# A test of *psbK-psbI* and *atpF-atpH* as potential plant DNA barcodes using the flora of the Kruger National Park as a model system (South Africa)

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## Introduction

DNA barcoding is a new technique that uses short, standardized DNA sequences (400-800 bp) of an organism to determine its identity. Because this sequence has to be variable enough to identify individual species, but not too variable within the same species so that a clear threshold can be defined between intra- and inter-specific diversities, it is very challenging to apply this technique to all species on the planet . A DNA barcode has been identified for animals, i.e. the mitochondrial gene *cox1*, which shows strong abilities in identifying cryptic species, accelerating biodiversity inventories and helping to identify species from degraded material (e.g. to control trade of threatened) . For plants, the identification of a suitable DNA barcode is more problematic. Cho et al. showed that mitochondrial DNA evolves too slowly in plants to provide a region variable enough to discriminate between species. Then the quest for the best suitable barcode started and is still ongoing .

Kress et al. opened the debates and suggested the use of multiple genes to identify plant species quickly and accurately. At the Second International Barcode of Life Conference in Tapei (September 2007), at least five different plant DNA barcodes were proposed, but no consensus reached. Among those, both *atpF-atpH* and *psbK-psbI* suggested by Kim et al. have not yet been tested. Here, we evaluate the use of these loci as DNA barcodes for plants by applying them to a wide range of plant species. The two new intergenic loci *atpF-atpH* and *psbK-psbL* are both localized in the large single copy (LSC) of the plastid genome. The genes *atpF* and *atpH* encode ATP synthase subunits CFO I and CFO III, respectively. Both genes *psbK* and *psbI* encode two low molecular mass polypeptides, K and I, respectively, of the photo-system II. These two new loci are conservative from algae to land plants and even in parasitic plants . In this study, we focus on the trees and shrubs from the Kruger National Park (hereafter KNP), part of the Maputaland-Pondoland-Albany hotspot in southern Africa. On a selected sampling from

the 2,700 taxa surveyed in the area, we applied several metrics following Lahaye et al. to evaluate the efficiency of combining *matK* either to *trnH-psbA* and/or *atpF-atpH* and/or *psbK-psbI* for DNA barcoding purposes.

#### **Material and Methods**

*Sampling*. In total 101 taxa from the KNP were sampled, covering 18 families from the monocotyledons to the euasterids II. This dataset included 31 species of trees and shrubs in which we had more than one representative per species, 3 species of Orchids, one of which with 2 representatives, and 3 parasitic plants, one of which is achlorophyllous. Parasitic plants have been sampled to test the universality of the potential DNA barcodes. We used *Amborella trichopoda* Baill. (complete genome GenBank accession AJ506156) as outgroup for the phylogenetic analyses. All specimens were collected in different ecosystems when possible (Figure 1) and voucher specimens are available as detailed in Table 1.

*Collection and preservation.* Collection of plant material was done in the KNP with the assistance of the park's rangers. Plants were sampled and pressed for herbarium voucher specimens in triplicate, one for the herbarium of the KNP, one for Kew Herbarium (K; United Kingdom), and one for the herbarium at Pretoria (PRE; South Africa). Information about the locality and habit of collected plants were entered on a palmtop-GPS to facilitate their further treatment, and also noted on hard copy for security. For each plant collected, leaf material was stored in silica for molecular studies, and flowers and fruit stored in ethanol when available.

**DNA sequencing.** Total DNA was extracted from dried leaf material using the standard method of Doyle and Doyle and cleaned with QIAquick silica columns (Qiagen, Helden,

Germany). Sequences of matK and trnH-psbA for each taxa were published in Lahaye et al. and their accession numbers are available from GenBank (Table 1). We amplified atpF-atpH and psbK-psbI using PCR as follows: 35 cycles, 30 sec denaturation at 94°C, 40 sec annealing at 51°C, and 40 sec extension at 72°C. Primers were kindly provided by Ki-Joong: atpF-atpH- atpF 5'-ACTCGCACACACTCCCTTTCC-3', atpH Kim 5'-GCTTTTATGGAAGCTTTAACAAT-3'; psbK-psbI: psbKand 5'-TTAGCCTTTGTTTGGCAAG-3', psbl- 5'-AGAGTTTGAGAGTAAGCAT-3'. After cycle sequencing using Big Dye terminator v3.1 and sequencing on a 3130xl genetic analyzer (Applied Biosystems, UK), electropherograms were edited using SEQUENCER 4.6 software (Genes Codes Corporation, USA) and DNA sequences aligned by eye in PAUP4b10\* (incomplete sequences at both ends were excluded from the analyses). Taxa with missing data (amplification or sequencing failed) were removed from the combined matrix in order to analyze complete matrices.

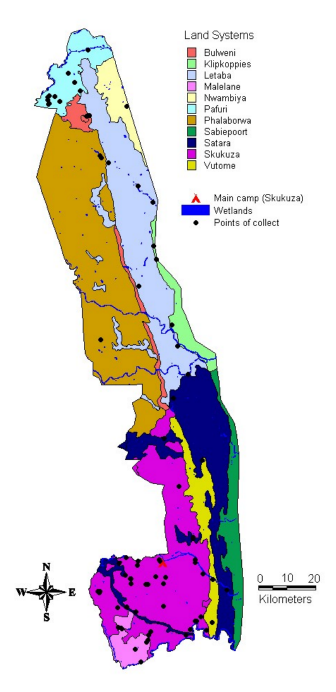


Figure 1. Map of the KNP with landsystems following Venter (1990) and collecting points from this study

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Plant Family	name Checked on IPNI	Voucher	Location	GPS	Altitude	matK	trnH-psbA	atpF-atpH	psbK-psbI
Fabaceae	Acacia exuvialis Verdoorn	OM260	KNP	S24 58 54.3 E31 34 26.3	284 m	EU214205	EU213781	-	EU626889
Fabaceae	Acacia exuvialis Verdoorn	RL1204	KNP	S25 29 35.4 E31 28 12.3	319 m	EU214206	EU213782	EU626796	EU626890
Fabaceae	Acacia exuvialis Verdoorn	RL1412	KNP	S25 21 41.5 E31 30 56.5	320 m	EU214207	EU213783	-	EU626891
Fabaceae	Acacia nigrescens Oliver	RL1111	KNP	S25 06 26.4 E31 30 24.5	452 m	EU214208	-	EU626797	EU626892
Fabaceae	Acacia nigrescens Oliver	RL1205	KNP	S25 29 35.4 E31 28 12.3	319 m	EU214209	EU213784	EU626798	EU626893
Fabaceae	Acacia nigrescens Oliver	RL1656	KNP	S22 41 29.6 E31 01 37.2	439 m	EU214210	EU213785	EU626799	EU626894
Fabaceae	Acacia tortilis Hayne	OM261	KNP	S24 59 20.9 E31 34 34.5	266 m	EU214213	EU213788	EU626800	EU626895
Fabaceae	Acacia tortilis Hayne	RL1483	KNP	S24 36 53.6 E31 40 51.4	333 m	EU214211	EU213786	EU626801	EU626896
Fabaceae	Acacia tortilis Hayne	RL1608	KNP	S22 57 38.1 E31 14 50.5	302 m	EU214212	EU213787	EU626802	EU626897
Orchidaceae	Acampe praemorsa ( Roxb. ) Blatt. & McCann	RBN203	KNP	S22 42 06.1 E30 58 14.4	504 m	EU214214	EU213789	EU626803	EU626898
Amborellaceae	Amborella trichopoda Baill.	-	-	-	-	AJ506156	AJ506156	AJ506156	AJ506156
Orchidaceae	Ansellia africana Lindl.	OM1163	KNP	S25 12 54.8 E31 35 36.0	280 m	EU214215	-	EU626804	EU626899
Orchidaceae	Ansellia africana Lindl.	OM531	KNP	S25 19 54.3 E31 44 28.5	225 m	EU214216	-	EU626805	EU626900
Orchidaceae	Bonatea speciosa Willd.	RL1158	KNP	S25 13 11.4 E31 23 41.8	472 m	EU214217	EU213790	EU626806	EU626901
Asteraceae	Brachylaena huillensis O.Hoffm.	OM1281	KNP	S23 28 54.6 E31 33 27.0	421 m	EU214218	EU213791	EU626807	EU626902
Asteraceae	Brachylaena huillensis O.Hoffm.	OM247	KNP	S25 06 12.7 E31 35 44.2	276 m	EU214219	EU213792	EU626808	EU626903
Asteraceae	Brachylaena huillensis O.Hoffm.	RBN360	KNP	S22 42 51.4 E31 23 46.3	507 m	EU214220	EU213793	EU626809	EU626904
Combretaceae	Combretum apiculatum Sond.	RL1100	KNP	S25 06 24.7 E31 30 41.4	389 m	EU214221	EU213794	EU626810	EU626905
Combretaceae	Combretum apiculatum Sond.	RL1185	KNP	S25 23 11.2 E31 30 42.1	391 m	EU214222	EU213795	EU626811	EU626906
Combretaceae	Combretum apiculatum Sond.	RL1355	KNP	S25 20 11.4 E31 49 48.0	213 m	EU214223	EU213796	EU626812	EU626907
Combretaceae	Combretum collinum Fresen.	OM722	KNP	S25 00 07.4 E31 21 07.0	378 m	EU214224	EU213797	EU626813	EU626908
Combretaceae	Combretum collinum Fresen.	RL1164	KNP	S25 14 44.5 E31 26 39.8	419 m	EU214225	EU213798	EU626814	EU626909
Combretaceae	Combretum collinum Fresen.	RL1392	KNP	S25 25 45.2 E31 26 26.4	334 m	EU214226	EU213799	EU626815	EU626910
Combretaceae	Combretum hereroense Schinz	RL1120	KNP	S25 06 28.6 E31 29 58.5	383 m	EU214227	EU213800	EU626816	EU626911
Combretaceae	Combretum hereroense Schinz	RL1183	KNP	S25 23 11.2 E31 30 42.1	391 m	EU214228	EU213801	EU626817	EU626912
Combretaceae	Combretum hereroense Schinz	RL1364	KNP	S25 17 18.5 E31 46 34.6	235 m	EU214229	EU213802	EU626818	EU626913
Euphorbiaceae	Croton gratissimus Burch	OM785	KNP	S23 48 24.9 E31 38 27.2	285 m	EU214230	EU213803	EU626819	EU626914
Euphorbiaceae	Croton gratissimus Burch	RL1619	KNP	S22 45 43.6 E31 10 50.8	379 m	EU214231	EU213804	EU626820	EU626915
Euphorbiaceae	Croton gratissimus Burch	RL1621	KNP	S22 45 52.1 E31 10 29.1	414 m	EU214232	EU213805	EU626821	EU626916

Plant Family	name Checked on IPNI	Voucher	Location	GPS	Altitude	matK	trnH-psbA	atpF-atpH	psbK-psbI
Euphorbiaceae	Croton megalobotrys Müll.Arg.	OM774	KNP	S24 03 13.4 E31 43 50.0	211 m	EU214233	EU213806	EU626822	EU626917
Euphorbiaceae	Croton megalobotrys Müll.Arg.	RL1540	KNP	S23 54 53.6 E31 40 18.7	201 m	EU214234	EU213807	EU626823	EU626918
Euphorbiaceae	Croton megalobotrys Müll.Arg.	RL1574	KNP	S23 11 37.5 E31 32 16.5	246 m	EU214235	EU213808	EU626824	EU626919
Euphorbiaceae	Croton pseudopulchellus Pax	RBN186	KNP	S22 39 57.7 E30 59 57.6	468 m	EU214236	EU213809	EU626825	EU626920
Euphorbiaceae	Croton pseudopulchellus Pax	RBN262	KNP	S22 26 00.7 E31 10 57.6	291 m	EU214237	EU213810	EU626826	EU626921
Euphorbiaceae	Croton pseudopulchellus Pax	RL1650	KNP	S22 40 09.2 E30 57 39.6	451 m	EU214238	EU213811	-	EU626922
Orchidaceae	Eulophia R.Br.	OM473	KNP	S25 03 40.0 E31 23 11.2	351 m	EU214239	EU213812	EU626827	EU626923
Proteaceae	Faurea rochetiana Chiov. ex Pic.Serm.	OM1413	KNP	S25 08 43.0 E31 14 33.4	726 m	EU214240	EU213813	EU626828	EU626924
Proteaceae	Faurea rochetiana Chiov. ex Pic.Serm.	OM1461	KNP	S25 08 43.6 E31 14 32.6	722 m	EU214241	EU213814	EU626829	EU626925
Proteaceae	Faurea rochetiana Chiov. ex Pic.Serm.	OM1463	KNP	S25 08 43.1 E31 14 33.1	727 m	EU214242	EU213815	EU626830	EU626926
Proteaceae	Faurea saligna Harv.	OM1438	KNP	S25 19 31.7 E31 21 42.3	486 m	EU214243	EU213816	EU626831	EU626927
Proteaceae	Faurea saligna Harv.	OM1443	KNP	S25 19 16.9 E31 20 59.5	523 m	EU214244	EU213817	EU626832	EU626928
Proteaceae	Faurea saligna Harv.	OM1445	KNP	S25 19 39.5 E31 22 08.8	473 m	EU214245	EU213818	EU626833	EU626929
Moraceae	Ficus abutilifolia Miq.	OM557	KNP	S25 04 41.4 E31 24 54.5	414 m	EU214248	EU213821	EU626834	EU626930
Moraceae	Ficus abutilifolia Miq.	RL1471	KNP	S24 52 32.9 E31 45 21.9	256 m	EU214246	EU213819	EU626835	EU626931
Moraceae	Ficus abutilifolia Miq.	RL1501	KNP	S24 22 39.3 E31 35 51.8	369 m	EU214247	EU213820	EU626836	EU626932
Moraceae	Ficus glumosa Delile	OM564	KNP	S25 04 36.8 E31 25 03.7	473 m	EU214249	EU213822	EU626837	EU626933
Moraceae	Ficus glumosa Delile	RL1407	KNP	S25 23 41.1 E31 30 02.4	466 m	EU214250	EU213823	EU626838	EU626934
Moraceae	Ficus glumosa Delile	RL1429	KNP	S25 08 29.6 E31 14 42.6	665 m	EU214251	EU213824	EU626839	-
Moraceae	Ficus sycomorus L.	RBN197	KNP	S22 40 53.4 E30 57 43.2	445 m	EU214252	EU213825	EU626840	EU626935
Moraceae	Ficus sycomorus L.	RL1598	KNP	S23 06 46.1 E31 27 16.5	264 m	EU214253	EU213826	EU626841	EU626936
Moraceae	Ficus sycomorus L.	RL1614	KNP	S22 45 43.1 E31 11 18.4	356 m	EU214254	EU213827	EU626842	EU626937
Malvaceae	Grewia bicolor Juss.	OM329	KNP	S25 04 18.8 E31 36 29.5	363 m	EU214255	EU213828	EU626843	EU626938
Malvaceae	Grewia bicolor Juss.	RL1545	KNP	S23 36 52.2 E31 27 36.5	290 m	EU214256	EU213829	EU626844	EU626939
Malvaceae	Grewia bicolor Juss.	RL1658	KNP	S22 41 29.6 E31 01 37.2	439 m	EU214257	EU213830	EU626845	EU626940
Malvaceae	Grewia flavescens Juss.	OM323	KNP	S25 04 18.8 E31 36 29.5	363 m	EU214258	EU213831	-	EU626941
Malvaceae	Grewia flavescens Juss.	RL1472	KNP	S24 52 32.9 E31 45 21.9	256 m	EU214259	EU213832	EU626846	EU626942
Malvaceae	Grewia flavescens Juss.	RL1604	KNP	S22 58 18.8 E31 15 13.5	305 m	EU214260	-	-	-
Malvaceae	Grewia villosa Willd.	RL1342	KNP	S24 58 56.5 E31 46 02.3	208 m	EU214261	EU213833	-	EU626943
Malvaceae	Grewia villosa Willd.	RL1523	KNP	S24 10 31.8 E31 38 53.8	255 m	EU214262	EU213834	EU626847	EU626944

Plant Family	name Checked on IPNI	Voucher	Location	GPS	Altitude	matK	trnH-psbA	atpF-atpH	psbK-psbI
Malvaceae	Grewia villosa Willd.	RL1569	KNP	S23 24 48.9 E31 32 21.8	363 m	EU214263	EU213835	-	EU626945
Apiaceae	Heteromorpha arborescens Cham. & Schltdl.	OM1430	KNP	S25 13 27.0 E31 20 34.3	655 m	EU214264	EU213836	EU626848	EU626946
Apiaceae	Heteromorpha arborescens Cham. & Schltdl.	OM1488	KNP	S24 59 58.3 E31 21 04.3	359 m	EU214265	EU213837	EU626849	EU626947
Apiaceae	Heteromorpha arborescens Cham. & Schltdl.	OM1516	KNP	S25 20 29.0 E31 31 25.8	426 m	EU214266	EU213838	EU626850	EU626948
Hydnoraceae	Hydnora johannis Becc.	OM534	KNP	\$25 21 37.5 E31 43 11.1	241 m	EU214267	-	-	EU626949
Arecaceae	Hyphaene coriacea Gaertn.	OM1184	KNP	S25 08 03.4 E31 56 37.7	167 m	EU214268	EU213775	EU626851	EU626950
Arecaceae	Hyphaene coriacea Gaertn.	OM1187	KNP	S25 17 45.4 E31 51 44.5	185 m	EU214269	EU213776	EU626852	EU626951
Arecaceae	Hyphaene coriacea Gaertn.	OM236	KNP	S25 03 08.3 E31 48 38.6	201 m	EU214271	EU213778	EU626853	EU626952
Arecaceae	Hyphaene coriacea Gaertn.	OM755	KNP	S24 29 10.7 E31 48 29.4	259 m	EU214270	EU213777	EU626854	EU626953
Arecaceae	Hyphaene petersiana Klotzsch ex Mart	OM1296	KNP	S22 38 18.4 E31 08 25.1	382 m	EU214272	EU213779	EU626855	EU626954
Arecaceae	Hyphaene petersiana Klotzsch ex Mart	OM908	KNP	S22 32 55.9 E31 04 25.5	347 m	EU214273	EU213780	EU626856	EU626955
Myrothamnaceae	Myrothamnus flabellifolia Welw.	OM1137	KNP	S25 06 15.4 E31 24 58.6	452 m	EU214275	EU213840	EU626857	EU626956
Myrothamnaceae	Myrothamnus flabellifolia Welw.	OM1209	KNP	S25 04 03.5 E31 33 04.7	485 m	EU214276	EU213841	EU626858	EU626957
Myrothamnaceae	Myrothamnus flabellifolia Welw.	OM285	KNP	S25 04 01.2 E31 33 04.8	577 m	EU214274	EU213839	EU626859	EU626958
Anacardiaceae	Rhus gueinzii Sond.	OM265	KNP	S24 59 25.4 E31 27 19.6	268 m	EU214277	EU213842	EU626860	EU626959
Anacardiaceae	Rhus gueinzii Sond.	RL1366	KNP	S25 17 23.1 E31 46 06.3	208 m	EU214278	EU213843	EU626861	EU626960
Anacardiaceae	Rhus gueinzii Sond.	RL1474	KNP	S24 52 08.3 E31 45 22.4	283 m	EU214279	EU213844	EU626862	EU626961
Anacardiaceae	Rhus leptodictya Diels	RBN205	KNP	S22 42 13.5 E30 57 56.4	487 m	EU214280	EU213845	EU626863	EU626962
Anacardiaceae	Rhus leptodictya Diels	RL1645	KNP	S22 42 06.5 E30 58 10.5	499 m	EU214281	EU213846	EU626864	EU626963
Anacardiaceae	Rhus leptodictya Diels	RL1655	KNP	S22 41 29.1 E31 01 38.4	448 m	EU214282	EU213847	EU626865	EU626964
Anacardiaceae	Rhus transvaalensis Engl.	OM282	KNP	S25 08 53.2 E31 14 38.3	664 m	EU214283	EU213848	EU626866	EU626965
Anacardiaceae	Rhus transvaalensis Engl.	OM943	KNP	S25 08 30.6 E31 14 07.8	610 m	-	EU213849	EU626867	EU626966
Anacardiaceae	Rhus transvaalensis Engl.	RL1427	KNP	S25 08 59.4 E31 14 35.0	630 m	EU214284	EU213850	EU626868	EU626967
Solanaceae	Solanum panduriforme Drège ex Dunal	OM1115	KNP	\$25 00 44.2 E31 27 13.7	341 m	EU214285	EU213851	EU626869	EU626968
Solanaceae	Solanum panduriforme Drège ex Dunal	OM326	KNP	S25 04 18.8 E31 36 29.5	363 m	EU214286	EU213852	EU626870	EU626969
Solanaceae	Solanum panduriforme Drège ex Dunal	OM350	KNP	S25 04 17.5 E31 36 29.2	354 m	EU214287	EU213853	EU626871	EU626970
Apiaceae	Steganotaenia araliacea Hochst.	OM1350	KNP	\$23 52 55.8 E31 15 00.9	422 m	EU214288	EU213854	EU626872	EU626971
Apiaceae	Steganotaenia araliacea Hochst.	OM1517	KNP	S23 52 56.3 E31 15 06.4	420 m	EU214289	EU213855	EU626873	EU626972
Apiaceae	Steganotaenia araliacea Hochst.	OM566	KNP	S25 04 36.8 E31 25 03.7	473 m	EU214290	EU213856	EU626874	EU626973
Orobanchaceae	Striga elegans Benth.	OM683	KNP	S25 04 02.4 E31 33 06.1	383 m	EU214291	-	EU626875	EU626974

Plant Family	name Checked on IPNI	Voucher	Location	GPS	Altitude	matK	trnH-psbA	atpF-atpH	psbK-psbI
Loganiaceae	Strychnos decussata ( Pappe ) Gilg	OM900	KNP	\$22 35 35.0 E31 06 37.5	329 m	EU214292	EU213857	EU626876	EU626975
Loganiaceae	Strychnos decussata ( Pappe ) Gilg	RL1560	KNP	\$23 24 53.0 E31 32 29.7	379 m	EU214293	EU213858	EU626877	EU626976
Loganiaceae	Strychnos decussata ( Pappe ) Gilg	RL1561	KNP	\$23 24 53.0 E31 32 29.7	379 m	EU214294	EU213859	EU626878	EU626977
Loganiaceae	Strychnos madagascariensis Spreng. ex Baker	RL1433	KNP	S25 08 24.1 E31 14 51.5	641 m	EU214295	EU213860	EU626879	EU626978
Loganiaceae	Strychnos madagascariensis Spreng. ex Baker	RL1460	KNP	S24 58 21.4 E31 23 21.8	342 m	EU214296	EU213861	EU626880	EU626979
Loganiaceae	Strychnos madagascariensis Spreng. ex Baker	RL1559	KNP	\$23 24 53.0 E31 32 29.7	379 m	EU214297	EU213862	EU626881	EU626980
Loganiaceae	Strychnos spinosa Lam.	OM220	KNP	S24 59 49.9 E31 46 10.3	208 m	EU214298	EU213863	EU626882	EU626981
Loganiaceae	Strychnos spinosa Lam.	RL1346	KNP	\$25 04 51.2 E31 51 53.2	185 m	EU214299	EU213864	EU626883	EU626982
Loganiaceae	Strychnos spinosa Lam.	RL1652	KNP	S22 39 39.3 E30 58 17.4	430 m	EU214300	EU213865	EU626884	EU626983
Loranthaceae	Tapinanthus Blume	OM825	KNP	S22 59 46.4 E31 17 32.6	312 m	EU214301	-	EU626885	EU626984
Velloziaceae	Xerophyta retinervis Baker	OM1213	KNP	\$25 08 32.4 E31 14 23.7	678 m	EU214302	EU213866	EU626886	EU626985
Velloziaceae	Xerophyta retinervis Baker	OM516	KNP	\$25 16 03.6 E31 47 53.3	267 m	EU214303	EU213867	EU626887	EU626986
Velloziaceae	Xerophyta retinervis Baker	OM562	KNP	\$25 04 36.8 E31 25 03.7	473 m	EU214304	EU213868	EU626888	EU626987

Table 1. Material sampled for this study, species checked in IPNI, voucher, GPS and altitude information, GenBank accession numbers. All vouchers have been collected in triplicate, one for Kew Herbarium, one for the herbarium of the KNP at Skukuza (South Africa), and one for the National Herbarium at Pretoria (South Africa).

Genetic analyses. Inter- and intra-specific genetic divergences were calculated using each potential DNA barcode following Meyer and Paulay. Three different metrics were used to characterize intra-specific divergence: (i) average pairwise distances between all individuals sampled within those species that had at least two representatives, (ii) 'mean theta', with theta being the average pairwise distances calculated for each species that had more than one representative, thereby eliminating biases associated with uneven sampling among taxa and (iii) average coalescent depth, i.e. the depth of a node linking all sampled extant members of a species, 'book-ending' intra-specific variability. Genetic distances between con-generic species were used to characterize inter-specific divergence. For each barcode, pairwise distances were calculated with the simplest K2P model following Lahaye et al. in which this model showed the best results. This model also utilizes the CBOL advises for distance calculations (barcoding.si.edu/). Wilcoxon Signed Rank Tests were performed to compare intra- and inter-specific variability for every pair of barcodes following Kress and Erickson. We evaluated 'DNA barcoding gaps' by comparing the distribution of intra- versus inter-specific divergences. Median and Wilcoxon Two-Sample Tests were used to evaluate whether these distributions overlapped.

*Phylogenetic analyzes.* To evaluate whether species were recovered as monophyletic with each barcode, we used standard phylogenetic techniques. Note that this is not to say that barcodes can be used to reconstruct phylogenies, because in this case we are disregarding the recovered inter-specific relationships. Trees were built with PAUP4b10\* using Maximum Parsimony (MP) and UPGMA, the two best algorithms in terms of percentages of species correctly identified . UPGMA trees were inferred with PAUP4b10\* from K2P distances. MP analyses were performed using tree bisection-reconnection (TBR), branch swapping and 1,000 random addition sequence replicates keeping 10 trees at each step. MP analyses have been performed with and without coding

indels as a 5<sup>th</sup> state in order to assess the impact of keeping this information for barcoding purposes.

*Coalescence analyses.* For each barcode, we identified those clusters that were derived from an independent coalescence process and asked whether they matched previously recognized taxonomic species, using methods developed by Pons et al. and Fontaneto et al. . The likelihood of waiting times between successive branching events on a DNA barcode-based tree was calculated under the null model that all terminals were derived from a single coalescence process, and under the alternative model that all taxa derived from a set of two independently evolving populations. With the alternative model, a threshold age T was calculated, at which point the older nodes represented inter-specific diversification events whereas the younger nodes represented separate coalescent processes typical of intra-specific clusters. We used DNA barcode-based trees from MP and transformed branch lengths with nonparametric rate smoothing to produce ultrametric trees, i.e. branch lengths reflecting time only. We also used the ultrametric UPGMA trees. Likelihood models were determined using an R script available from TGB.

#### **Results & Discussion**

*Molecular characteristics and PCR success.* Amplification was generally successful for each potential barcode tested with more than 92% of taxa successfully amplified and sequenced (Table 2). The best percentage was given by *matK* with 99% of taxa sequenced and the lowest percentage was obtained for *trnH-psbA* with 92%. The potential DNA barcode *psbK-psbI* showed PCR and sequencing performances very close to those of *matK* with 98% of taxa successfully amplified. Both *atpF-atpH* and *trnH-psbA* failed to amplify the parasitic/non-chlorophytic plant *Hydnora johanis*. Alignment of sequences was unproblematic for *matK* and *psbK-psbI*, but *trnH-psbA* and *atpF-atpH* 

presented significant difficulties due to a high level of length variation (225 to 758 bp and 218 to 847 bp, respectively). Because its alignment was not reliable by Clustal X, we performed a first visual alignment between congeneric species and then aligned all taxa by adding as many gaps as necessary to keep the homology between congeneric species for inter- and intraspecific calculations. The alignment of *trnH-psbA* revealed a highly conservative intron only for the Orchidaceae and Amaryllidaceae which has been identified previously . Combining *matK* to one of the other potential barcodes allowed building a matrix including sequences for all taxa (Table 2).

matK	99%
psbK-psbl	98%
trnH-psbA	92.1%
atpF-atpH	93.1%
matK+trnH-psbA	100%
matK+trnH-psbA+atpF-atpH	100%
matK+trnH-psbA+psbK-	
psbl	100%
matK+atpF-atpH	100%
matK+psbK-psbl	100%
matK+atpF-atpH+psbK-psbI	100%
4 loci	100%

Table 2. Percentages of taxa represented in each matrix by at least one sequence.

*Intra- and Inter-specific diversities*. Performances of each DNA barcode was assessed by means of inter- and intra-specific diversity calculated from K2P (Kimura's two parameters) pairwise distance matrices (barcoding.si.edu/; Table 3). The highest interspecific diversity was reached by *atpF-atpH* (3.45%) followed by *trnH-psbA* (2.55%) and the lowest was given by *psbK-psbI* (1.06%) with *matK* between these (1.34%). Regarding the different metrics to infer the intra-specific differences, the mean theta was in most cases similar to the average of overall intra-specific distances because there is no bias associated with species over-sampled in our study with the majority of the species represented by three specimens. The mean coalescent depth was slightly superior to the average of overall interspecific distances because it takes into consideration only the highest distance between specimens sampled for a species. Results showed the highest mean of intraspecific differences for *trnH-psbA* regardless of the metric used (Table 3). The lowest values were obtained for both *atpF-atpH* and *psbK-psbI*. Wilcoxon rank tests performed on the different distance matrices showed with very high significance that *trnH-psbA* had by far the highest inter-specific variability, followed by *matK* and *atpF-atpH*, which had a similar divergence (Table 4). The highest intra-specific distances were also significantly reached by *trnH-psbA* whereas the three other loci presented almost similar values (Table 5).

		matK	trnH-	atpF-	psbK-	4 loci	matK+	matK+atpF-	matK+psbK-	matK+	matK+	matK+psbK- psbl+
					<b>P</b>		trnH-	atpH+trnH-	psbl+trnH-	atpF-	psbK-	
			psbA	atpH	psbl		psbA	psbA	psbA	atpH	psbl	atpF-atpH
Mean of all inters	pecific											
distances	poono	0.0134	0.0255	0.0345	0.0106	0.0172	0.0175	0.0189	0.0157	0.0168	0.0118	0.0150
St. deviation	+/-	0.0127	0.0227	0.0665	0.0096	0.0151	0.0154	0.0180	0.0121	0.0201	0.0092	0.0159
Mean of all intras	pecific											
distances		0.0012	0.0017	0.0004	0.0005	0.0009	0.0012	0.0009	0.0011	0.0007	0.0009	0.0007
St. deviation	+/-	0.0040	0.0041	0.0015	0.0012	0.0015	0.0026	0.0017	0.0021	0.0020	0.0026	0.0016
Mean Theta		0.0012	0.0015	0.0007	0.0005	0.0008	0.0012	0.0009	0.0010	0.0007	0.0009	0.0007
St. deviation	+/-	0.0037	0.0032	0.0023	0.0010	0.0013	0.0023	0.0015	0.0018	0.0018	0.0024	0.0015
Mean coalescent	depth	0.0017	0.0023	0.0008	0.0008	0.0013	0.0017	0.0014	0.0016	0.0012	0.0013	0.0011
St. deviation	+/-	0.0050	0.0047	0.0023	0.0016	0.0018	0.0032	0.0021	0.0026	0.0026	0.0033	0.0021
Number of measure	ments for	93	90	84	91	95	95	95	95	95	95	95
all intraspecific dis	stances											
Number of measure	ments for	200	194	168	194	206	206	206	206	206	206	206
all interspecific dis	stances			-								

Table 3. Measures of inter- and intra-specific K2P distances for four potential barcodes and different combinations applied to a selective sampling from the KNP.

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	Wilcoxon Signed-Ranks Test –	
	Interspecific pair-distances	
matK vs trnH-psbA	W+ = 1462, W- = 14648, N = 179, p <= 2.216e-21	matK< <trnh-psba< th=""></trnh-psba<>
matK vs atpF-atpH	W+ = 4977, W- = 5608, N = 145, p <= 0.5341	matK = atpF-atpH
matK vs psbK-psbl	W+ = 8655, W- = 6051, N = 171, p <= 0.0447	matK > psbK-psbl
trnH-psbA vs atpF-atpH	W+ = 8482, W- = 3608, N = 155, p <= 1.345e-05	trnH-psbA > atpF-atpH
trnH-psbA vs psbK-psbl	W+ = 13538, W- = 2572, N = 179, p <= 2.88e-15	trnH-psbA >> psbK-psbl
atpF-atpH vs psbK-psbl	W+ = 7663, W- = 2922, N = 145, p <= 2.902e-06	atpF-atpH > psbK-psbl
4 loci vs matK+trnH-psbA	W+ = 7286, W- = 12217, N = 197, p <= 0.002095	4 loci < matK+trnH-psbA
4 loci vs matK+trnH-psbA+atpF-atpH	W+ = 5244, W- = 14259, N = 197, p <= 1.859e-08	4 loci < matK+trnH-psbA+atpF-atpH
4 loci vs matK+trnH-psbA+psbK-psbl	W+ = 6661, W- = 12060, N = 193, p <= 0.0005137	4 loci < matK+trnH-psbA+psbK-psbl
4 loci vs matK+atpF-atpH	W+ = 14310, W- = 5193, N = 197, p <= 1.284e-08	4 loci > matK+atpF-atpH
4 loci vs matK+psbK-psbl	W+ = 15830, W- = 3673, N = 197, p <= 3.333e-14	4 loci > matK+psbK-psbl
4 loci vs matK+psbK-psbI+atpF-atpH	W+ = 15351, W- = 4152, N = 197, p <= 2.807e-12	4 loci < matK+atpF-atpH+psbK-psbl
matK+trnH-psbA vs matK+trnH-psbA+atpF-atpH	W+ = 12287, W- = 6434, N = 193, p <= 0.0001661	matK+trnH-psbA > matK+trnH-psbA+atpHF
		matK+trnH-psbA > matK+trnH-psbA+psbK-
matK+trnH-psbA vs matK+trnH-psbA+psbK-psbI	W+ = 13374, W- = 6129, N = 197, p <= 6.174e-06	psbl
matK+trnH-psbA vs matK+atpF-atpH	W+ = 13379, W- = 6124, N = 197, p <= 5.995e-06	matK+trnH-psbA > matK+atpF-atpH
matK+trnH-psbA vs matK+psbK-psbl	W+ = 16218, W- = 3285, N = 197, p <= 7.1e-16	matK+trnH-psbA >> matK+psbK-psbl
		matK+trnH-psbA > matK+atpF-atpH+psbK-
matK+trnH-psbA vs matK+atpF-atpH+psbK-psbI	W+ = 13179, W- = 6324, N = 197, p <= 1.894e-05	psbl

Table 4. Wilcoxon signed rank tests of inter-specific divergence among loci.

	Wilcoxon Signed-Ranks Test -	
	Intraspecific pair-distances	
matK vs trnH-psbA	W+ = 298, W- = 605, N = 42, p <= 0.05574	matK < trnH-psbA
matK vs atpF-atpH	W+ = 334, W- = 162, N = 31, p <= 0.09384	matK = atpF-atpH
matK vs psbK-psbl	W+ = 299, W- = 229, N = 32, p <= 0.5189	matK = psbK-psbl
trnH-psbA vs atpF-atpH	W+ = 340, W- = 95, N = 29, p <= 0.008339	trnH-psbA > atpF-atpH
trnH-psbA vs psbK-psbl	W+ = 375, W- = 121, N = 31, p <= 0.01318	trnH-psbA > psbK-psbl
atpF-atpH vs psbK-psbl	W+ = 89, W- = 142, N = 21, p <= 0.3662	atpF-atpH = psbK-psbl
4 loci vs matK+trnH-psbA	W+ = 450, W- = 981, N = 53, p <= 0.01898	4 loci < matK+trnH-psbA
4 loci vs matK+trnH-psbA+atpF-atpH	W+ = 486, W- = 945, N = 53, p <= 0.04263	4 loci < matK+trnH-psbA+atpF-atpH
	W+ = 319, W- = 1007, N = 51, p <=	
4 loci vs matK+trnH-psbA+psbK-psbI	0.001283	4 loci < matK+trnH-psbA+psbK-psbl
4 loci vs matK+atpF-atpH	W+ = 923, W- = 508, N = 53, p <= 0.06687	4 loci = matK+atpF-atpH
4 loci vs matK+psbK-psbl	W+ = 901, W- = 530, N = 53, p <= 0.1015	4 loci = matK+psbK-psbl
4 loci vs matK+psbK-psbI+atpF-atpH	W+ = 906, W- = 525, N = 53, p <= 0.09256	4 loci = matK+atpF-atpH+psbK-psbl
matK+trnH-psbA vs matK+trnH-psbA+atpF-atpH	W+ = 810, W- = 271, N = 46, p <= 0.003294	matK+trnH-psbA > matK+trnH-psbA+atpHF
		matK+trnH-psbA > matK+trnH-psbA+psbK-
matK+trnH-psbA vs matK+trnH-psbA+psbK-psbI	W+ = 833, W- = 392, N = 49, p <= 0.02864	psbl
	W+ = 924, W- = 252, N = 48, p <=	
matK+trnH-psbA vs matK+atpF-atpH	0.0005795	matK+trnH-psbA > matK+atpF-atpH
matK+trnH-psbA vs matK+psbK-psbI	W+ = 854, W- = 371, N = 49, p <= 0.01652	matK+trnH-psbA > matK+psbK-psbI
	W+ = 1068, W- = 363, N = 53, p <=	matK+trnH-psbA > matK+atpF-atpH+psbK-
matK+trnH-psbA vs matK+atpF-atpH+psbK-psbI	0.001832	psbl

Table 5. Wilcoxon signed rank tests of intra-specific difference among loci.

In a multi loci approach for DNA barcoding purposes, the highest mean of inter-specific variability was achieved by *matK* combined with *trnH-psbA* and *atpF-atpH* whereas the highest mean of intra-specific distances were given by combining *matK* with *trnH-psbA* (Table 3). Wilcoxon statistical rank tests showed the combination *matK* + *trnH-psbA* having the highest inter-specific pair-distances (Table 4). They revealed also that all the combinations including *trnH-psbA* had a higher intra-specific variability than combinations without it (Table 5).

*Distribution of distances.* Accuracy of each DNA barcode was assessed by looking at the distribution of inter- and intraspecific K2P distances to infer the barcoding gap. Distributions were similar for each single potential barcode with two peaks of inter- and intraspecific variability that could be distinguished (Figure 2).

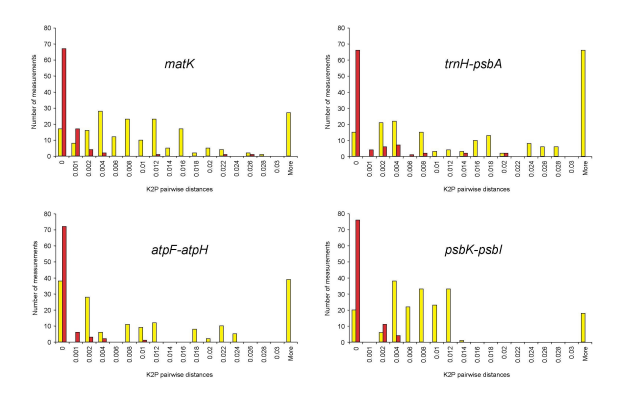


Figure 2. Relative distributions of inter-specific divergence between con-generic species (yellow) and intra-specific K2P distances (red) for four single loci: *matK*, *trnH-psbA*, *psbK-psbI* and *atpF*-

*atpH*. Barcoding gaps were assessed with Median tests and Wilcoxon Two-Sample tests, and all were highly significant (p<0.0001).

Each distribution also showed a slight overlap between intra- and inter-specific distances, but to a lesser extent for *matK* and *trnH-psbA*. Combining the different loci showed distributions with a slight decrease of this overlap (Figure 3).

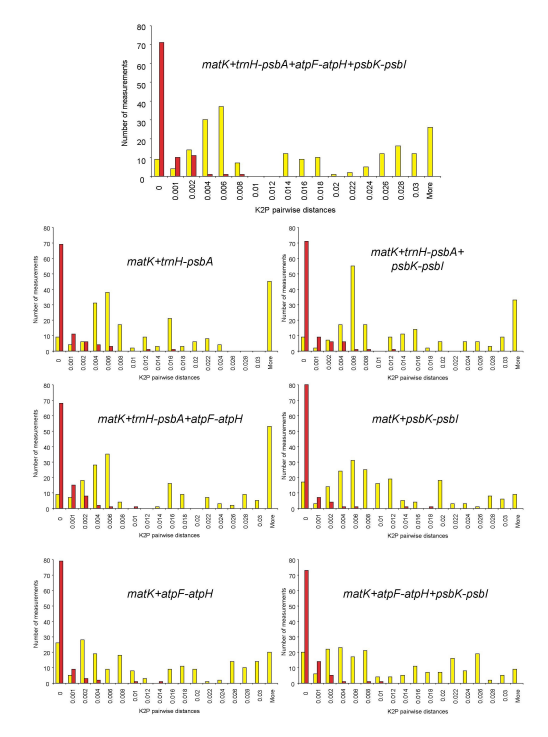


Figure 3. Relative distributions of inter-specific divergence between con-generic species (yellow) and intra-specific K2P distances (red) for 7 different combinations keeping *matK* for each. Barcoding gaps were assessed with Median tests and Wilcoxon Two-Sample tests, and all were highly significant (p<0.0001).

Two clear peaks were still distinguishable and a slight overlap still occurred between low classes of intra- and inter-specific distances, but the overlap observed was less than that

for the single locus approach. These observations were confirmed by median and Wilcoxon two samples statistical tests differentiating the medians for the former and the medians plus the distributions between the inter- and intra-specific distances for the latter. For each distribution, Median and Wilcoxon two sample tests were significant (Table 6). In a single locus approach, the highest significances were given by *matK* and *psbK-psbI*. Combining the loci made the significance increasing with the highest significance given by the combination matK+trnH-psbA+psbK-psbI.

K2P distributions	median test	Wilcoxon Two Sample Test
matK	#A = 199 #B = 93, Median = 0.00524, p <= 1.11e-26	#A = 200 #B = 93, W = 6020.5, p <= 9.314e-30
trnH-psbA	#A = 194 #B = 90, Median = 0.00799, p <= 1.11e-22	#A = 194 #B = 90, W = 5634, p <= 6.125e-29
atpF-atpH	#A = 168 #B = 84, Median = 0.00216, p <= 1.52e-23	#A = 168 #B = 84, W = 5526, p <= 8.996e-21
psbK-psbl	#A = 194 #B = 91, Median = 0.00509, p <= 1.44e-29	#A = 194 #B = 91, W = 5333, p <= 2.524e-32
4 loci	#A = 206 #B = 95, Median = 0.00608, p <= 1.23e-28	#A = 206 #B = 95, W = 5507, p <= 2.394e-36
matK+trnH-psbA	#A = 206 #B = 95, Median = 0.00648, p <= 8.07e-28	#A = 206 #B = 95, W = 5675, p <= 4.825e-35
matK+trnH-psbA+atpF-		
atpH	#A = 206 #B = 95, Median = 0.00574, p <= 5.11e-29	#A = 206 #B = 95, W = 5642.5, p <= 2.711e-35
matK+trnH-psbA+psbK-		
psbl	#A = 206 #B = 95, Median = 0.00676, p <= 5.11e-29	#A = 206 #B = 95, W = 5540, p <= 4.338e-36
matK+atpF-atpH	#A = 206 #B = 95, Median = 0.00401, p <= 1.2e-26	#A = 206 #B = 95, W = 6318, p <= 2.802e-30
matK+psbK-psbl	#A = 206 #B = 95, Median = 0.00607, p <= 8.07e-28	#A = 206 #B = 95, W = 6064, p <= 4.064e-32
matK+atpF-atpH+psbK-		
psbl	#A = 206 #B = 95, Median = 0.00493, p <= 2.92e-28	#A = 206 #B = 95, W = 6026.5, p <= 2.151e-32

Table 6. Median and Wilcoxon two sample statistical tests applied to the distributions of intraand inter-specific K2P distances for each potential DNA barcode.

*Species identification*. The performance of each DNA barcode in identifying and delineating species was assessed by the percentage of monophyletic species recovered by MP and UPGMA analyses (Table 7). Because *trnH-psbA* and *atpF-atpH* were highly variable and their alignment showed many indels, MP analyses were performed with and without coding the gaps as 5<sup>th</sup> state to infer whether this information could be useful for barcoding purposes. The highest values of species monophyly were obtained from UPGMA reconstruction. The UPGMA analysis of *trnH-psbA* gave 90.3% of species monophyletic but only 77.4% supported by BS>50%. Although *matK* and *psbK-psbI* 

grouped 87.5% of the species under UPGMA reconstruction, they gave 78.1% of monophyletic species with a BS>50%, a value higher than *trnH-psbA*. *MatK* showed the best percentage of species correctly identified using MP reconstruction. Coding the gaps as 5<sup>th</sup> state in the MP analysis did not significantly affect the results obtained for *matK* and *psbK-psbI*, but it increased the percentages of species correctly identified by 6% and 7% given by the more variable *atpF-atpH* and *trnH-psbA*, respectively. In a multi-loci approach, it is noteworthy that combining all potential barcodes did not result in 100% monophyly for species whatever the reconstruction method. Each barcode failed in grouping the two different species of *Faurea*. That can be done by using the intergenic locus *atpF-atpH* and by coding the gaps in the matrix as 5<sup>th</sup> state of character, but this decreases the total percentage of monophyletic species. In a multi-loci approach, combining *matK* and *psbK-psbI* gave the highest percentage of monophyletic species (Table 7).

	UPGMA	MP	MP+5th state character
trnH-psbA	90.3 (77.4)	71 (71)	77.4 (74.2)
matK	87.5 (78.1)	75 (75)	75 (75)
psbK-psbl	87.5 (78.1)	62.5 (68.8)	53.1 (53.1)
atpF-atpH	82.8 (69)	65.5 (65.5)	72.4 (69)
matK+psbK-psbI	93.8 (87.5)	81.3 (81.3)	59.4 (56.3)
matK+trnH-psbA+psbK-psbI	93.5 (90.3)	87.1 (87.1)	80.6 (80.6)
matK+atpF-atpH+psbK-psbI	93.1 (86.2)	86.2 (86.2)	82.8 (82.8)
matK+trnH-psbA+atpF-			
atpH+psbK-psbI	92.9 (89.3)	85.7 (85.7)	82.1 (82.1)
matK+trnH-psbA	90.3 (87.1)	83.9 (83.9)	77.4 (77.4)
matK+atpF-atpH	89.7 (82.8)	79.3 (79.3)	79.3 (79.3)
matK+trnH-psbA+atpF-atpH	89.3 (85.7)	82.1 (82.1)	82.1 (82.1)

Table 7. Proportion (%) of monophyletic species (with BS > 50% in brackets) recovered with UPGMA and MP analyses with gaps not coded and coded as a fifth character state.

*Coalescence*. The accuracy of the DNA barcode can be assessed by evaluating the ability of each candidate to give genetic clusters that are derived from an independent coalescence process and that corresponds to a recognized taxonomic species . The highest number of genetic clusters corresponding to taxonomic species was given using the UPGMA trees. Transforming MP trees by NPRS for coalescence analysis gave half the genetic clusters corresponding to taxonomic species compared to the UPGMA trees (Table 7). In a single barcode approach, *matK* gave the highest numbers of genetic clusters corresponding to taxonomic species (Table 7). In a single barcode approach, *matK* gave the highest numbers of genetic clusters corresponding to taxonomic species (Table 8). When *matK* was combined with *psbK-psbI* the value increased from 22 to 23 genetic clusters corresponding to recognized species. Molecular evolutionary rates of both *matK* and *psbK-psbI* showed higher abilities to differentiate independently evolving entities corresponding to taxonomic species than the high variable *trnH-psbA* and *atpF-atpH*.

	UPGMA	MP	Nos. of potential genetic clusters
matK	22	11	32
psbK-psbl	20	15	32
atpF-atpH	18	12	29
trnH-psbA	16	12	31
matK+psbK-psbI	23	8	32
matK+atpF-atpH+psbK-psbl	20	4	29
matK+atpF-atpH	20	6	29
matK+trnH-psbA+psbK-psbI	3	7	31
matK+trnH-psbA+atpF-atpH+psbK-			
psbl	3	1	28
matK+trnH-psbA	3	8	31
matK+trnH-psbA+atpF-atpH	3	5	28

Table 8. Coalescence analyses indicating the number of independent genetic clusters corresponding to taxonomically recognized species.

Our results showed that combining *matK* to *trnH-psbA* and *psb-psbI* can slightly increase its performance in identifying species. However we still support the conclusion of Lahaye et al., i.e. that *matK* should be used for DNA barcoding of plants in a single locus approach and that case-by-case additional barcodes are developed for problematic groups.

### **Literature Cited**