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DNA barcoding identifies cryptic animal tool materials

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Some animals fashion tools or constructions out of plant materials to aid foraging, reproduction, self-maintenance, or protection. Their choice of raw materials can affect the structure and properties of the resulting artifacts, with considerable fitness consequences. Documenting animals' material preferences is challenging, however, as manufacture behavior is often difficult to observe directly, and materials may be processed so heavily that they lack identifying features. Here, we use DNA barcoding to identify, from just a few recovered tool specimens, the plant species New Caledonian crows (Corvus moneduloides) use for crafting elaborate hooked stick tools in one of our long-term study populations. The method succeeded where extensive fieldwork using an array of conventional approaches-including targeted observations, camera traps, radiotracking, bird-mounted video cameras, and behavioral experiments with wild and temporarily captive subjects-had failed. We believe that DNA barcoding will prove useful for investigating many other tool and construction behaviors, helping to unlock significant research potential across a wide range of study systems.

animal construction behavior \mid DNA barcoding \mid nest building \mid New Caledonian crow \mid tool use

here is increasing interest in the plant materials selected by nonhuman animals to manufacture foraging tools and constructions (1, 2). Animals' raw-material preferences can affect the structural and functional properties of artifacts, and, in some cases, appear to be socially transmitted, contributing to rudimentary material "cultures" (3–5). Two complementary approaches are available for identifying plant materials used by wild animals: direct observation of manufacture behavior ("animal-centered") and examination of artifacts in isolation from the behavior that created them ("artifact-centered"). The latter, adopted by necessity in archaeology, is particularly useful when animals cannot be habituated or are otherwise difficult to observe, but can present considerable challenges. Artifacts are often heavily processed (lacking features that aid identification, such as leaves or flowers), may be physically distanced from the raw materials from which they were produced (because the animal transported them), and may comprise a complex assemblage of materials from different sources (such as in bird nests). In these cases, material identification has so far relied on expert knowledge, which may be difficult and expensive to acquire (6). Here, we demonstrate that DNA barcoding-the use of standardized DNA regions to identify organic material to species level (7)-provides a robust, cost- and time-efficient solution to these problems.

New Caledonian crows (*Corvus moneduloides*) are renowned for their ability to manufacture complex foraging tools (8). When making a hooked stick tool, they select a forked plant stem, remove a suitable branch, trim off any leaves and twiglets, and often refine the tool by sculpting the remains of the nodal joint into a neat terminal hook, stripping bark near the functional end, and bending the tool shaft (9). These processing steps substantially alter the appearance of the plant material (Fig. 1*A*). Importantly, properties of the raw material affect the morphology of the resulting tools, which in turn affects foraging efficiency (9–11). New Caledonian crows are highly selective when choosing plants for hooked stick tool manufacture: we recently discovered that three study populations target different species despite living just a few kilometers apart (12). While we managed to identify raw materials at two sites (site-1 and site-2), we failed at the third (site-3), even after employing a wide range of well-established field methods aimed at observing tool manufacture directly (Fig. 1 *B* and *C* and *SI Appendix*).

Given the importance of identifying the crows' preferred plant species at site-3 for our overall research program, we tried an innovative artifact-centered approach (Fig. 1D). We extracted DNA from seven hooked stick tools recovered at the site during 2016-2017 and amplified two DNA barcoding regions: trnL-UAA [~500 bp (13)] and internal transcribed spacer (ITS) [~600 bp (14)]. All samples produced identical haplotypes. Comparison against the National Center for Biotechnology Information (NCBI) Nucleotide nonredundant database indicated the tool samples belonged to the family Sapotaceae, most likely the genus Mimusops or Manilkara (trnL: >98% identity; ITS: >95% identity). The ITS region exhibited greater resolution and indicated Mimusops elengi as a candidate (96 to 99% identity). With a putative source identified, we collected reference leaf samples of M. elengi and Planchonella cinerea, the only closely related species known to occur locally, and analyzed them using the same method. The tool samples and M. elengi reference samples produced identical haplotypes for both DNA barcodes. Furthermore, the maximum likelihood phylogenetic tree for ITS produced well-supported lineages that clustered tool samples and M. elengi reference samples together, within a wider clade of non-New Caledonian M. elengi sequences (Fig. 1 D, iii). P. cinerea reference samples clustered within the Sapotaceae, but outside the Minusops genus, as expected, confirming that the crow tools from site-3 were made from M. elengi. We subsequently verified that wild-caught, temporarily captive New Caledonian crows readily manufacture hooked stick tools from this material (this work was conducted at site-1, since birds from site-3 proved too difficult to work with in field aviaries).

The use of DNA barcoding has led to an important breakthrough for our research program. Reliable raw-material identification is key to uncovering the drivers of the striking regional divergence we observed in an important aspect of New Caledonian crows' hooked stick tool-making behavior (15). Specifically, with a set of three study populations established, and the ability to conduct rapid surveys across additional replicate sites, it will

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С Initial investigations

2012

Collection

discarded tools

begins at site-3

with help from

local residents.

Recovery of

Animal-centered

D

Artifact-centered

Raw material of NC crow tools at site-3 is shown to differ from that at site-2, but field observations, aviary experiments and camera traps fail to identify the plant species

Bird-mounted video loggers

Miniature cameras are attached to NC crows at site-3 to document foraging behavior - no tool manufacture recorded.

0

Research focus on site-2

Work focuses on site-2 where wild and temporarily captive NC crows make hooked stick tools from the non-native perennial shrub D. virgatus

Aviary experiments

Temporarily captive NC crows from site-3 are presented with candidate plant species M. elengi (along with other locally sourced plants) - low task engagement.





Behavioral confirmation

Temporarily captive NC crows from site-1 are found to readily make hooked stick tools from supplied M. elengi forks and use them for extracting bait from a



Fig. 1. Identifying the raw material used by wild New Caledonian (NC) crows for manufacturing hooked stick tools. (A) NC crow holding a hooked stick tool manufactured from Desmanthus virgatus at site-2. (B) Satellite photograph showing study sites on the west coast of Grande Terre, New Caledonia. Map image credit: © 2019 Google Maps/CNES/Airbus, TerraMetrics, Data SIO, NOAA, U.S. Navy, NGA, GEBCO. (C) Timeline of animal-centered approaches employed while attempting to identify the tool material used at site-3, with short explanations for their limited success. (D) Timeline of the artifact-centered, DNA barcoding approach which ultimately led to successful material identification. (i) Hooked stick tool recovered at site-3 (5 × 5 mm background). (ii) M. elengi trees, with close-up of a forked terminal branch, which would be suitable for tool manufacture (shape [in white] of potential tool overlaid on image). (iii) Simplified maximum-likelihood ITS phylogenetic tree detailing clustering of tool samples and a subset of reference data (data for the full tree are deposited in Dryad). Symbols denote samples from M. elengi (diamonds), crow tools (triangles), Mimusops spp. (squares; from top: M. zeyheri, M. caffra, M. comorensis, M. obovata, M. kummel, M. sp., M. coriacea, M. lecomtei, M. perrieri, M. membranacea), and Sapotaceae spp. (circles; from top: Tieghemella heckelii, Autranella congolensis, Labourdonnaisia spp., Faucherea spp., Labramia spp., Manilkara spp., Baillonella toxisperma, Vitellaria paradoxa, Vitellariopsis spp. and two samples from P. cinerea collected in this study). OG is an outgroup (Sarcosperma laurinum). Filled symbols denote samples collected from the study site, open symbols denote those from outside of New Caledonia accessed through GenBank, and hatching denotes a species potentially introduced to New Caledonia (but not sampled there). Asterisks indicate >70 bootstrap support, and the scale bar shows substitutions per site. A single Manilkara hexandra sequence from GenBank (JX856473), which resolved with Mimusops, is omitted here, as it was most likely a misidentification (all other sequences from this genus clustered elsewhere, as shown). (iv) "Refit" of a hooked stick tool made from M. elengi material by a temporarily captive crow from site-1: the tool is displayed along with stem and plant debris which were discarded by the crow during the manufacture process (scale in millimeters).

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now be possible to examine whether crows' raw-material preferences are related to environmental variation in the availability of different plant species and foraging opportunities; for example, birds may simply use a locally common tool material, or they may choose a material that is mechanically well suited to targeting local prey resources (in fact, DNA barcoding could potentially also be used to determine prey identity, using trace DNA left on tool tips). Such ecological work is of critical importance for informing our understanding of technological (cultural) evolution in this model species (16).

Perhaps more importantly, we believe that genetic approaches will be useful for many other study systems where traditional observational methods are not feasible or would cause undue disturbance, and/or where plant materials are routinely transported or heavily modified. For example, DNA barcoding could facilitate the identification of raw materials used by chimpanzees (Pan troglodytes) for making tools for termite fishing and other tasks, avoiding the need to regrow plants and curate herbarium specimens (4-6). We also envisage studies that identify-from small samples-individual components of complex composite structures such as bird nests and bowers, replacing time-consuming destructive investigation. There are also exciting opportunities for further methodological refinement. For example, it should be possible to recover DNA from artifacts held in museum and research collections, potentially enabling productive retrospective analyses (17). Furthermore, targeting more variable regions of the genome, such as single-nucleotide polymorphisms, could help identify, more precisely, where an animal collected plant materials (18), providing valuable information on search and transport costs. Reliable, cost- and time-efficient raw-material identification will

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facilitate detailed investigation of how animals source plant materials from the environment, and how the properties of these materials affect the function of the resulting artifacts.

Materials and Methods

Methods are summarized in the main text and Fig. 1 *C* and *D*. *SI Appendix* contains extended methods, detailing our unsuccessful animal-centered (observation-focused) approaches and successful artifact-centered (DNA barcoding) approach. The latter includes DNA sequencing of samples, the search strategy and recovery of sequences from the NCBI Nucleotide non-redundant database, and the subsequent phylogenetic analyses. All sequences produced in this study are deposited in GenBank, and all data for the full maximum-likelihood phylogenetic trees (ITS and *trnL*) that were used for raw-material identification are deposited in Dryad (for details, see *Data Availability*).

Data Availability. DNA sequence data have been deposited in GenBank (MT366813–MT366824 and MT366951–MT366962). Sequence alignments and resulting phylogenetic trees are deposited in Dryad (https://doi.org/10.5061/dryad.d7wm37q1v) (19), including GenBank accession numbers.

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Supplementary Information for

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This PDF file includes:

Extended methods SI References

Extended methods

Animal-centered approaches (see Fig. 1C in the main text)

Initial investigations. In September and October 2012, five New Caledonian (NC) crows were caught at site-3 (for study site locations, see Fig. 1*B*), housed in temporary on-site field aviaries, and provided with assorted, locally sourced plant materials and extraction tasks (logs with drilled holes baited with small pieces of meat). None of these subjects manufactured hooked stick tools. For full details and methodology, see (1), and for husbandry protocols, see (2).

Between September and November 2012, a baited setup was presented to wild NC crows at site-3. The setup consisted of either an extraction log (presented on the ground or raised on wooden tripods), or a raised platform with a water bath and/or food bait (meat, fruit, or cat food). The setup was monitored using an autonomous camera trap (Bushnell Trophy Cam, Trophy HD, model no: 119447) that recorded 60-second-long videos at each trigger. NC crow visits to the setup were captured in 345 videos, and tool use was seen in 153 of these. Four visits contained tool manufacture episodes, in which crows sourced plant material from small shrubs immediately beside the extraction log (the plant species could not be identified). We were able to confirm that three of the tools manufactured were non-hooked sticks, but for the fourth, it was impossible to determine tool type. In all four manufactures, however, crows performed only minimal processing (e.g., removing leaves and twiglets) after detaching material from the plant and before using it as a tool. We did not observe any of the processing that is commonly associated with the manufacture of hooked stick tools, such as sculpting of the tool's functional end, bark stripping, or tool-shaft bending (3). A similar setup was used at site-3 between September and November 2013, but only a baited, raised platform was presented (no extraction logs). A total of 3,343 videos were recorded by autonomous camera traps but, since tool use was only observed in conjunction with an extraction log in 2012, we have not formally reviewed these videos.

In addition to these recordings, during occasional targeted observation sessions in a private garden at site-3 (see below), a wild NC crow was once seen manufacturing a stick tool (likely non-hooked) in a large tree, which was later identified as a Banyan (genus *Ficus*) growing around an established *Cerbera manghas* tree.

Bird-mounted video loggers. In October 2012, we fitted three NC crows at site-3 with miniature video loggers (two of these birds had been held in temporary captivity for the aviary experiments described above). Following standard protocols, the birds were sexed morphologically [all males; (4)] and their age was estimated by gape coloration [one immature and two adults; (5)]. For more details on video logger specifications and deployment methods, see (6, 7). We managed to recover two loggers: one unit was removed from the bird upon re-capture, but was badly damaged and no video footage was salvaged; the other unit, which detached as planned, had recorded video footage, and although one instance of tool use was recorded (probing with a tool in a hole in a tree), no clear view of the tool itself was captured, and it could not be established if the tool was hooked or non-hooked. No tool-manufacture episodes were recorded.

Field experiments. A baited extraction log was presented to wild NC crows at site-3 in August and September 2017 and monitored by either human observers in nearby hides or autonomous camera traps (as above). Locally sourced plant material was occasionally presented alongside the log. During a total of 12.1 hours (over nine non-consecutive days) of in-person monitoring, we observed four NC crow visits to the log and one instance of tool use. Of the 97 camera-trap videos which recorded NC crow visits, 54 contained tool use. Where crows were recorded sourcing a tool, they appeared to take sticks from the ground nearby and no tool manufacture was observed. Visits to the log were rare, presumably due to the good availability of food in surrounding gardens (e.g., chicken feed, cat food, bones left out for dogs, and coconuts cracked open by residents to feed wild birds) and the presence of stray cats.

Aviary experiments. Between September and October 2018, four NC crows (all males; one juvenile, one immature, and two adults) were trapped at site-3 and kept in temporary captivity in field aviaries (as above). To facilitate birds' habituation to aviary confinement and experimental procedures, and to avoid corrupting potential local NC crow "cultures" [captive and wild birds may socially learn tool behaviors from each other that they do not naturally express; (1, 8)], we always erect field aviaries near the site of capture. After careful consideration, we made an exception on this occasion. Due to lack of suitable space and the large number of free-roaming cats and dogs at site-3, we transported crows to field aviaries at nearby site-2. In order to prevent any potential visual interaction between captive site-3 subjects and local wild crows from site-2, all aviaries were covered in a double layer of UV screen for the duration of the experiments (allowing sufficient light in but preventing a clear view). All subjects were released at site-3 where they had been caught.

Subjects were presented with a choice of plant materials for tool manufacture and a food extraction task both in their housing aviaries and in the experimental chamber. Informed by our DNA barcoding analyses of tools recovered at site-3 (see below), plants presented always included *Mimusops elengi* and, when sufficient source material was available (6 of 15 presentations), *Planchonella cinerea*, the only closely related species known to occur locally. Other plant species presented could not be identified reliably but were sourced locally and had a branching structure that was judged suitable in principle for hooked stick tool manufacture. Plants were displayed in pseudo-randomized positions standing upright on presentation logs either at ground level or raised to ~1 m on wooden tripods, to better approximate how NC crows may encounter material in larger tree species (like *M. elengi* and *P. cinerea*) in the wild. All subjects were reluctant to engage with extractive foraging tasks, and when they did, were only observed to forage with supplied sticks and non-hooked stick tools they had manufactured from *Desmanthus virgatus*. One individual briefly used a small twig of *M. elengi* to probe in its metal ID ring.

NC crows normally habituate very quickly to our field aviaries, and readily participate in behavioral experiments. On this occasion, our impression during husbandry activities and experimental work was that birds were unsettled. We suspect that transportation between sites, and unfamiliarity with site-2 (including the calls of unfamiliar wild crows visiting the aviaries), may have contributed to the subjects' lack of engagement with tasks. These observations caution that NC crows should not be routinely translocated for behavioral experiments.

VHF radio-tracking. Six NC crows (two females, four males; one juvenile, two immatures, and three adults) were caught at site-3 between September and October 2018 and fitted with VHF radio-tags (picopips, Biotrack Ltd., UK; tag mass: 1.8–2.1 g). Tags were attached with epoxy adhesive to the base of the birds' two central tail feathers, and the antenna was held in place along the shaft of one of these with two to three knots of dental floss fixed with superglue. Four of the six individuals released with tags had previously been held in field aviaries and tested in behavioral experiments (see above).

Following release, crows were allowed to habituate to the tag for at least one full day before tracking commenced. The order in which we searched for tagged crows each day was pseudo-randomized. At the start of each tracking session, a sweep over the entire study site was carried out by car and, if an individual's signal was detected, we attempted to get closer to that individual on foot for focal following. We confirmed a bird's identity by either reading its ID ring or, where this was impossible, establishing a line of sight coupled with a strong, directional radio signal. The positions of any individuals which could not be directly observed were determined by cross-triangulation where possible.

We spent a total of 48.25 hours radio-tracking over 10 days between 30 September and 15 October 2018. As birds were trapped and released throughout the study period, tracking effort varied substantially each day (median 2.67 hours per session, range 0.17–6.17 hours). Radio-tags were verified to be working on four of the six crows. One individual was never detected after release, possibly due to tag malfunction or the bird leaving the study area, and another tag was recovered detached in a garden seven days after the bird's release (the bird was subsequently seen alive and well). The remaining birds proved difficult to find and follow: only for 23 of a total of 199 radio fixes were we able to home-in on the location of the tagged bird and achieve a sighting. This was due to two main reasons: accessing many private properties in the study area proved difficult due to the presence of unrestrained guard dogs; and crows regularly crossed open sections of the river behind the residential area (20–100 m wide) and spent a large proportion of their time on small islands in the mangrove swamp of the river delta, which were inaccessible to us during the study period.

On one occasion, a crow was recorded feeding on *M. elengi* fruits, but no tool use or tool manufacture were observed throughout our radio-tracking study. After our tracking study had ended, we were able to briefly visit one of the small islands in the mangrove swamp, where we observed a radio-tagged crow (ID ring CE8) holding a tool as it flew through the canopy. We were unable to determine the tool type from this brief sighting.

Artifact-centered approach (see Fig. 1D in the main text)

Tool collection. Between 2012 and 2017, a total of 105 tools were collected in a private garden at site-3 by both researchers from our team and local residents (1). Tool collection was largely opportunistic, as NC crows visiting a bird water bath in one of the gardens would regularly leave tools behind after drinking. The "startle" method (9) was also used at least once, to distract a crow observed holding a tool. The time taken until tool recovery ranged from a few minutes (for the startle method) to possibly several days (for tools recovered opportunistically). In 2016 and 2017, tools were stored in re-sealable plastic freezer bags as soon as possible following their discovery, along with sufficient silica gel powder to cover them [n = 19; (10)]. Stored tools were transported to the UK for further analysis and archiving in the team's NC crow tool collection.

The recovered tools were all of broadly similar appearance [see Fig. 1*D* in the main text, and Figure 1C in (1)], exhibiting features characteristic of hooked stick tools collected elsewhere (a terminal hook, curved tool shaft, and bark stripped from the functional end). Most tools showed signs of foraging wear around the hook, but presented no morphological features that would have enabled reliable plant species identification [see main text, and (3, 9, 11)]. Having never observed hooked stick tool manufacture by wild crows at site-3, it was also not possible for us to recover and examine plant debris discarded by crows during raw-material processing. That said, some morphological features were preserved on a small subset of tools, including bark along the tool shaft with scars suggesting an alternate leaf arrangement and, less frequently, a small amount of leaf material or a small leaf bud on the non-functional end of the tool. While the appearance of tools suggested that they all originated from the same plant source, we were unable to identify the species through visual examination alone, despite also consulting several expert botanists [present study, and see also (1)].

DNA barcoding identifies candidate species. In May 2018, we selected for further analysis a subset of seven tools from the 19 tools collected at site-3 between 2016 and 2017 (see above). The first five tools were selected specifically to maximize our chances of successful DNA recovery, as they had fragments of bark remaining on the tool shaft. Once we had confirmed that DNA extraction was possible from these tools, we randomly selected an additional two tools with little bark remaining on the tool shaft, to diversify our sample. Given that visual inspection suggested all tools were of common material origin, and in order to preserve as much of our tool collection as possible for other research (since sampling was destructive), we limited our analysis to this subset of seven tools. Two subsamples were taken from each tool, and these were homogenized in 2 ml tubes with two tungsten beads using the FastPrep-24[™] 5G Benchtop Homogenizer for up to one minute, or with QIAGEN Tissuelyser II for 6 minutes at 25 Hz. DNA extraction then proceeded according to the DNeasy Plant Mini Kit (QIAGEN, Manchester UK), except the lysis step was extended to 60 minutes.

For species identification, we amplified ~500 bp regions of the chloroplast *trnL-UAA* intron [*trnLc-d*; (12)] and a ~600 bp nuclear ribosomal region containing the internal transcribed spacer regions ITS1 and ITS2 [ITS5p-8p; (13)]. Amplification was in 20 µl reactions using approximately 100 ng of genomic DNA, 10 x reaction buffer (Bioline BIOTAQ Reagent Buffer), 30 nmol MgCl₂, 4 nmol dNTPs, 7.5 pmol primers, and Bioline BIOTAQ DNA polymerase (0.5 units). Negative controls and DNA extraction blanks were included in each PCR to check for potential contamination. Thermocycling was performed on a Bio-Rad Tetrad 2 (Bio-Rad, Hamburg, Germany) under the following conditions. For *trnL*: initial denaturation (94 °C for 4 min), followed by 35 cycles of denaturation (94 °C for 45 s), annealing (55 °C for 45 s) and extension (72 °C for 120 s), and a final extension of 10 min at 72 °C; and for ITS: initial denaturation (94 °C for 2 min), followed by 30 cycles of denaturation (94 °C for 60 s), annealing (55 °C for 60 s) and extension (72 °C for 90 s), and a

final extension of 5 min at 72 °C. Two independent PCRs were carried out for each subsample. Successfully amplified PCR products were cleaned using ExoSap-IT[©] (USB Corporation, Cleveland, Ohio, USA). Sequencing was resolved on an AB 3730xl Sequencer at Edinburgh Genomics (Scotland, UK). Sequences were checked and edited with reference to chromatograms using Sequencher v 5.4.1 (Gene Codes Corporation, Ann Arbor, MI, USA).

Sequence data were obtained for all seven crow tools (14 subsamples, with 2 replicates per tool), yielding identical haplotypes for all samples for both *trnL* and ITS. Sequences obtained for all tool samples were lodged with GenBank under accession numbers MT366813–MT366819 (*trnL*) and MT366951–MT366952, MT366955–MT366959 (ITS). There was no amplification in negative controls. The resulting sequence haplotype for each region was used for performing BLASTn searches against the National Center for Biotechnology Information Nucleotide nonredundant database (4 July 2018) to obtain a putative identification. Searches for both *trnL* and ITS indicated that the unknown plant material belonged to the family Sapotaceae, most likely the genus *Mimusops* or *Manilkara (trnL*: >98% identity; ITS: >95% identity). Specifically, the best 100 matches (excluding environmental samples) for *trnL* all possessed 98–99% identity and belonged to the Sapotaceae, with the most likely match being *Mimusops capuronii*, followed by *Manilkara zapota*. The ITS region showed similar results with all species in the top 100 matches belonging to the Sapotaceae and the unknown samples exhibiting 96–99% identity to *M. elengi*, followed by several species within this and one other genus (*M. caffra, M. zeyheri, M. comorensis, M. obovata, M. kummel*, and *Manilkara hexandra*) with 95–96% identity.

Based on these results, we carried out a qualitative comparison of the remaining tools with preserved samples of *M. elengi* (as the closest match and the only species of *Mimusops* native to New Caledonia) in three herbarium collections – the Royal Botanic Garden Edinburgh, Scotland [accessed physically and online; (14)], the Muséum National d'Histoire Naturelle, France [accessed online only; (15)], and the Institut de Recherche pour le Développement, New Caledonia [accessed online only through JSTOR Global Plants; (16)]. Morphological features preserved on a small subset of the tools (see above) were also observed in the terminal twigs of some *M. elengi* herbarium samples, but it was not possible at this stage to definitively confirm a match between these samples and the tools.

Material identity confirmed. In September 2018, we located and examined *M. elengi* trees at site-3 and in the surrounding area. Some terminal branches of these trees indeed appeared suitable for hooked stick tool making and possessed structural features we believe to be important in other plants used by NC crows [e.g., *D. virgatus*, Fabaceae; (2, 3, 17)]. These features include an acute branching angle less than 45°, a strong but flexible tool shaft, and a hook shaft of similar diameter to the tool shaft. In particular, we found "reiterations" – young epicormic shoots emerging from mature branches that replicate a tree's overall architecture (18) – on some *M. elengi* trees that appeared very similar in structure to the *D. virgatus* stems preferred by NC crows at site-2 (3).

Following these observations, we collected reference leaf samples of *M. elengi* (n = 3) and its closest locally occurring relative, *P. cinerea* (n = 2). Leaves were collected from separate trees at or near site-3 and site-1. In March 2019, DNA extraction, PCR amplification and sequencing for the reference samples were performed exactly as described above for the tool samples. The resulting DNA sequences for the reference data were lodged with GenBank under accession numbers MT366823–MT366824 (*trnL*) and MT366953–MT366954 (ITS) for *P. cinerea*, and MT366820–MT366822 (*trnL*) and MT366960–MT366962 (ITS) for *M. elengi*. Phylogenetic trees were generated using these samples plus the top 100 BLASTn matches (excluding environmental samples). Since taxa from both Sapotaceae and Theaceae were present in the top matches for *trnL*, *Acanthogilia gloriosa* (GenBank accession number EU348374), a polemonioid Ericale was included as the outgroup for *trnL*. Since all the taxa in the top matches for ITS were from the Sapotaceae, subfamily Sapotoideae, *Sarcosperma laurinum* (GenBank accession number AM408055), which has previously been shown to be sister to the rest of the family, was included as an outgroup for ITS (19).

Sequences were aligned using the MUSCLE algorithm (20). Maximum-likelihood (ML) trees were constructed using the .pml function in the package phangorn (21), implemented in R (22). The most appropriate model of DNA substitution was selected using the AIC and the function modelTest. We used the HKY model with the proportion of invariant sites and rate variation opti-

mized using the function optim.pml. Support for the branching topology was evaluated with 1,000 bootstrap replicates. Separate trees were generated for chloroplast (*trnL*) and nuclear (ITS) regions.

Sequence comparisons revealed that M. elengi reference material produced identical sequences to the tool samples for both DNA barcodes. The ML phylogenetic tree for trnL (data deposited in Drvad: see main text) showed poor resolution at the genus or species level within the Sapotaceae family (it showed an exact match between the trnL sequence of the M. elengi reference samples and the tool material, but this haplotype was also shared with a Madagascan species, M. capuronii, the only other Mimusops species with sequence data available). The ML tree for ITS produced the most informative data, with well-supported lineages largely corresponding to species/ genera (data deposited in Dryad; see main text). The unknown plant material from tool samples clustered with sequences from species belonging to the Mimusops genus and was nested within a cluster of samples corresponding to *M. elengi*. The unknown tool samples had an identical ITS sequence to the New Caledonian *M. elengi* reference samples, and this sequence cluster was nested within a wider clade of *M. elengi* samples from Indonesia (GenBank accession number KF686246) and Thailand (GenBank accession numbers KF686245 and HF542849). Importantly, M. elengi is the only native Minusops species in New Caledonia (23, 24), adding additional confidence to the identification. One other *Mimusops* species is recorded from New Caledonia as an introduction (*M. coriacea*); this shows a clearly distinct ITS sequence (GenBank accession number KM370965), which clusters separately to the tool samples and the reference material of M. elengi. The reference samples for P. cinerea clustered within the Sapotaceae, but outside the Mimusops genus as expected. A single Manilkara hexandra sequence from GenBank (GenBank accession number JX856473) is resolved within *Mimusops* in the ITS phylogeny (separate from the tool samples and *M. elengi*), although this is most likely a misidentification, as all other sequences from Manilkara clustered elsewhere in the tree. Therefore, based on an identical sequence match to reference material gathered from the study location, the unknown plant material used by NC crows to produce hooked stick tools at site-3 is identified as *M. elengi*.

Behavioral confirmation. In August and September 2019, 13 NC crows at site-1 were held onsite in temporary captivity for behavioral experiments, following the protocols described above. Birds were initially presented with a range of locally available plant material alongside extraction tasks in both their housing aviaries and in experimental chambers. The materials included two plant species known to have been used by crows at site-1 [*Acacia spirorbis* and *Melaleuca quinquenervia*; (7)], the two species flagged by our DNA barcoding analyses (*M. elengi* and *P. cinerea*), the preferred tool material of crows at site-2 [*D. virgatus*; (1)], as well as two other locally sourced plants with a structure judged suitable for hooked stick tool making.

Two of these crows made hooked stick tools (the others either made only non-hooked stick tools or did not make tools at all in captivity) and regularly chose *M. elengi* forks (in the presence of other plant species) to make these tools, using them to extract bait. This provided the first direct observations of NC crows processing *M. elengi* forks to manufacture hooked stick tools. Importantly, these tools manufactured (in captivity) from *M. elengi* by crows from site-1 appeared morphologically very similar to the tools we had previously recovered at site-3 (see above).

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