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RESEARCH ARTICLE

Monitoring large and complex wildlife aggregations with drones

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Abstract

- 1. Recent advances in drone technology have rapidly led to their use for monitoring and managing wildlife populations but a broad and generalised framework for their application to complex wildlife aggregations is still lacking.
- 2. We present a generalised semi-automated approach where machine learning can map targets of interest in drone imagery, supported by predictive modelling for estimating wildlife aggregation populations. We demonstrated this application on four large spatially complex breeding waterbird colonies on floodplains, ranging from c. 20,000 to c. 250,000 birds, providing estimates of bird nests.
- 3. Our mapping and modelling approach was applicable to all four colonies, without any modification, effectively dealing with variation in nest size, shape, colour and density and considerable background variation (vegetation, water, sand, soil, etc.). Our semi-automated approach was between three and eight times faster than manually counting nests from imagery at the same level of accuracy.
- 4. This approach is a significant improvement for monitoring large and complex aggregations of wildlife, offering an innovative solution where ground counts are costly, difficult or not possible. Our framework requires minimal technical ability, is open-source (Google Earth Engine and R), and easy to apply to other surveys.

KEYWORDS

aerial vehicle, automated detection, bird, breeding, colonial, ecology, machine learning, waterbird

1 | INTRODUCTION

Recent advances in technology offer the potential to improve field methods for rapidly and effectively monitoring biodiversity (Pimm et al., 2015). Among these advances is the use of aerial vehicles, or drones, that can carry remote sensing instruments to capture extremely high spatial resolution imagery with temporal flexibility (Anderson & Gaston, 2013). Drones are relatively easy to use and their increasing 'off the shelf' application to wildlife research has been innovative and exciting (Chabot & Bird, 2012, 2015). There are increasing novel applications for monitoring both populations and

behaviours of different fauna, including birds (Chabot & Francis, 2016; Hodgson et al., 2018), elephants (Vermeulen, Lejeune, Lisein, Sawadogo, & Bouche, 2013), crocodiles (Evans, Jones, Pang, Saimin, & Goossens, 2016), chimpanzees (Van Andel et al., 2015) and marine mammals (Seymour, Dale, Hammill, Halpin, & Johnston, 2017).

Given the ability of drones to collect high quality data over large aggregations of wildlife, they offer an attractive opportunity for improving methods and increasing cost effectiveness of monitoring wildlife populations. The relative advantages of aerial countingboth in-air and from aerial imagery-for wildlife monitoring is long established, including reduced detection error, increased precision, reduced observer effects and retrospective analysis of data. For example aerial counting was more accurate and precise than ground counting using aerial images of penguin colonies (Fraser, Carlson, Duley, Holm, & Patterson, 1999) and geese (Boyd, 2000). Similar advantages of image-based counts over ground-based counts have been demonstrated for drone-acquired imagery too (Hodgson et al., 2018).

At large spatial scales (km) and for large aggregations (e.g. >5,000–10,000 individuals), in-air aerial surveys provide cost effective information on counts of individuals, breeding-pairs and nests (Caughley, 1977; Kingsford & Porter, 2009), although sometimes suffering high variability and imprecision (Kingsford, 1999). High-altitude imagery from aeroplanes allows large areas, if not whole aggregations, to be captured in single images (e.g. in Boyd (2000) *c*. 30 photos captured flocks of *c*. 10,000 geese). Owing to the fact that similar areas require many thousands of drone images and to the extra complexity from increased spatial resolution, drone use for monitoring wildlife aggregations (i.e. <5–10,000 individuals), though there are some notable exceptions (Afán, Máñez, & Díaz-Delgado, 2018; Chabot & Bird, 2012; Chabot, Craik, & Bird, 2015).

Manually counting targets of interest (e.g. individual animals, breeding-pairs, nests) from aerial images, regardless of capture platform, is laborious. This has driven the development of automated or semi-automated counting approaches (Chabot & Francis, 2016; Hollings et al., 2018), aided by the widespread availability of increased computing power, growing computer literacy and new methods. Current approaches typically involve spectral thresholding (Chabot & Bird, 2012; Seymour et al., 2017), point process algorithms (Descamps, Bechet, Descombes, Arnaud, & Zerubia, 2011) or combinations of spectral properties and predictive modelling (Hodgson et al., 2018). These methods rely on high contrast (i.e. dark animals on light backgrounds or light animals on dark backgrounds) and consistency in the shape and colour of the targets (Hollings et al., 2018). They are generally only applicable when the spectral and structural characteristics of the animals (in the images) are unique compared to the rest of the image (Chabot & Francis, 2016). More recently, remote sensing-based methods have been used to overcome challenges with low contrast and high variation among target objects (Afán et al., 2018; Chabot, Dillon, & Francis, 2018; Drever et al., 2015; Groom, Petersen, Anderson, & Fox, 2011; Liu, Chen, & Wen, 2015).

Despite the interest in automated methods for counting aggregations of birds, their use by ecologists and managers for monitoring complex wildlife aggregations remains limited (Chabot & Francis, 2016), with manual approaches still dominating (Buckland et al., 2012; Drever et al., 2015). There are three key reasons that have been highlighted for the disconnect between new methods and their ecological application: (a) most methods have only been demonstrated at small spatial scales relative to real-world applications (even if the number of individuals is very large) and in homogenous areas with little environmental complexity (Hollings et al., 2018); (b) ecological complexity and outcomes are not appropriately considered with respect to the mobility of individuals and variation in the types of target features of interest (Baxter & Hamilton, 2018); and (c) there is a high technical threshold for implementing most methods (Chabot & Francis, 2016).

In this paper, we develop a semi-automated framework for monitoring large complex wildlife aggregations using drone-acquired imagery. We use the case study of colonial waterbird breeding colonies because they present the key challenges currently inhibiting uptake of drone-based methods; the colonies cover large spatial extents and can have range of density of animals across these extents; there are many thousands of highly mobile individuals that cannot be contained to single drone images: the target features of interest are nests, which can exhibit significant differences in structure and colour across space and time (e.g. empty nests, adult/juvenile/chick/egg occupied nest, variable nest material, variable nest shape and arrangement); and considerable variation in background environment (mud, sand, water, live/dead vegetation). We developed a set of generalised methods, that could be transferred directly between colonies without modification, and required relatively little technical ability to apply. We captured imagery over four breeding waterbird (mostly Straw-necked Ibis) colonies in New South Wales, Australia, ranging in size from c. 20,000 to over 200,000 birds, including the largest ever waterbird colonies to be surveyed by drone. We detail flight planning, image acquisition and processing, manual and automated methods for mapping and counting nests. We include the Google Earth Engine and R code required for our analyses, along with a web-app to explore drone data, intermediate machine learning predictor and nest map layers.

2 | MATERIALS AND METHODS

Our primary motivation was mapping and counting nests for breeding colonial waterbirds, with wide applicability. The methodology needed to work on both small (c. 10,000–20,000 birds) and large (200,000+ birds) colonies and be transferable across different environments and applications, requiring limited technical modification or ability. We developed a modular approach that included: (a) drone image surveys of four large breeding colonies; (b) manual counting of nests for training and validation; (c) a machine learning mapping method to map nests from drone imagery; and (d) a predictive modelling method to estimate total nest numbers.

2.1 | Study location and bird colony details

Straw-necked Ibis *Threskiornis spinicollis* are an Australian nomadic waterbird species which form very large breeding colonies, sometimes mixed with other waterbird species, when ecological conditions are favourable (Brandis, Koeltzow, Ryall, & Ramp, 2014). We surveyed four colonies: *Merrimajeel, Zoo Paddock, Eulimbah and Block Bank* (Table 1). We surveyed the colonies at around their maximum size, determined by progression of breeding (Brandis, Kingsford, Ren, & Ramp, 2011). Straw-necked Ibis typically make their nests in flooded wetlands and floodplains, using inundated vegetation as nesting material raised above ground/water level. The

TABLE 1 Location and information on drone-surveyed waterbird colonies. All colonies were in New South Wales, Australia. Nests were manually counted from the drone-based imagery. Nest count error was calculated from in situ ground counts cross-referenced with manual nest counts from drone imagery

Location (Colony name)	Date	Approx. colony size	Manual nest count	Manual nest count error	Estimated number of birds ^a
Lachlan River (Merrimajeel)	Oct 2016	60-65 Ha	96,989	±6.1%	200-250,000
Macquarie Marshes (Zoo Paddock)	Nov 2016	60-65 Ha	20,411	±8.8%	40-50,000
Murrumbidgee River (Eulimbah)	Nov 2016	15-20 Ha	13,343	±8.4%	30-40,000
Lachlan River (Block Bank)	Sep 2017	7–10 Ha	7717	±12.1%	15-20,000

^aFrom (Lyons, Brandis et al., 2018) – the estimated number of birds incorporates site-specific information.

vegetation at the sites was dominated by Lignum shrubs Duma florulenta and Common Reed Phragmites australis. Nests can be isolated nests or 'clumped' (10-200 nests). The nests are generally round or oblong in shape, but are often irregular in large clumps, with trampled vegetation, forming a dark green to brown colour, which increasingly whitens with guano (Figure 1), until nests are abandoned either when offspring are lost or chicks fledge; at the latter stages, nests begin to lose structure and colour. At any point, nests may be empty, occupied by adults, eggs or juveniles, or a combination depending on parental foraging and care and chick mortality and fledging (Figure 1). All four colonies we surveyed had a mix of nest and juvenile ages, along with foraging and in-flight birds away from nests. Most (>90%-95%) of birds in the colonies were Straw-necked Ibis, a dark glossy blue-black bird on their back and wings, and with a white underside (black when viewed from above). The remaining 5%-10% of birds were composed of Australian White Ibis T. Molucca, Glossy Ibis Plegadis falcinellus and Spoonbills Platalea spp. All the bird species are <1 m tall and <50 cm in length when roosting, and most nests are between 20 and 40 cm in diameter.

2.2 | Drone data and processing

Drone image data were collected during Spring and Summer of 2016 and 2017 (Table 1), using a DJI Phantom 3 Professional quadcopter, with the stock single sensor red/green/blue (RGB) camera (4,000 \times 3,000 image size, lens FOV 94° 20 mm). Colonies were within large flooded extents (km's wide), so multi-rotor drones were the only option, with no landing area for fixed-wing drones. We launched a drone from an amphibious vehicle or canoe used to enter the colonies. Flights were conducted using parallel flight lines, at c. 100 m and speed of 5–10 $\rm ms^{-1}$ (Lyons, Brandis et al., 2018; Lyons et al., 2019). We aimed to acquire imagery with c. 70% forward and lateral overlap to ensure adequate coverage for post-processing. We acquired imagery over the entire extent of the four colonies, including a 200-300 m buffer around the colony edges. Depending on weather and environmental conditions, we surveyed 5-20 hectares per flight, requiring multiple flights to survey each colony. There were no obvious negative interactions with the waterbirds; further animal ethics considerations can be found in Lyons, Brandis et al. (2018), and a more detailed protocol for drone-based monitoring of waterbird colonies in Lyons et al. (2019).

The drone imagery was processed using the commercial software PIX4DMAPPER (v4.19, Pix4D SA), which uses a photogrammetry technique called 'structure from motion' to identify points in overlapping images, ultimately generating a 3D point cloud reconstruction of the landscape. The 3D information is then used to generate a digital surface model and an orthorectified image mosaic. Only standard accuracy GPS (5–10 m accuracy) was used for georeferencing. This resulted in some error in absolute geographic location, but was not important, given our focus on identification and relative position of nests in the image mosaics.

2.3 | Semi-automated approach for nest counting

Once the imagery was acquired, we needed to effectively identify nests which were highly variable in shape and colour, and sometimes had low contrast to the surrounding environment (Figure 1). We initially tested a point process algorithm (Descamps et al., 2011) but it could not handle large data sizes; an object-based image analysis routine (sensu Chabot et al. (2018)) but it had difficultly identifying more than 3,000-5,000 nests with one ruleset; and a machine learning/modelling approach (Hodgson et al., 2018) but it could not identify more than c. 1,000 nests with one parameterisation (see Data accessibility for modified Matlab routine). No particular technique worked effectively within or between the colonies, supporting similar findings on the limitations of automated and semi-automated methods (Hollings et al., 2018). So, we developed a modular approach, adaptable to variable target properties and scalable to large spatial extents, applicable to multiple colonies. This involved first mapping the area of nests using a remote sensing approach, and then estimating the number of nests using a predictive modelling approach.

2.3.1 | Manual counts for training and validation data

A comprehensive training and validation dataset was critical for developing counting methods. So, we first manually and systematically counted all the nests in the imagery over all colonies. We imposed a 50×50 m grid of quadrats on each colony, and digitally annotated every visible nest. We used this gridded method for two reasons: (a) it enabled an observer to sequentially work



FIGURE 1 Example drone imagery showing the variation in nest types and environments across four breeding colonial waterbird (mostly Straw-necked lbis) colonies. Images from top row to bottom row are from the following colonies: *Merrimajeel, Zoo Paddock, Eulimbah* and *Block Bank* (details in Table 1)

through the whole colony, while reducing distraction (and computer memory overhead) from surrounding areas; and (b) it reflected real-world practices when users choose only a limited number of training quadrats to manually count nests. During the field work, we also counted nests (in situ) at several GPS-tagged locations at each colony, which we used to test the accuracy of the drone-based manual counting.

2.3.2 | Machine learning mapping

We applied a supervised machine learning approach to map nests at each colony. We defined nests as any material or bird that constituted a nest or nest clump, based on our experience in the field. Motivated by its robustness to redundant predictor variables, we used a random forest classifier (Breiman, 2001), which enabled us to use a training dataset of nest locations and a suite of relevant image predictor variables to predict nests out into image pixels across the entire study area. The random forest procedure works by taking many different bootstrapped samples of the training data and fitting a classification tree to each of them, with each candidate split along the decision tree considering a random selection of the predictor variables. This ensures uncorrelated trees and stops the decision trees from overfitting (Breiman, 2001). The final prediction is taken as the mode of the classification tree results, with the classification being robust to collinear and redundant predictor data. This allowed us to include many different image-based predictor variables without altering the approach for different colonies.

All nests were manually identified, so we sampled a subset to train the random forest classifier to simulate the scenario of manually counting only a small proportion of the nests/image area. To simulate the 50 \times 50 m guadrats, we placed a grid of points across the colony at least 30 m apart and within 1 m of an identified nest (to avoid choosing training areas where there were no nests), and randomly chose a number of those points as a classifier training location. To approximate the 50×50 m quadrat area, a 30 m buffer was placed around each chosen training locations, within which all manually counted nests were selected for training. Each 30 m buffer would select between c. 100-1,000 nests. We trialled between 5 and 20 training locations for each of the colonies. The classifier also requires non-target features (non-nest) randomly spread across the colonies: 1,000 points for the smaller colonies (Eulimbah and Block Bank) and 10,000 points for the larger colonies (Merrimajeel and Zoo Paddock).

We derived arithmetic and textural metrics from the red, green and blue channels (*r*, *g*, *b* respectively) in the drone data to use as predictor variables in the random forest classification. These included: a 'white' index $\frac{b+g}{r}$; a Laplacian-8 edge-detection kernel on the 'white' metric; an RGB vegetation index $\frac{g-r}{g+r}$ (Bendig et al., 2015); a 'green brightness' index $\frac{g}{b+g+r}$; the 'contrast', 'variance', 'inverse difference moment' and 'shade' texture metrics from the Gray Level Co-occurrence Matrix (Haralick, 1979), applied to each of the 'white' index and blue band; the standard deviation within a 2 m and 7 m radius of each pixel applied to the 'shade' metric and vegetation index; and a 1st and 2nd order difference of Gaussians (Polakowski et al., 1997) on the 'shade' metric.

The training dataset was compiled by extracting the pixel values for each image metric layer within a 10 cm buffer (minimum nest diameter 20 cm), around each training nest and non-nest point, so the random forest classifier was a binary nest and non-nest classification. The algorithm was parameterised with 500 trees and a minimum leaf population of 10. We implemented the predictor variable calculation and random forest classification in Google Earth Engine (Gorelick et al., 2017), allowing a seamless prototyping, visualisation and production environment for processing the large high-resolution image datasets. Any contiguous areas <0.03 m² (minimum nest size of 20 cm diameter) were deemed to be noise and removed before exporting. Google Earth Engine is freely available to anyone, and we provide the JavaScript code required to run the classifications, along with an interactive web-app to explore some drone data, predictor layers and nest classification interactively (link in Data accessibility section).

We assessed the nest mapping accuracy via a standard error matrix approach, using the full manually counted dataset, along with an extended collection of background points. The background points were compiled from random points spread across the colony area (approx. as many points as there are nests), constrained to be >20 cm from a nest point. This provided a balanced validation dataset, to get a good estimate of overall mapping accuracy, as well as omission and commission error for nest mapping. We used an empirical bootstrapping approach (Lyons, Keith, Phinn, Mason, & Elith, 2018) to get confidence intervals for overall accuracy. Accuracy metrics were calculated in R version 3.5.1 ((R Core Team 2018); see Data accessibility section).

2.3.3 | Predictive model estimation

To estimate the number of nests as a function of the mapped nest area for each colony, we used predictive modelling. We first summarised the number of manually counted nests and the mapped nest area within each 50×50 m quadrat. We then predicted the number of nests in each quadrat, with the whole colony count being the sum of the quadrat estimates. We used two simple approaches: (a) an assumption that the number of nests was directly proportional to the mapped nest area (linear area:count ratio); and (b) a generalised linear model (GLM; Poisson error distribution) of nest count as a function of nest area and local nest density. We expected that the local density of nests would have a relationship with the number of nests. Density was calculated as the percentage of the 50×50 m quadrat mapped as nests. Using a GLM with a negative binomial error distribution or a generalised additive model with smoothers for nest area and density provided no appreciable gains, so neither was pursued.

We used a resampling procedure to examine the number of manually counted 50 × 50 m quadrats needed to accurately estimate the number of nests for a whole colony. This involved repeated random sampling of *n* quadrats, estimating the number of nests using the area ratio and GLM approaches described above. We used 800 iterations without replacement (i.e. Monte Carlo resampling, not a bootstrap) for each of 1, 2, ..., n_{max} quadrats. This resulted in a sampling distribution of 800 whole-colony nest count estimates at each *n*.

To simulate the scenario of limited resources for manual counting, we implemented another resampling approach to determine whether a given sample of the manually counted quadrats provided an accurate estimate (plus a confidence interval) of nest count for a whole colony. This involved a random draw of *n* quadrats (i.e. scenario of choosing a set of quadrats for training), and applying a repeated *k*-fold cross-validation using the area ratio and GLM estimation approaches. Each random draw of quadrats was stratified by mapped nest area density, to simulate choosing a range of nest density quadrats to count. We used *k* = 10 and 10 repeats for the crossvalidation, and varied *n* from *c*. 10%–40% of the total number of manually counted quadrats. This resulted in a sampling distribution of 100 nest count estimates for each random draw of quadrats, and we took the mean as the resampling estimate and 2.5 and 97.5 percentiles as a 95% confidence interval. We decided on *k*-fold resampling as a good approach to reduce bias for the small sample sizes, but a range of resampling options are available (Lyons, Keith, et al., 2018). All statistical analysis was performed in R version 3.5.1 ((R Core Team 2018); see Data accessibility section).

3 | RESULTS

3.1 | Manual training and validation nest counts

The four study colonies varied widely in size, number of nests and bird density (Table 1). The flying height of *c*. 100 m generated orthomosaic imagery with a pixel size between 3 and 4 cm. It took 5–15 min to manually count the nests in a 50 × 50 m quadrat, with higher nest density on the upper end of that time. Ibis nests and the flooded colony environment were so variably complex that it was often not possible to accurately manually count nests, even from 3 to 4 cm pixel drone imagery. Occasionally, artefacts from drone imagery processing also obstructed counting. The accuracy of the manual counting was estimated using the on-ground field counts, which ranged from $\pm6\%$ to $\pm12\%$ (Table 1). The smallest colony had a manual count of 7,717 nests and the largest colony had 96,989 nests, and with an estimated population of over 200,000 birds at the time (Lyons, Brandis et al., 2018).

3.2 | Semi-automated approach

The same Google Earth Engine code was applied to each colony, showing that the nest area mapping routine was robust to differing background environments and nest characteristics within and among each of the colonies. Around 10 of the 30 m training buffer locations were required for consistent classification of the large extent colonies (Merrimajeel, Zoo Paddock; c. 5% total area), and around 5 for the smaller extent colonies (Eulimbah, Block Bank; c. 10% total area). The chosen predictor variables did a good job at extracting the salient features of the bird colonies (Figure 2) and the machine learning classification appeared to identify nests and birds appropriately (Figure 3). The accuracy assessment showed the nest mapping routine performed well, with overall accuracy between 94% and 96% for Merrimajeel, Eulimbah and Block Bank and 86% for Zoo Paddock (Table 2). For all colonies, omission error was low and commission error was higher (Table 2), demonstrating that the mapping routine rarely missed nests, but included too much background as nest material. This was particularly the case for Zoo Paddock, which had a very high commission error (27.3%) and a lower overall accuracy (86%). Supporting Information Table S1 contains the raw accuracy assessment error matrices from which the accuracy measures were derived.

The first resampling routine demonstrated that considerable variation in nest estimates was likely given any random draw of quadrats, but only a small subset of the quadrats was required to capture most of the variation and provide estimates within the manual count error range (Supporting Information Figure S1). For the three colonies with >50 quadrats, the variability increased relative to the population size component of a finite population corrected standard error, $\sqrt{\left(1-\frac{n}{N}\right)/N}$, where *n* is the number of sampled quadrats and *N* is the total available quadrats. So for larger colonies, users could use this relationship with choose the size of the manual count needed based on either a desired count error or estimate where it starts to tail off towards zero. There was no noticeable improvement in using the GLM estimation method over the straight area ratio method. Comparing the results of the nest count estimates for individual quadrats showed that there was a large amount of variation among estimates for individual quadrats, the primary motivation for the use of a resampling-based estimate (Supporting Information Figure S2).

For the k-fold nest count estimation, we decided that an adequate number of quadrats (n) to use would be signified by most of the estimates from each k-fold cross-validation falling within the error margin of the manual nest counting (Figure 4, Table 3). For the two largest colonies, Merrimajeel and Zoo Paddock, we used 30 quadrats (c. 12% of all 50 × 50 m quadrats) to provide accurate estimates. For the two smaller colonies, Eulimbah and Block Bank, we used 15 and 10 quadrats respectively (c. 20% and c. 30% of total quadrats respectively). The manual effort time saving was best for the larger colonies-the nest counts were eight times faster for the two larger colonies (Merrimajeel and Zoo Paddock), but only five and three times faster for Eulimbah and Block Bank respectively (Table 3). The estimation was most accurate for the smaller two colonies, and there was some over estimation for the larger colonies, particularly Zoo Paddock (Figure 4), that could not be rectified with more training data. Again, there was no noticeable gain in using the GLM estimation method over the straight area ratio method; the gain from stratifying the random draw by mapped nest density was far more appreciable.

4 | DISCUSSION

We developed a generalised approach for monitoring complex wildlife aggregations, demonstrated through semi-automated analysis of drone-based remote sensing imagery over four large and complex waterbird colonies. The approach accurately mapped nests, and subsequently provided accurate estimates of nest numbers. The method offers significant time savings compared to manual counts from the imagery. In our study, we obtained accurate maps and estimates of nests for one of Australia's more extensive breeding of colonial waterbirds, and some of the largest waterbird colonies ever surveyed via drone. Our methodology is simple and robust enough to be applied in multiple environments, and works for both simple and complex target features. Continued development will see drone-based monitoring become integrated into waterbird monitoring (Lyons et al., 2019), and used to help quantify salient biological features like nesting success (Sarda-Palomera, Bota, Padilla, Brotons, & Sarda, 2017). There are clear benefits for monitoring some of

FIGURE 2 An example of image metrics derived from drone imagery over a waterbird (mostly Strawnecked Ibis) colony (Eulimbah), used as predictor variables in the random forest classification: (a) raw drone imagery; (b) 'white' brightness image; (c and d) Gray Level Co-occurrence Matrix (GLCM) 'shade' and 'contrast' of the 'white' metric; (e) GLCM 'shade' of the blue reflectance; (f) RGB vegetation index; (g) difference of gaussians applied to the GLCM 'shade' on the 'white' metric; (h) an RGB composite of the 'white' metric and the standard deviation within a 2 m and 7 m radius for the GLCM 'shade' of the 'white' metric



the many other species of birds that form complex aggregations, and our methods will be easy to test on other wildlife in different environments.

Our approach is modular, and the nest mapping and counting is implemented on free open source platforms, allowing users to readily change parameters or substitute their own or more appropriate methods. The semi-automated approach applied a machine learning classifier to high-resolution drone imagery to identify nests (Figures 2 and 3, Table 2), supported by modelling to estimate nest counts (Figure 4). The methods were effectively applied across four different waterbird colonies, that contained highly variable target features on variable backgrounds. The colonies ranged in size from



FIGURE 3 Example nest area classifications for four colonial waterbird (mostly Straw-necked Ibis) colonies, surveyed via drone and classified using a random forest classifier in the Google Earth Engine. Images from top row to bottom row are from the following colonies: Merrimajeel, Zoo Paddock, Eulimbah and Block Bank. Full details in Tables 1 and 2

around 7,000 nests to almost 100,000 nests (Table 1), and our semi-automated method required only relatively small amount of training data to produce comparable accuracy to manually counting from the drone imagery (Figure 4, Table 3). Here we further discuss the cost-benefit aspects, opportunities for wider uptake, current challenges, and finish with some recommendations moving forward.

4.1 | Cost-benefit of the semi-automated approach

The two key motivators for drone-based automated methods are reducing (on-ground) human observer bias and reducing cost (Baxter & Hamilton, 2018; Chabot & Bird, 2015; Hodgson et al., 2018; Hollings et al., 2018). For large and complex wildlife aggregations, such as our waterbird colonies, it is rarely possible to perform comprehensive

TABLE 2 Accuracy assessment results for mapping of nests in four different waterbird colonies, based on drone imagery and a machine learning algorithm. Measures include the overall accuracy (plus a bootstrapped 95% confidence interval) and the omission and commission error for nests

Colony name	Overall accu- racy (%) (95% Cl)	Nest omis- sion error (%)	Nest commis- sion error (%)
Merrimajeel	94.1 (94-94.2)	1.2	10.4
Zoo Paddock	86 (85.6-86.3)	0.9	27.3
Eulimbah	95.8 (95.5–96)	1.7	6.9
Block Bank	94.7 (94.3–95)	4.7	5.9

on-ground counts and so drone-use provides an attractive option, and coupled with semi-automated methods, presents significant time savings too. In our case the large colonies were eight times quicker to survey (Table 3), representing a cost (time or money) saving of almost 90% compared to full manual counts. Cost-benefit analysis will continue to vary with user ability and conditions, including data acquisition in the field, drone image processing, modelling and programming, and even the level of detail and accuracy required for monitoring outcomes. Benefits will also vary with the nature of application with one-off monitoring perhaps better achieved using manual methods, until the technology and processes become routine. The efficiency increases in Table 3 are probably more likely to represent the potential time savings of further application of the methods. For new applications in novel environments, large manual counts are probably still required to understand the potential sources of error.

4.2 | Opportunities for uptake of semiautomated methods

Transferability across environments and spatial scales has prevented widespread adoption of semi and fully automated methods in wildlife monitoring (Chabot & Francis, 2016; Hollings et al., 2018). We successfully implemented our semi-automated approach, using the same routine/code, on four different waterbird colonies, providing immediate applicability for other avian applications, and opportunities more broadly for large complex aggregations of wildlife. The key challenge we overcame was identification of target features with high spatial and spectral variation, on high variability backgrounds, across large spatial extents. Most current detection approaches rely on methods that require high consistency in the spatial and spectral organisation of target and background features. Our use of a random forest classifier efficiently handled redundant predictor data (Breiman, 2001), allowing inclusion of many different colour, spatial and textural metrics as predictor layers. This helped capture more of the spatial and spectral variation in target features, compared to just using the image colour, as well as potential image blur and illumination artefacts (Figure 3, top row).



FIGURE 4 Resampling estimates of nest counts for four breeding waterbird (mostly Straw-necked Ibis) colonies surveyed via drone, trained using a classification of nest area and manually counted nests. Each black dot represents the mean of the sampling distribution ($10 \times$ repeated *k*-fold *k* = 10 cross-validation) for a different subset of the manually counted training nests (corresponding lines denote 95% percentile), and the red horizontal lines denote the manual estimate for the whole colony, and the 95% error margin calculated from on-ground counts

Colony name	50 × 50 m quad- rats in grid	Manual nest count (±manual error)	Mean and range of <i>k</i> -fold nest estimates	Full count ef- fort (hours)	k-fold count effort (hours and speed-up)
Merrimajeel	233	96,989 (91,073-102,905)	99,645 (90,383–106,727)	40	5 (8×)
Zoo Paddock	244	20,411 (18,615–22,207)	21,432 (16,627–27,361)	42	5 (8×)
Eulimbah	71	13,343 (12,222-14,464)	13,479 (12,212-14,879)	12	2.5 (5×)
Block Bank	33	7,717 (6,783-8,651)	7,777 (7,152–8,425)	5.5	2 (3×)

TABLE 3 Manual and semi-automated counting results for drone-surveyed waterbird colonies. Colonies were divided into a grid of quadrats and nests were manually counted with accuracy from in situ counts. *k*-fold nest estimates were derived from our semi-automated approach, using 40 different random subsets of quadrats

Research is increasingly tackling this issue of consistency across target and background features, adapting methods from remote sensing mapping (Afán et al., 2018; Chabot et al., 2018). Application of a single consistent detection routine to many different applications will provide significant opportunity for broad uptake across scientific and management applications (Hollings et al., 2018). Although we found that existing methods (Chabot et al., 2018; Descamps et al., 2011; Hodgson et al., 2018) were not directly able to deal with the level of complexity in our case studies, our method ultimately required more training data, which reduces the overall cost saving. Thus continued development of a range of methods will provide opportunities for significant time and cost savings when applied over large spatial extents, over time.

Detection approaches from imagery, such as drone imagery, are increasingly benefiting from the remote sensing disciplines (Chabot et al., 2018), due to innovations in dealing with large volumes of data efficiently. Existing detection methods typically deal with image tiles in the order of 1–10 Mb. Our waterbird colonies involved 500 Mb to 5 Gb of data, requiring significantly improved data management and analysis. Use of the Google Earth Engine platform (or similar platforms) enables handling of large data, and will facilitate future expansion into web-based tools where users only supply imagery and training data, reducing local expertise and computing resource requirements.

We successfully identified both nests and individual birds when they were away from their nests (see Figure 3, 3rd row). This demonstrates the opportunity to use our mapping driven approach to identify and count individual waterbirds. Indeed Chabot et al. (2018) used an object-based mapping approach for identifying and counting individual Snow Geese. If only individual birds were of interest, and they did not form complex spatial aggregations, the mapping process would be sufficient to identify and count individuals (i.e. the *k*-fold estimation process would be unnecessary). This represents an additional cost saving because it took less training data to train the machine learning mapping (e.g. for Merrimajeel, c. 5% of the guadrats were needed to train the random forest, but 12% were needed to train the k-fold estimation). For small and simple tasks (e.g. counting just a few thousand birds or nests) our k-fold estimation process could also be replaced with simple thresholding or classification of the predictor metrics. For example thresholding and vectorizing the predictor layers we used (e.g. Figure 2, bottom row) produces accurate nest counts, but these

thresholds become increasingly variable as spatial scale increases, making consistent application difficult.

4.3 | Challenges for drone-based monitoring

Our main challenge was converting nest maps to nest counts. A remote sensing approach allowed us to make very accurate nest maps (three colonies c. 95% accurate, one colony c. 86%; Table 2), but the complex organisation of nests (i.e. different shapes and sizes, irregular overlapping and aggregation) prevented a direct conversion to singular pin-pointed nest locations. We used the k-fold estimation approach to overcome this limitation to estimate nest counts from nest maps, but note that while the overall colony counts were accurate, nest counts for some individual quadrats contained significant error (Figure 4, Supporting Information Figure S2). For the Zoo Paddock colony, the high commission error (Table 2), led to some over estimation of nest count (Figure 4, Table 3). This colony had a large spatial extent but was only sparsely populated, compared to the other large colony (Merrimajeel; Table 1). Improved modelling of density effects may reduce this problem. As it was, only five out of the 40 scenarios we ran would be considered a sizable overestimation (Figure 4), and even then these numbers would be unlikely to affect local management decisions (Brandis et al., 2011), but this may vary depending on the application. We randomly selected quadrats, so a more judicious initial choice of quadrats for training (e.g. choosing a range of nest densities) may also rectify this issue to some degree.

Another challenge is the potential impact of uncertainty (c. 6%–12%, Table 1) in manual counting that can propagate through to the mapping and estimations. Moving semi-automated methods to increasing spatial scales or more complex environments requires dealing with more variation in image quality and limitations in the resolution able to be captured (Hollings et al., 2018). In our surveys, image resolution and quality was a challenge, affected by our ability to access appropriate remote points for take-off and landing, along with environmental and ethics considerations that limited time available to collect imagery. This led to varying incident sun angles and wind conditions during image collection, resulting in sun glint and image blur that sometimes obscured manual counting. Identifying old nests (e.g. Figure 1, top and bottom rows) was difficult, potentially further increasing manual counting errors. Our cross-validation approach was motivated by the need to account for uncertainty,

and generally accounted well for this error (Figure 4, Supporting Information Figures S1 and S2), but had a cost in terms of increased training data requirements.

Another challenge is the potential antipathy towards use of drones, because sometimes the literature presents them in terms of taking over the role of surveyors. This is unjustified, because equally large amounts of human effort will continue to be required for collection and processing of drone imagery, collecting and curating training and test data, and developing detection routines. Just as Fraser et al. (1999), almost two decades ago, demonstrated improved aerial counting from a kite-mounted camera, drones are now becoming part of the toolkit. Furthermore, researchers and managers can be excited about access to fast and accurate counting, without adequately considering the potential uncertainty, labour and skills required for effective use of drones for monitoring large and complex wildlife aggregations, and that drones still cannot produce all the required biodiversity metrics for monitoring (Callaghan, Brandis, Lyons, Ryall, & Kingsford, 2018).

4.4 | Recommendations

There are major improvements in data collection, interpretation and understanding which can come through using drone imagery, including cost savings and potentially improved accuracy. Applications will continue to grow, assisted by development of semi-automated methods such as ours. Drones should be viewed as a tool to complement ecological and environmental monitoring practitioners, rather than a replacement option. We suggest development of semi-automated approaches should focus on adaptability to deliver key monitoring indicators (Baxter & Hamilton, 2018), and that detection methods themselves should aim for three main properties: (a) use predictor data that is easily derived from common drone-based (or airborne) imagery; (b) minimal parametrisation among environments, ensuring any parametrisation should be accessible to non-expert users; and (c) implementation on widely available platforms, not requiring significant local computing resources but able to manage large volumes of image data.

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AUTHORS' CONTRIBUTIONS

All authors contributed to study design; M.B.L., C.T.C., J.A.M. and K.J.B. carried out field work; M.B.L., J.H.W. and N.J.M. led the data processing and statistical analysis; and all authors wrote the manuscript.

DATA ACCESSIBILITY

Our nest mapping routines were implemented in Google Earth Engine, and the code we provide can be directly run in the JavaScript API (https://earthengine.google.com/). All statistical analyses, including nest counting and accuracy assessment, were performed in the R programming environment (R Core Team 2018). The Earth Engine and R code are available on Github (https://github.com/mitch est/bird-colony-count-drones) and archived on Zenodo (https:// doi.org/10.5281/zenodo.2594095). The Merrimajeel, Zoo Paddock and Block Bank colonies are on private/sensitive land, over which raw drone data cannot be released publicly, however, the code and data provided includes the summarised data and spatial data files required for our analyses. The Eulimbah colony is on public land, and we have developed a web-app (https://mitchest.users.earthengine. app/view/ibis-drone-count) through which users can explore drone data, predictor variables and the nest map classification interactively, and the raw orthorectified image mosaic can be downloaded from Dryad Digital Repository (https://doi.org/10.5061/dryad. m4r0cn0). Data available from Lyons MB, Brandis KJ, Murray NJ, et al. Monitoring large and complex wildlife aggregations with drones. https://doi.org/10.5061/dryad.m4r0cn0

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SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of the article.

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