W. Maier | – – |
| KR-M-43406 | Vachellia permixta | South Africa, Western Cape, Worcester | 20 Dec 2004 | M.J. Wingfield | MG945974 MG946006 |
| KR-M-43657 | Vachellia permixta | South Africa, Limpopo, Mokopane; 24°08'52.4"S; 29°02'21.9"E | 23 Feb 2015 | M. Ebinghaus | MG945982 MG946014 |
| PREM61862 | Vachellia natalitiae | South Africa, Limpopo, Steelport; 24°41'32.3"S; 30°12'32.3"E | 19 Feb 2015 | M. Ebinghaus | MG945980 MG946012 |
| PREM61218 | Vachellia natalitiae | South Africa, Mpumalanga, Nelspruit | 10 Jan 2005 | W. Maier | MG945979 MG946011 |
| PREM61219 | Vachellia natalitiae | South Africa, Mpumalanga, Nelspruit | 10 Jan 2005 | W. Maier | MG945977 MG946009 |
| PREM61888 | Vachellia natalitiae | South Africa, Mpumalanga, Nelspruit | 21 June 2005 | W. Maier | – – |
| PREM 61216 | South Africa, Eastern Cape, Port St. John | 28 Dec 2005 | W. Maier | MG945978 MG946010 |
| PREM61226 | Vachellia natalitiae | South Africa, Mpumalanga, South of Nelspruit | 16 Mar 2010 | M. Ebinghaus | – – |
| PREM61214 | Vachellia xanthophloea | South Africa, Mpumalanga, East of Barberton | 2 June 2012 | M. Ebinghaus | MG945981 MG946013 |
| PREM61215 | Vachellia xanthophloea | South Africa, Mpumalanga, Barberton; 25°46'52.5"S; 30°03'10.7"E | 3 Jul 2012 | M. Ebinghaus | MG945985 MG946017 |
| PREM61213 | South Africa, KwaZulu-Natal, Mount Moreland; 29°38'21.6"S; 30°05'27.3"E | 16 June 2012 | M. Ebinghaus | MG945983 MG946015 |
| PREM61213† | South Africa, KwaZulu-Natal, Mount Moreland; 29°38'21.6"S; 30°05'27.3"E | 16 June 2012 | M. Ebinghaus | MG945986 – |
| PREM61000 | South Africa, Mpumalanga, Komatiporto | 9 Apr 2013 | M. Ebinghaus | MG945984 MG946016 |
| PREM1935† | Vachellia karroo | South Africa, KwaZulu-Natal, Winkelspruit | 29 Nov 1911 | I. B. Pole Evans | – – |
| PREM2514† | Vachellia karroo | South Africa, KwaZulu-Natal, Winkelspruit | 6 Jul 1912 | E. M. Doidge | – – |
| PREM2575† | Calpurnia sylvatica | South Africa, KwaZulu-Natal, Mulden | 1 May 1912 | P. MacOwan | – – |
| PREM10698 | not known | not known | not known | P. MacOwan | – – |
| PREM20727† | South Africa, Western Cape, Somerset East | 1875 | P. MacOwan | – – |

1Specimens were only morphologically investigated.
2Specimen representing the aecial infection state.
protocol. DNA sequencing was carried out in both directions using the same primers as those used for PCR on a 3130XL Genetic Analyzer (Applied Biosystems) at the sequencing service of the Faculty of Chemistry and Biochemistry of the Ruhr University Bochum, Germany.

Phylogenetic analyses

Sequences were screened against the NCBI GenBank using the BLASTn algorithm (Altschul et al. 1990) to exclude erroneously amplified contaminants from further processing. Forward and reverse strands of the rust sequences were assembled using Sequencher 5.0 software (Gene Codes Corp., Ann Arbor, MI, USA) and, where necessary, manually edited. In total, 32 sequences were used to construct an alignment of the nrITS and LSU rDNA sequence data, respectively, using MAFFT v6.832b (Katoh and Standley 2014) applying the L-INS-i strategy. Maximum likelihood analyses were conducted in RAxMLGUI v.1.3 (Silvestro and Michalak 2012) using RAxML 8.0.26 (Stamatakis 2014) using the general time reversible model of nucleotide substitution (Lanave et al. 1984) with gamma distributed substitution rates. The analyses were run with a rapid bootstrap analysis using 1000 bootstrap replicates. The analyses were first conducted for each dataset separately and topological congruence was checked visually. As no conflict of supported phylogenetic groupings was observed, the final phylogeny was inferred by combining both datasets of the nrITS and LSU rDNA sequences applying the same methodology as for individual datasets.

Parsimony network analyses were performed using TCS v1.21 (Clement et al. 2000) and the same sequence alignments that were used for the phylogenetic analyses. Gaps were deleted from calculations and the default connection limit of 95% was used.

Light- and electron microscopic investigations

The spores of the dried herbarium specimens (Table 1) were scraped from leaf surfaces and mounted in lactophenol on microscope slides. A minimum of ten teliospores and 30 urediniospores per specimen were examined. Minimum, maximum and mean values are provided in Table 2. The specimens PREM2211 (R. evansii), PREM2403 (R. evansii), PREM2539 (R. evansii), PREM6807 (R. evansii), PREM7105 (R. evansii), PREM1935 (R. natalensis), PREM2514 (R. natalensis), PREM2375 (R. glabra), PREM10698 (R. glabra) and PREM20727 (R. glabra) were examined at the facilities of the ARC-Plant Protection Institute (ARC-PPRI), Roodeplaat, South Africa using a Leica Dialux 22 EB microscope and a ColorView III CCD colour camera. Measurements were made using analySIS LS software (LS Research Software GmbH, Germany). The remaining specimens were studied at the Ruhr University Bochum, Germany, using a Zeiss Axioplan light microscope. Morphological characteristics were measured
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Table 2. Measurements of morphological characters of Ravenelia evansii, R. macowaniana and R. xanthophloeae sp. nov., separately performed for each host species. All size measurements are given in μm. †Data taken from original descriptions by Doidge (1927).

| Ravenelia on host species | Teliospore diameter | Probasidial cell length | Probasidial cell width | Epispore thickness | Ornamentation length | Cell numbers in diam. |
|---------------------------|---------------------|-------------------------|------------------------|--------------------|----------------------|----------------------|
| **R. evansii**            |                     |                         |                        |                    |                      |                      |
| V. robusta†               | 50–80               | 25–30                   | 17–23                  | 4–6                | 4–6                  | 4–6                  |
| V. borleae                | (52)72–83(92)       | (21)23–28(40)           | (16)22–26(34)          | (2.5)3–4.5(6.0)    | (3)3.5–4.5(6)        | 3–6                  |
| V. davyi                  | (83)95–105(115)     | (19)25–30(34)           | (12)19–24(32)          | (2.5)4–5(6.5)      | (2.0)3.5–5(7)        | 5–7                  |
| V. exuvialis              | (47)70–85(101)      | (19)23–28(33)           | (15)21–26(30)          | (2.5)3.5–4.5(6)    | (3)4.5–6(7.5)        | 3–7                  |
| V. luedercizii            | (79)94–100(115)     | (18)23–30(36)           | (15)19–25(32)          | (2.5)3–5(7)        | (1.5)3–5(6)          | 5–7                  |
| V. robusta                | (81)90–110(118)     | (20)22–29(35)           | (14)19–25(33)          | (3)3.5–5(6.5)      | (1.5)3.5–6(8)        | 5–8                  |
| V. sieberiana             | (63)85–97(112)      | (19)23–29(35)           | (15)21–24(33)          | (2.5)3.5–4.5(6)    | (2.5)4.5–6(8)        | 4–8                  |
| V. swazica                | (71)85–105(124)     | (16)22–29(39)           | (12)18–27(32)          | (2)3–5(7)          | (2.5)3.5–6(7)        | 5–8                  |
| **R. macowaniana**        |                     |                         |                        |                    |                      |                      |
| V. karroo†                | 60–130              | up to 45                | 18–28                  | not stated         | –                    | 4–7                  |
| V. karroo                 | (75)82–105(118)     | (21)23–31(35)           | (14)17–24(29)          | (2)3–5.5(6.5)      | –                    | 4–7                  |
| V. natalitia              | (50)80–105(114)     | (19)24–34(45)           | (14)19–26(34)          | (2)3–4.5(5.5)      | –                    | 3–6                  |
| **R. xanthophloeae**      |                     |                         |                        |                    |                      |                      |
| V. xanthophloeae          | (40)65–75(84)       | (19)22–28(40)           | (12)18–25(29)          | (2)3–4(6)          | (0.5)1–2(3)          | 3–6                  |

using Cell^D v. 3.1 imaging software (Olympus Soft Imaging Solutions GmbH, Germany) and Zen2 lite (Blue Edition) V. 2.0.0.0 (Carl Zeiss Microscopy, 2011, Jena, Germany). Photographs were obtained using a Color View microscope camera (Olympus Soft Imaging System, Germany). For detailed investigations of the spore-surface structures, scanning electron microscopy (SEM) was used. For this purpose, infected leaflets from the herbarium specimens were mounted on double-sided adhesive carbon tape on metal stubs and coated with gold in a sputter coater BAL-TEC SCD OSO (Capovani Brothers Inc, USA). Subsequently, the samples were examined using a ZEISS Sigma VP scanning electron microscope.

Principal Component Analyses (PCA)

For principal component analyses (PCA) the morphological data collected for all examined rust individuals were separated into sub-sets based on preliminary species assignments representing R. evansii, R. macowaniana and Ravenelia sp. (Groups A, B1 and B2). PCA for all subsets was conducted separately using the R-packages plyr and ggplot2 implemented in R (www.R-Project.org). Six characteristics of teliospores providing numeric data were defined and measured: teliospore diameter, probasidial cell length and probasidial cell width, number of cells in diameter, epispore thickness and ornamentation length. Mean values were calculated for the individual teliospore measurements, scaled and missing values were deleted from analyses.
Results

Phylogenetic Analyses

Sequence data from the nrITS and LSU rDNA gene regions were obtained for all 31 newly collected specimens. The alignment of the nrITS sequence dataset had a total length of 764 bp with 133 variable sites of which 131 positions were parsimony informative. The aligned sequences of the LSU rDNA dataset had a length of 922 bases and comprised 31 sequences with 45 variable sites and 40 parsimony informative positions. The combination of the nrITS and LSU rDNA datasets resulted in an alignment with a total length of 1686 nucleotides comprising 32 sequences. The sequence alignment and phylogenetic tree of the combined rDNA sequence data set was deposited at TreeBASE (http://purl.org/phylo; submission IDS22307).

Maximum likelihood analysis of the combined dataset resulted in a phylogenetic tree that consisted of three highly supported groups representing *R. evansii*, *R. macowaniana* and a novel *Ravenelia* species described below (Fig. 1A).

The phylogenetic group representing *R. evansii* consisted of 17 sequences. Three of these were obtained from *Vachellia sieberiana* var. *woodii* (PREM61223, PREM61228, PREM61881) and three from *V. robusta* ssp. *robusta* (KR-M-43649, PREM61208, PREM61209). These are tree species that had previously been reported as hosts of *R. evansii*. Rust specimens collected from the following five *Vachellia* species also clustered in this group: *V. borleae* (ME384 and PREM61869), *V. davyi* (PREM61005), *V. exuvialis* (PREM61868, PREM61876), *V. hebeclada* (PREM61227) and *V. swazica* (PREM61002, PREM61008, PREM61028). These are all newly reported hosts for *R. evansii*.

A second group included the sequences of eleven specimens, five of which originated from *V. karroo* and were identified as *R. macowaniana* (KR-M-43657, PREM61222, PREM61221, PREM61210, KR-M-43406). Five specimens were collected from *V. natalitia* (PREM61214, PREM61862, PREM61218, PREM61219, PREM61216) and one originating from *V. permixta* (PREM61875) also clustered in this group. The latter two hosts are newly reported for *R. macowaniana*.

A distinct clade, nested within the *R. macowaniana* group, was represented by three *Ravenelia* specimens that were isolated from *V. xanthophloea* (PREM61215, PREM61213, PREM61000) suggesting that it represents a novel taxon. For PREM61213, two identical sequences were obtained, one derived from aeciospores and one from teliospores.

The parsimony network analysis, based on the combined set of nrITS and LSU rDNA sequence data, separated three distinct groups each comprising the same specimens representing *R. evansii*, *R. macowaniana* and the novel *Ravenelia* species in our phylogenetic analysis, respectively (Fig. 1B). Network analysis relying on LSU alone could not separate *R. macowaniana* from the novel taxon, while separation of these two groups was observed based on nrITS alone (not shown). The *R.
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**Figure 1.** Phylogenetic reconstruction of *Ravenelia* species on different *Vachellia* hosts

**A** Maximum likelihood tree with 1000 bootstrap repeats based on combined nrITS and LSU rDNA sequence data. Bootstrap values below 75 are not shown. Three highly supported groups represent *R. evansii*, *R. macowaniana* and *R. xanthophloeae* sp. nov., respectively. Specimens that originated from formerly unreported host species are highlighted in bold.

**B** Parsimony network analysis based on the same dataset as in the ML-analysis. Each line represents one base substitution while small circles represent intermediate but missing sequences. Numbers next to lines indicate the positions of the substitutions in the alignment. Sequences in rectangular boxes were inferred as ancestral by this analysis.

evansii group consisted of six haplotypes of 17 sequences that differed by a maximum of two substitutions from the inferred ancestral sequences (PREM61005 and PREM61227). In *R. macowaniana*, sequence divergence was more pronounced and consisted of nine haplotypes in a total of eleven sequences. The highest rate of six substitutions was observed for specimen PREM61221 relative to the inferred ancestral sequence (KR-M-43657). Specimens collected on *V. xanthophloeae* had only one substitution.
Morphological analyses

**Ravenelia evansii**

The teliospore morphology of *R. evansii* specimens showed a considerable overall variability in all six investigated teliospore characteristics (Suppl. material 1: Fig. S1, Table 2). The voucher specimens PREM61869 (on *V. borleae*), PREM61876 and PREM61868 (both on *V. exuvialis*) had significantly smaller teliospores compared to the remaining specimens, but variation in this trait could also be observed within single host associations, e.g. within those from *V. davyi* and *V. robusta* (Suppl. material 1: Fig. S1).

The principal component analysis (PCA) of teliospore characteristics clustered several individuals derived from specific hosts into distinct groups (Fig. 2C, D). Individuals that originated from *V. borleae* and *V. exuvialis* clustered more closely and could be separated from those individuals that were collected from *V. davyi* (Fig. 2D). The separation of these individuals was supported by PC1 which could explain 37.6% of the overall variability. The traits ‘cells in diameter’ and ‘teliospore diameter’ were characteristics that corresponded best with this axis (Fig. 2C). These results indicated, that the latter two characters were the most variable traits to separate these individuals with different host associations. However, individuals from *V. luederitzii* var. *retinens*, *V. robusta* var. *robusta*, *V. sieberiana* var. *woodii* and *V. swazica* showed only weak separation, i.e. less clear patterns of morphological separation.

**Ravenelia macowaniana**

Specimens representing *R. macowaniana* were morphologically more homogeneous compared to *R. evansii*. Here, only spore characteristics such as ‘probasidial cell width’ and ‘epispore thickness’ were often significantly different between investigated specimens (Suppl. material 2: Fig. S2). There was little variation for the characters ‘teliospore diameter’, ‘probasidial cell length’ and the number of ‘cells in diameter’. These characters varied distinctly only between single specimens, e.g. PREM61226 originating from *V. natalitia* had significantly larger teliospores in comparison to specimen PREM61888 that was also collected from this host (Suppl. material 2: Fig. S2).

For the specimens of *R. macowaniana*, PC1 and PC2 could explain 36.2% and 22.4% of the similarity, respectively. However, unlike in *R. evansii*, single teliospore characteristics did not differ significantly in terms of host association (Fig. 2A).

**Ravenelia sp. nov.**

Due to similar teliospore characteristics, *R. macowaniana* was compared using PCA to individuals of the undescribed *Ravenelia* species collected on *V. xanthophloea* in order to characterise and, if possible, to contrast both morphologies. The PCA separated two groups that corresponded well with *R. macowaniana* and the novel *Ravenelia* species and
showed very little overlap in morphological characteristics (Fig. 2B). Separation between both species groups was mostly seen in PC1 that could explain 42.5% of the overall variability with the ‘ornamentation length’ corresponding best to this axis (Fig. 2B). Although less distinct in the PCA, mean values of teliospore characters also revealed that the characters ‘epispore thickness’, ‘probasidial cell length’ and especially the ‘teliospore diameter’ and ‘cells in diameter’ are valuable characters for the discrimination of both species (Fig. 3). All spore measurements, derived from the six defined spore characteristics that were used for PCA, are available as an excel-file in the Suppl. materials 3, 4: tables S1, S2: ‘PCA-table $R.\ evansii$’ and ‘PCA-table $R.\ macowaniana$', respectively.

**Taxonomy**

**Ravenelia xanthophloeae** Ebinghaus, W. Maier & Begerow, sp. nov.
Mycobank: MB824073
Figure 4A–H

**Type.** South Africa, KwaZulu-Natal, 29°38’21.6”S; 31°05’27.3”E, on leaves and gall-transformed inflorescences of *Vachellia xanthophloea* (Benth.) P.J.H. Hurter (Fabaceae:
Mimosoideae), 16 June 2012, M. Ebinghaus ME188, (holotype: PREM61213); South Africa, Mpumalanga, 25°46'52.5"S; 31°03'10.7"E, on leaves of Vachellia xanthophloea (Benth.) P.J.H. Hurter (Fabaceae: Mimosoideae), 3 July 2012, M. Ebinghaus ME174, (paratype: PREM61215); South Africa, Mpumalanga, 25°26'10.0"S; 31°57'48.6"E, on leaves of Vachellia xanthophloea (Benth.) P.J.H. Hurter (Fabaceae: Mimosoideae), 9 April 2013, M. Ebinghaus ME248, (paratype: PREM61000)

**Etymology.** The name refers to the host tree, Vachellia xanthophloea.

**Description.** Spermogonia not found. Aecia on rust-induced galls, which are formed instead of inflorescences. Aeciospores globose to sub-globose, often angular, yellowish-transparent in light microscopy, light brown when dry (19.0)21.0–24.0(28.5) × (14.5)18.0–20(21.5) μm, cell wall (1.0–)2.0(–3.0) μm thick, densely verrucose, germ pores numerous, scattered (Figure 4C). Peridia and peridial cells could not be described because only disintegrated aecia were present in the dried herbarium material.

Uredinia amphigenous on leaves, but mostly on the abaxial side of the leaflets, scattered or in small groups, minute, 0.1–0.2 mm, erumpent and surrounded by the torn epidermis; ellipsoidal to roundish, light-brown to blackish; paraphyses numerous, scattered within sorus; capitulate, thickened end ovoidal, about 19–20 × 11–13 μm, cell wall 2–3 μm, light-brown, smooth; urediniospores ovoidal to broadly ellipsoidal or sometimes subglobose, (18)23–26(38) × (13)16–20(25) μm, spore wall evenly 1.5–2.0 μm thick with densely echinulate aculei (Figure 4D), germ pores 5–6, subequatorial to equatorial (Figure 4E).
New host associations and a novel species for the genus *Ravenelia*

**Figure 4.** Infected host organs and spore images of *R. xanthophloeae* (A–H), *R. natalensis* (I), *R. evansii* (J) and *R. macowaniana* (K)  
A Infected individual of *V. xanthophloea*. Leaves were prematurely shed in comparison with uninfected trees  
B Telia on leaflets of *V. xanthophloea*  
C SEM of an aeciospore showing scattered germpores  
D SEM of an urediniospore  
E Urediniospores seen in LM  
F SEM view of a teliospore of *R. xanthophloeae*. The arrows indicate irregularly arranged verrucose ornamentations  
G Telium of *R. xanthophloeae* seen in SEM  
H LM view of a teliospore. The arrow indicates irregularly arranged verrucose ornamentations  
I Teliospores of *R. natalensis* with long pedicels  
J SEM picture of median section of a teliospore of *R. evansii*. Arrows indicate 2-celled probasidial cells  
K LM picture of teliospores of *R. macowaniana*.  
Scale bars: 1 mm (B), 4 μm (C), 2 μm (D), 20 μm (E), 20 μm (F–H, J–K), 40 μm (I).
Telia replacing the uredinia; teliospores often irregularly shaped from top view; single probasidial cells distinctly arched upwards (Figure 4F–H), orange-brown to pale brown, teliospore surface in general smooth but with single and irregularly arranged small verrucae, (Figure 4F–I); teliospores varying in size from (40)65–75(84) μm in diameter with mostly 4–5 probasidial cells in a cross-section, rarely 3 or 6 cells; central cells in two layers, 32–40 μm in lateral view; marginal cells in a single layer, (19)22–28(33) μm in lateral view and (12)18–25(29) μm from above; upper cellwall (2)3–4(6) μm thick; verrucose ornamentations (0.5)1–2(3) μm in height (Figure 4F, H), rarely with protuberances of up to 7 μm in height; cysts smooth and hyaline, of variable number but often appear in same number as marginal probasidial cells, swelling in water but only slightly in lactophenol, pedicels compound, not persisting.

Notes. In South Africa, *R. macowaniana*, *R. glabra* Kalchbr. & Cooke and *R. deformans* (Maublanc) Dietel are the only known species that exhibit two-layered probasidial cells and smooth teliospores. While the first character is shared by *R. xanthophloeae*, the teliospore surface bears small and irregularly arranged small warts clearly visible in SEM (Fig. 4F). However, these can easily be overlooked in light microscopy (Fig. 4H) and could potentially lead to misidentification. Specifically, *R. macowaniana* differs from *R. xanthophloeae* in the overall size of its teliospores (Table 2; Figure 4K) and the urediniospores have four equatorial germ pores whereas those of *R. xanthophloeae* have five to six equatorial germ pores. The teliospores of the microcyclic *R. glabra* Kalchbr. & Cooke are about twice the size (120–160 μm) of those of *R. xanthophloeae* and its oblong urediniospores are significantly larger (32–48 × 14–21 μm). This rust has also been reported only from *Calpurnia sylvatica* (Burch.) E. Mey (Fabaceae) (Doidge 1927). The demicyclic *R. deformans* (Maublanc) Dietel was synonymised with the neotropical *R. hieronymi* Speg. based on nearly identical morphology and congruent life cycle characteristics (Hernandez and Hennen 2003; Hennen et al. 2005) but conspecificity of these two rust fungi is doubtful as they infect distinct host species and occur each on different continents. However, both species produce aecia that induce malformations in young branches, which is a characteristic similar to the newly described *R. xanthophloeae*. With a size of 60–120 μm and 75–120 μm, respectively, the teliospores of *R. deformans* and *R. hieronymi* are, however, on average significantly larger and develop intermingled with the aecia (Doidge 1927), while *R. xanthophloeae* is macrocyclic and aecia, uredinia and telia are produced in spatially separated sori.

The teliospores of *R. xanthophloeae* may also be confused with those of *R. natalensis* Syd., P. Syd & Pole-Evans, but they are significantly smaller in size (30–50 μm diam.) and possess extraordinarily long and persistent pedicels (up to 110 μm; Sydow 1912, Fig. 4I). In *R. natalensis*, the aparaphysate uredinia and telia are confluent and cover large areas on the branches of the host (Sydow and Sydow 1912; own observations), while the specimens of *R. xanthophloeae* examined in this study have minute uredinia with numerous paraphyses and telia not exceeding 200 μm in diameter.

New host records. Morphological and molecular phylogenetic analyses based on nrITS and nrLSU data confirmed new host records for *R. macowaniana* and *R. evansii* that will be reported in the following section. An emended species description for *R. evansii* is also provided.
**Ravenelia macowaniana** Pazschke & Hedwigia, 23: 30 and 59 (1894)

Figure 4K

**Ravenelia macowaniana** Pazschke & Hedwigia, 23: 30 and 59 (1894). On leaves and in gall-transformed inflorescences of *Vachellia natalitia* (E.Mey) Kyal. & Boatwr. and on leaves of *V. permixta* (Burtt Davy) Kyal. & Boatwr. (Fabaceae: Mimosoideae).

**Specimens examined.** South Africa, Limpopo, Steelport, 24°41’32.3”S; 30°12’32.3”E, on leaves of *Vachellia permixta*, 23 February 2015, M. Ebinghaus ME424 (PREM61875); South Africa, Limpopo, Steelport, 24°41’32.3”S; 30°12’32.3”E, on leaves of *V. natalitia*, 19 February 2015, M. Ebinghaus ME416 (PREM61862); South Africa, Mpumalanga, East of Barberton, on leaves of *V. natalitia*, 2 June 2012, M. Ebinghaus ME158 (PREM61214); South Africa, Mpumalanga, Nelspruit, on leaves of *V. natalitia*, 10 January 2005, W. Maier WM3292 (PREM61218) and WM3294 (PREM61219); South Africa, Mpumalanga, South of Nelspruit, on leaves of *V. natalitia*, 10 March 2010, W. Maier WM3590 (PREM61226); South Africa, Mpumalanga, Nelspruit, on leaves of *V. natalitia*, 21 June 2005, W. Maier WM3423 (PREM61888); South Africa, Eastern Cape, Port St. Johns, on leaves of *V. natalitia*, 28 December 2005, W. Maier WM3453 (PREM61216); South Africa, North-West Province, Hartebeesspoort Dam, on leaves of *V. karroo* 23 June 2005, W. Maier WM3433 (PREM61222); South Africa, North-West Province, Hartebeesspoort Dam, on leaves of *V. karroo* June/July 2005, W. Maier WM3448 (PREM61221); South Africa, Eastern Cape, Haga Haga, on leaves of *V. karroo*, December 2004, W. Maier WM3485 (PREM61210); South Africa, on leaves of *V. karroo*, 15 May 2006, W. Maier WM3512 (PREM61220); South Africa, Western Cape, Worcester, on inflorescences of *V. karroo*, 20 December 2004, W. Maier WM3577 (KR-M-43406); South Africa, North-West Province 25°30’08.2”S; 27°21’32.4”E, on leaves of *V. karroo*, M. Ebinghaus ME433 (KR-M-43657).

**Notes.** Morphological as well as phylogenetic analyses, based on nrITS and nrLSU sequence data of the specimens PREM61214, PREM61216, PREM61218, PREM61219, PREM61862 and PREM61875 collected from *V. natalitia* and *V. permixta*, respectively, supported conspecificity and their placement in *R. macowaniana*. Therefore, we report *V. natalitia* and *V. permixta* as new hosts for *R. macowaniana*.

**Ravenelia evansii** Sydow, Ann. Mycol. 10, p. 440, Monograph. Ured. 3, p. 234

Figure 4J

**Ravenelia evansii** Sydow, Ann. Mycol. 10, p. 440, Monograph. Ured. 3, p. 234 On *Vachellia borleae* (Burtt Davy) Kyal. & Boatwr., *V. davyi* (N.E.Br.) Kyal. & Boatwr., *V. exuvalidis* (I. Verd.) Kyal. & Boatwr., *V. hebeclada* (DC.) Kyal. & Boatwr., *V. luederitzii* var. *retinens* (Sim.) (JH Ross & Brenan) Kyal. & Boatwr. and *V. swazica* (Burtt Davy) Kyal. & Boatwr. (Fabaceae: Mimosoideae).
**Emended description.** Telia subepidermally erumpent, dark brown to blackish, scattered or in loose groups on the abaxial side of leaflets, sori on the comparatively large leaflets of *V. robusta* ssp. *robusta* often forming concentric rings of 2.2–3.3 mm in diameter, single sori (120)230–500(710) μm in diameter with the largest telia appearing to develop in concentric arranged groups, subcircular to elongated; paraphyses lacking; teliospores circular to subcircular from topview, topside convex to almost hemispherical from lateral view, chestnut brown, (47)74–103(124) μm in diameter with (3)5–7(8) probasidial cells in a cross-section; single probasidial cells mostly single-layered, sometimes central cells and in rare events single cells two-layered, (16)23–30(39) μm from lateral view and (11)18–25(34) μm from top view; each probasidial cell with 3–5(8) spines; cysts hyaline and smooth, uniseriate and each cyst appears to be divided by a faint constriction, of the same number as peripheral probasidial cells, swelling in water but only slightly in lactophenol, pedicels compound, not persisting.

**Specimens examined.** All specimens examined for the emended species description of *R. evansii* representing new host associations are given in Table 1.

**Notes.** Rust infections on specimens of *Vachellia borleae* (ME384, PREM61869), *V. davyi* (PREM61845, PREM61005), *V. exuvialis* (PREM61876, PREM61868), *V. bebeclada* (PREM61227), *V. luderitzii* var. *retinens* (PREM61846) and *Vachellia swazica*. (PREM61002, PREM61028, PREM61008) were identified using morphological characters of the teliospores. These generally matched those of *R. evansii* Syd. & P. Syd. given in Doidge (1927) and Shivas et al. (2013) and were supported by molecular phylogenetic analyses of nrITS and nrLSU sequence data. These *Vachellia* species are reported as new hosts for *Ravenelia evansii*. Despite major congruence of teliospore characters, the range of the teliospore diameter observed by our examinations exceeded the size range of 50–80 μm and 63–90 μm given in Doidge (1927) and Shivas et al. (2013), respectively. Furthermore, the occurrence of two-layered probasidial cells is reported here for the first time (Figure 4J). Therefore, we present an emended description of the telial stage of *R. evansii* Syd. & P. Syd. emend. Ebinghaus, W. Maier and Begerow.

**Table 3.** Summary of morphological characters and spore sizes of *Ravenelia evansii*, *R. macowaniana* and *R. xanthophloeae*. All size measurements are given in μm, minimum and maximum values in parentheses. †Measurements and observations according to Doidge (1939).

| Teliospores | Diameter | Probasidal cell size | Ornamentation | No. of Cells in diameter |
|-------------|----------|-----------------------|---------------|--------------------------|
| *R. evansii* | (47)70–95(124) | (16.24–29(40) × (12)18–25(34)) | (1.5)4–6(8) | (3)5–7(8) |
| *R. macowaniana* | (50)82–105(118) | (19)23–32(40) × (13.5)18–24(34) | – | (3)5–6(7) |
| *R. xanthophloeae* | (40.5)65–75(82) | (19)22–25(39.5) × (12.5)19–23(32) | (0.5)1–2(3) | (3)4–5(6) |

| Aeciospores | Urediniospores |
|-------------|----------------|
| **Size** | **Size** | **Germ pores** |
| **Number** | **Arrangement** |
| *R. evansii* | 23–40 × 16–26† | 25–35 × 18–24† | 4† | equatorial† |
| *R. macowaniana* | 24–35 × 17–28† | 20–30 × 12.5–15† | 4† | equatorial† |
| *R. xanthophloeae* | (19)21–24(29) × (14)17–19(23) | (18)23–26(38) × (13)16–20(25) | 5–6 | equatorial |
New host associations and a novel species for the genus *Ravenelia*  

**Discussion**

The new rust species, *R. xanthophloeae*, was found only on the fever tree *Vachellia xanthophloea*. In South Africa, this tree species is naturally confined to habitats with shallow water tables in low-altitude areas of the northeastern KwaZulu-Natal, Mpumalanga and Limpopo Provinces (Coates Palgrave 2005, Smit 2008). However, it is frequently planted as ornamental at higher altitudes throughout Southern Africa, where infections by *R. macowaniana* on *V. karroo* are common. Yet, despite extensive sampling efforts, no rust has been reported from *V. xanthophloea* in these regions, suggesting that *R. xanthophloeae* might currently be restricted to the native range of its host tree. Furthermore, *V. xanthophloea* is apparently resistant to infection by the frequently co-occurring and closely related *R. macowaniana*. This observation lends additional support to the separation of *R. macowaniana* and *R. xanthophloeae* as distinct species.

Sequence divergence was smaller amongst the specimens of *R. evansii* than within *R. macowaniana* (Fig. 1). This is in contrast to teliospore morphology, where the six examined teliospore traits showed considerable variability in *R. evansii*, but very little variation in *R. macowaniana*. Specifically, an effect of the host association on teliospore morphology could be demonstrated and this was most pronounced in specimens of *R. evansii* collected from *V. borleae*, *V. dasyi* and *V. exuvialis*. It has been demonstrated in other fungal and oomycetous plant pathogens that infraspecific variation of spore traits might correlate with host species (Savile 1976, Lutz et al. 2005, Runge and Thines 2011). However, mechanisms leading to such host-associated differences in rust fungi remain obscure. Savile (1976) hypothesised that differences in host compatibility of rust fungi potentially lead to differences in nutrient supply and could consequently influence morphological features. It was also speculated that host anatomy such as the thickness of the cuticle and epidermis might influence spore morphology (Scholler et al. 2011). Clearly, experimental studies that focus on the differential effects of the host and environment on morphological character expression in the rust fungi are needed to resolve this question.

In the present study, the host ranges of *R. macowaniana* and *R. evansii* were expanded from one to three and from four to nine hosts, respectively. Thus, these two rust species have a broader host range than previously reported and parasitise several co-occurring acacia species in the South African savannah biome. This is in contrast to recent findings in the genus *Endoraecium* that infect Australian wattles (*Acacia* s. str., formerly *Acacia* subg. *Phyllodineae*). Based on morphological and molecular phylogenetic studies, species previously thought to have a broad host range were split into several species infecting mostly a single tree species (Berndt 2011, McTaggart et al. 2015). In this case, a general pattern of co-speciation of the parasites with their hosts was suggested (McTaggart et al. 2015). In the South African acacia rusts however, the shared distribution ranges of their hosts may prevent the rusts from speciation by recurrent gene flow between metapopulations. In contrast, *Ravenelia xanthophloeae* appears to infect only *V. xanthophloea*. In the phylogenetic analyses, this rust was closely related to *R. macowaniana*, which suggests a more recent speciation of both species. The host of *R. xanthophloeae* is eco-geographically clearly separated from the hosts of *R. macowaniana* and the different niches of the host most likely contributed considerably to the divergence of the parasite species.
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Supplementary material 1

Figure S1
Authors: Malte Ebinghaus, Wolfgang Maier, Michael J. Wingfield, Dominik Begerow
Data type: statistical data
Explanation note: Boxplot of measurements of the six defined teliospore characters of *R. evansii*. Values were obtained from teliospores derived from seven different host species of in total 18 individual trees. The boxplots are based on mean values calculated for all investigated teliospores for each specimen, respectively.
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Link: https://doi.org/10.3897/mycokeys.43.25090.suppl

Supplementary material 2

Figure S2
Authors: Malte Ebinghaus, Wolfgang Maier, Michael J. Wingfield, Dominik Begerow
Data type: statistical data
Explanation note: Boxplot of measurements of the six defined teliospore characteristics of *R. macowaniana* and *R. xanthophloeae*. Values were obtained from teliospores derived from three different host species of in total 10 individual trees. The boxplots are based on mean values calculated for all investigated teliospores for each specimen, respectively.
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Link: https://doi.org/10.3897/mycokeys.43.25090.suppl2
Supplementary material 3

Table S1
Authors: Malte Ebinghaus, Wolfgang Maier, Michael J. Wingfield, Dominik Begerow
Data type: species data
Explanation note: List of measurements of teliospore characters of *R. evansii*. Obtained values were sorted by voucher and by individual teliospores. All measurements are given in μm.
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Link: https://doi.org/10.3897/mycokeys.43.25090.suppl3

Supplementary material 4

Table S2
Authors: Malte Ebinghaus, Wolfgang Maier, Michael J. Wingfield, Dominik Begerow
Data type: species data
Explanation note: List of measurements of teliospore characters of *R. macowaniana* and *R. xanthophloeae*. Obtained values were sorted by voucher and by individual teliospores. All measurements are given in μm.
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