Combining live and lethal trapping to inform the management of alien invasive rodent populations in a tropical montane forest

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Academic editor: S Bertolino | Received 30 April 2020 | Accepted 20 October 2020 | Published 8 December 2020

Citation: Duron Q, Cornulier T, Vidal E, Bourguet E, Ruffino L (2020) Combining live and lethal trapping to inform the management of alien invasive rodent populations in a tropical montane forest. NeoBiota 63: 101–125. https://doi.org/10.3897/neobiota.63.53811

Abstract

On large inhabited islands where complete eradication of alien invasive rodents through the use of poison delivery is often not practical or acceptable, mechanical trapping may represent the only viable option to reduce their impact in areas of high biodiversity value. However, the feasibility of sustained rodent control by trapping remains uncertain under realistic operational constraints. This study aimed to assess the effectiveness of non-toxic rat control strategies through a combination of lethal and live-trapping experiments, and scenario modelling, using the example of a remote montane rainforest of New Caledonia. Rat densities, estimated with spatially-explicit capture-recapture models, fluctuated seasonally (9.5–33.6 ind.ha⁻¹). Capture probability (.01–.25) and home range sizes (HR₉₅, .23–.75 ha) varied greatly according to trapping session, age class, sex and species. Controlling rats through the use of lethal trapping allowed maintaining rat densities at ca. 8 ind.ha⁻¹ over a seven-month period in a 5.5-ha montane forest. Simulation models based on field parameter estimates over a 200-ha pilot management area indicated that without any financial and social constraints, trapping grids with the finest mesh sizes achieved cumulative capture probabilities > .90 after 15 trapping days, but were difficult to implement and sustain with the
local workforce. We evaluated the costs and effectiveness of alternative trapping strategies taking into account the prevailing set of local constraints, and identified those that were likely to be successful. Scenario modelling, informed by trapping experiments, is a flexible tool for informing the design of sustainable control programs of island-invasive rodent populations, under idiosyncratic local circumstances.

Keywords
Invasive species, island conservation, predator control, Rattus, trap

Introduction

Human activities, such as agriculture and international trade, modify habitats and disturb the composition, richness and diversity of animal and plant communities (Garrott et al. 1993; Vitousek et al. 1997). Alien invasive mammals cause detrimental ecological effects on native taxa through predation, competition and spread of diseases (Garrott et al. 1993; Towns et al. 2006; Wäber et al. 2013; Medina et al. 2014; Lieury et al. 2015), and further alter ecosystem structure and functions through trophic cascades (Chollet and Martin 2013; Thoresen et al. 2017). In areas where complete removal of invasive mammals is not feasible or appropriate due to technical, social or ethical issues, the permanent control of their populations is a widely used approach to restore degraded ecosystems (Goodrich and Buskirk 1995; Doherty and Ritchie 2017; Lambin et al. 2020) and has proven successful in enhancing the recovery of a wide range of endangered taxa (Fletcher et al. 2010; Smith et al. 2010; Jones et al. 2016). Control programs by trapping, shooting or poison delivery are, however, a lengthy endeavour, with ongoing expenditures due to compensatory demographic effects leading to reinvasion (Gundersen et al. 2001). Density-dependent dispersal and changes in competitive interactions in the removal area, especially, enable target populations to recover through immigration from uncontrolled surrounding areas (Efford et al. 2000; Gundersen et al. 2001; Beasley et al. 2013; Lieury et al. 2015). A fundamental applied question is therefore how to identify control strategies that deliver maximal benefits to biodiversity, while reducing reinvasion risks and operational costs (Doherty and Ritchie 2017).

Among the world’s most pervasive invasive species, rodents (Rattus spp. and Mus musculus) greatly contribute to ecosystem degradation and biodiversity loss on oceanic islands (Atkinson 1985; Towns et al. 2006; Harper and Bunbury 2015). Rodent control programs are a common practice to achieve biodiversity conservation targets worldwide (Bomford and O’Brien 1995; Armstrong et al. 2014); yet these programs still represent a significant challenge for conservation scientists and practitioners (Duron et al. 2017). Improvement of control methods, local community engagement and adaptive management have been identified as key to improve the cost-effectiveness and sustainability of such projects (Duron et al. 2017). In addition, rat management techniques should benefit from collating baseline data on the population structure and dynamics of the target rodent populations, as well as a better understanding of the processes influencing their demographic responses to culling.
When rodent control needs to be conducted in the long term, poison delivered as bait is the most cost-effective measure, especially when treated areas are large and dominated by rugged terrain (Russell et al. 2008). However, repeated use of rodenticides can be deleterious to non-target native species and the environment (Hoare and Hare 2006; Pitt et al. 2015; Duron et al. 2017) and lead to rat resistance to toxins (e.g. King et al. 2011) or objections from individuals or organizations concerned about toxicity issues. Therefore, selective lethal trapping may be considered as a more ecologically-friendly option, and in areas where the use of rodenticides is unlawful or not accepted by local communities, the only viable management approach (Ogden and Gilbert 2008). While control by trapping has proven successful to suppress rodent numbers to low densities in a short-term commitment (Stokes et al. 2009; King et al. 2011; Pender et al. 2013), we believe that there is a scope for optimise rodent trapping strategies in order to enhance project sustainability and achieve long-term ecological outcomes.

In the South Pacific archipelago of New Caledonia, Black and Pacific rats (Rattus rattus and R. exulans) are among the main predators of endemic and micro-endemic species in the dense evergreen rainforests (Thibault et al. 2017; Duron et al. 2019). As eradication of the 15 971 km² main island, which is inhabited and composed by complex ecosystems, is not currently practical, rat population control in targeted areas over prolonged periods represents a candidate option to restore native ecosystems. Most of the pristine remnants of New Caledonian rainforests lie in remote, mountainous areas, which renders the logistics of achieving effective management over significant spatial scales highly challenging. Although poisoning should have been the most cost-effective option in this context (Russell et al. 2008), in New-Caledonia, local native Kanak communities as well as local communities descending from European settlers, and local environment NGOs, have expressed strong disapproval regarding the use of poisoning for invasive species control due to the risk of toxicity to bushmeat, fish or drinking water (Cassan JJ, Northern Province Environmental Service, pers. comm.; see also Groseil 2010). Given these legitimate concerns, local communities and NGOs have expressed their interest in an ecologically-based rat management program that involves mechanical trapping without rodenticide application. Rat populations dwelling in the mountainous rainforests of New Caledonia have not been studied in depth (Rouys and Theuerkauf 2003; Thibault et al. 2017). Therefore, their biology and population dynamics need to be better understood if cost-effective, sustainable management strategies are to be developed in these pristine remnants of rainforest biodiversity hotspot.

Our study aimed at evaluating the effectiveness of mechanical trapping for maintaining rat numbers at low levels using a study site in a remote montane rainforest of New Caledonia as a case study. We combined a capture-mark-recapture (CMR) study with a small-scale (5.5 ha) intensive lethal trapping experiment to i) estimate rat densities and home ranges across species, sex, age and seasons; ii) assess the effects of a knock-down removal trapping experiment on the rat population structure; and iii) identify trapping grid layouts that maximise rat capture probability, given workforce availability constraints, using simulation models parameterized with rat population parameters derived from CMR data. This work will help conservation managers make evidence-based decisions for the management of invasive species.
Methods

Study site

This study was conducted in a dense evergreen rainforest located between 550 and 950 meters a.s.l. in the wilderness reserve of Mont Panié (20°37'30"S, 164°46'56"E, 5400 ha) in New Caledonia, South Pacific (Fig. 1). This area has been protected since 1950 on the basis of the cultural importance of Mont Panié and the high micro-endemism rates observed for most taxa (plants, insects, reptiles) (Tron et al. 2013). The climate is moist subtropical, with a hot season between December and mid-April and a cool season between mid-May and September. Mean annual precipitations in the Mont Panié region are around 3 000–4 000 mm, while mean temperatures range from about 18 °C to 25 °C throughout the year, and minimum temperatures can approach 0 °C (Tron et al. 2013). Two invasive rat species, *Rattus rattus* and *R. exulans*, are present in the study area. *R. exulans* was introduced in New Caledonia by the first Austronesian colonisers ca. 3 000 years ago, while *R. rattus* was introduced later by European settlers ca. 150 years ago (Beauvais et al. 2006). Both rat species are omnivorous, eating mainly plants, invertebrates and, to a lesser extent, Squamata, with *R. rattus* being more frugivorous and *R. exulans* more omnivorous (Duron et al. 2019).

Field sampling

Rat capture-mark-recapture (CMR)

Rats were live-trapped between September 2014 and September 2015 in the vicinity of the small removal area described below, in order to i) study rat population dynamics within the CMR area, and ii) monitor possible movements between both CMR and removal areas in response to ongoing lethal trapping in the adjacent removal area. Ninety-six standard wire cage rodent live-traps (with spring door) were set 20 m apart on a 12 × 8 trapping grid (3.08 ha, Fig. 1). Six CMR sessions (of five to eight consecutive days each) were conducted during the periods of 28 September-02 October 2014 (session 1), 26 November-02 December 2014 (session 2), 18–23 March 2015 (session 3), 21–26 May 2015 (session 4), 4–10 July 2015 (session 5) and 02–09 September 2015 (session 6). Traps were baited with fresh coconut chunks and checked every morning. Bedding material and leaf roof were provided to reduce mortality due to occasional rainy and cold conditions. Newly captured individuals were individually marked with small subcutaneous PIT-tags (type Tiny chip iso 1.4 × 8 mm size; Biolog-id, Paris, France). Each new individual was identified to species, sexed, weighed to the nearest 2.5 g, and checked for sexual maturity (females: perforate vagina and teat development status, males: presence of a scrotal sac). Since most black rat individuals weighing > 120 g and most Pacific rat individuals weighing > 55 g were sexually mature (respectively 94% and 98%, based on autopsies; \( n = 166 \)), body mass was used as a proxy for rat sexual maturity.
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Small-scale rat removal trapping

Rat removal trapping was performed between May and November 2015 in a 5.5-ha forest area immediately adjacent to the CMR trapping grid (Fig. 1) in order to 1) assess the effectiveness of lethal trapping in maintaining rat populations at low densities, and 2) study the reinvasion process over time. Rats were caught with the “Snap E Rat Trap” (Kness Mfg. Co., Albia, IA, USA) set 20 m apart on a 460 × 180 m trapping grid (5.5 ha). As the grid was crossed by a river, a total of 209 snap traps were set up (Fig. 1). Four lethal trapping sessions took place in 2015: 28 May-05 June (9 nights), 11–15 July (5 nights), 08–12 September (5 nights) and 28 October-01 November (5 nights). Traps were baited with coconut chunks and checked at sunrise. Animals caught were identified to species, sexed, weighed, and classified as adults or juveniles (as for the CMR experiment). We also recorded individuals that had previously been PIT-tagged, indicating movements between the CMR and removal grids.

Data analysis

Rat population densities and home ranges inferred from spatially-explicit capture recapture

Rat densities, home ranges and movements were estimated with spatially-explicit capture-recapture (SECR) analysis (Efford 2004; Borchers and Efford 2008). SECR was shown to outperform other methods for estimating density, and is preferable when capture probabilities are low (Blanc et al. 2013; Ivan et al. 2013). The number of range centers in an arbitrary area is Poisson-distributed, and range centers are independent and identically distributed uniform random variables. The probability of capturing an
animal at a particular trap location is assumed to be a half-Gaussian function of the Euclidian distance between the range center and the trap location, and is defined by two parameters: $g_0$ or the probability of capture at the center of the home range, and $\sigma$ or the distance to inflection point of the half-normal function describing the decay of capture probability with distance from an individual’s notional home range center. Models were fitted by maximising the full likelihood over all individuals observed, and by implementing a 60 m buffer ($3 \times \sqrt{\text{pooled square variance}}$; Efford 2004) around the trapping area. Accidental deaths ($n = 42$) during CMR trapping and PIT-tagged individuals killed in the removal area ($n = 17$) in May and July were accounted for in capture histories, with recapture probability fixed at 0 following death. As no general likelihood adjustments are currently available for single-catch devices, we implemented a multi-catch device likelihood, which appears only slightly biased for single-catch data (Efford et al. 2009). We explored differences in $g_0$ and $\sigma$ between rat species, age and sex categories, and between trapping sessions. We also tested for changes in $\sigma$ in response to the rat removal experiment in the immediate vicinity of the CMR grid, which began in May 2015, following the fourth CMR session.

Our dataset did not allow to test for the effect of individual sessions, rat species and age classes on $\sigma$ due to limited numbers of rat recaptures within each group category. We therefore grouped some sessions and individual categories together based on similarities in the shape of the raw data distribution, as well as environmental seasonality. As a result, sessions 1, 2, 6 (austral summer; $n = 44$ recaptures) and sessions 3, 4, 5 (austral winter; $n = 160$ recaptures) were merged, and each rat was assigned to one of the four following groups: $R. \text{exulans}$ ($n = 38$), $R. \text{rattus}$ juveniles ($n = 68$), $R. \text{rattus}$ adult males ($n = 52$), $R. \text{rattus}$ adult females ($n = 46$). Due to the limited number of data points, we did not test for interactions between covariates in SECR models.

Our final model combination allowed for testing the effects of a learned response to trapping, species, age, sex, session as well as group of sessions and group of individuals on $g_0$; and group of sessions, group of individuals and rat removal efforts on $\sigma$. The best model was selected by comparing AICc of all defined models (Hurvich and Tsai 1989). Significant effects were identified by investigating model outputs (parameter estimates and their 95% confidence intervals). From the best SECR model, we derived rat density estimates for each group of sessions and group of individuals. Then, based on estimated $\sigma$ and using the quantities of a circular Gaussian distribution, we calculated the 95% and 50% probability density areas of detection as surrogates for rat home range ($\text{HR}_{95}$) and home range core area ($\text{HR}_{50}$), as follows (see e.g. Ringler et al. 2014 for a Hazard function):

$$\text{HR}_{95} = \pi \times (2.45 \times \sigma)^2$$
$$\text{HR}_{50} = \pi \times (1.18 \times \sigma)^2$$

All the analyses described above were performed with the R package ‘secr’ 4.3.1 (Borchers and Efford 2008; Efford 2020) in R version 4.0.3.
Effects of the rat removal experiment on rat population densities and structure

Rat abundance \( (A_{\text{rat}}) \) in the removal area was estimated with the “Zippin removal” method, which assumes closed population within sessions and no heterogeneity in capture probability between individuals (Zippin 1958). Population densities before \( (D_{\text{before}}) \) and after \( (D_{\text{after}}) \) removal were estimated for each session, species, sex and age:

\[
D_{\text{before}} = \frac{A_{\text{rat}}}{\text{ETA}}
\]

\[
D_{\text{after}} = \frac{(A_{\text{rat}} - N_{\text{removed}})}{\text{ETA}},
\]

with ETA (effective trapping area: 9 ha) estimated as the size of the removal area (5.5 ha) plus a boundary strip (132.4 m) of the radius of the average SECR-derived rat home ranges from this study (0.42 ha), and \( N_{\text{removed}} \) the total number of rats killed during the trapping session. These analyses were performed with the R package ‘FSA’ (Ogle 2016) in R version 2.15.3.

Insights on reinvasion processes

We expected to observe a change in rat population structure in the removal area in response to the removal of a large number of resident individuals (Brown and Tuan 2005; King et al. 2011). To detect these potential changes, we compared the sex ratio, adult:juvenile ratio, average adult body mass and proportion of breeding individuals i) among the four removal trapping sessions, and ii) between the removal and CMR areas in sessions when both occurred subsequently (i.e. May, July and September 2015).

To detect temporal changes in spatial patterns of captures on the removal trapping grid, for each trapping occasion of each removal session we first calculated the average distance between trapping stations that had captured a rat and the nearest edge of the removal trapping grid \( (\text{Dist}_{\text{rats}}) \), and then compared \( \text{Dist}_{\text{rats}} \) to the average distance of the 209 traps to the nearest edge of the removal trapping grid \( (\text{Dist}_{\text{traps}} = 29.28 \text{ m}) \). We further expected that the home range centers of rats dwelling in the CMR area would shift towards the adjacent removal area in response to a sink effect. To test this hypothesis, from the best SECR model, we applied the fxi function in ‘secr’ (probability density function of home range center) to estimate, for each session, the home range center (HRC) coordinates of all rats recaptured at least once in the CMR area \( (n = 27) \). We then calculated, for each individual, the barycenter of its HRC coordinates before and after removal had started. Differences in barycenter locations before/after removal were tested by performing a paired Samples Wilcoxon Test.

Simulation of capture probabilities and control effort

The aim of this modelling exercise was to identify the rat management strategies that would yield the highest probability of rat capture within a single trapping session while being economically viable and socially acceptable in the remote area of the Mont Panié
wilderness reserve. We simulated the capture probability of one individual rat for a range of contrasted grids layouts over 200 hectares (10 different layouts; Table 1). This grid size already had local stakeholder approval as a pilot management unit and was defined as a compromise between operational constraints and likely benefits to local biodiversity. Grid mesh size varied according to distance between transects (15, 25, 50, 75 or 100 m) and distance between traps (15, 25, 50 or 75 m). For each of the ten different grid layouts, we simulated 10 000 rat home range center locations assuming complete spatial randomness and calculated the cumulative probability of capture of one individual rat over n trapping days.

In the absence of competition, the probability of capture of one rat with home range center at location i by trap j at time t is defined as follows (Efford 2004):

\[
P(\text{capt})_{ij} = g_0 e^{-\frac{d_{ij}^2}{2\sigma^2}}
\]

where \( d \) is the distance between i and j, \( \sigma \) is the scale parameter of the detection function and \( g_0 \) is the probability of rat capture at trap location j. Parameters \( g_0 \) and \( \sigma \) were estimated from our best SECR model based on the specific wire cage type of rat live-trap used in this study (see Results section). Parameters \( g_0 \) and \( \sigma \) are not known for snap traps; however, since rats can display neophobic behaviour (Clapperton 2006), \( g_0 \) is likely to be lower for cage traps than for snap traps. With this simulation exercise, we used cage trap parameters as a pessimistic series of scenarios. It is worth noting that the proposed approach could be applied to other trapping methods and devices, and other study sites and settings, once key parameters such as capture probability and home range sizes are calibrated.

The probability that one rat would be captured by any one of the b traps of a given grid layout over a n-day trapping session (or cumulative probability) is then:

\[
P(\text{capt})_i = 1 - \prod_{t=1}^{n} (1 - P(\text{capt})_{ijt})
\]

To assess how trapping efficiency varied across grid layouts as the trapping session progresses, we calculated the cumulative rat capture probability against time and project expenditure for each of the ten different grid layouts for one single trapping session. We estimated the average number of hours required to do a complete coverage of each grid over the entire trapping session, accounting for the decline in the number of rats captured as the trapping session progressed. The average time required to check and bait each trap was taken as 37 seconds, estimating that an empty trap that only needs rebaiting would take 30 s, and a trap where a rat had been captured would take 60 s. Based on our own field experience, we considered that it would take 10 minutes to walk 100 m through the rainforest while looking for traps. We constructed our simulation models based on the reasonable assumptions that a maximal number of 10 people would be willing to be away from their own villages for no more than 15 days, and each person would be willing to work 4 hours daily (fieldwork is rough and physically demanding) for a wage of 10 euros per hour. Our model accounted for a non-linear increase in project expenditure as the trapping session progresses, due to
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Table 1. Characteristics of the ten different removal trapping layouts tested in our simulation exercise. These calculations account for local social constraints in the Mont Panié area (i.e. 10 people available for 15 days and willing to work 4 hours a day).

| Layout | Dist. Transects | Dist. Traps | Nb. Traps | Nb. Hours | Nb. People | Nb. Splits |
|--------|-----------------|-------------|-----------|-----------|------------|------------|
| 1      | 15              | 15          | 8889      | 316.9     | 79         | 8          |
| 2      | 25              | 25          | 3200      | 169.6     | 42         | 5          |
| 3      | 25              | 50          | 1600      | 86.4      | 22         | 3          |
| 4      | 25              | 75          | 1067      | 58.7      | 15         | 2          |
| 5      | 25              | 100         | 800       | 44.9      | 11         | 2          |
| 6      | 50              | 50          | 800       | 78.2      | 20         | 2          |
| 7      | 50              | 75          | 533       | 53.3      | 13         | 2          |
| 8      | 50              | 100         | 400       | 40.8      | 10         | 2          |
| 9      | 75              | 75          | 356       | 51.4      | 13         | 2          |
| 10     | 75              | 100         | 267       | 39.4      | 10         | 1          |

Abbreviations: Dist. Transects: distance in meters between trapping grid transects; Dist. Traps: distance in meters between traps; Nb. Traps: number of traps used in each trapping layout; Nb. Hours: number of hours required to complete each grid as part of one trapping occasion; Nb. People: number of people required to complete each grid as part of one trapping occasion; Nb. Splits: number of splits required to complete each grid given number of people available.

some additional helicopter provisioning required every 15 days in this remote part of New Caledonia (provisioning costs for 10 people and 15 days were set to be 6 000 euros). Trapping equipment and grid cutting were not accounted for in the simulations as they would need to be costed separately, for example as initial investments (i.e. before the first trapping session commences) and running costs (i.e. maintenance) over multiple sessions.

Given economic and social constraints (i.e. 10 trappers available for 4 hours/day), some of the 200-ha layouts (i.e. with the finest mesh sizes) could not be completed within one day. In our calculations, we therefore allowed the grids to be trapped as adjacent separate management units (e.g. layout 1 was treated as 8 smaller units of 25 ha), each unit being trapped at a time. This strategy inevitably required extending the overall trapping session by n times ($n =$ number of management units). We assumed rat movements from non-controlled to controlled units to be minimal within this small temporal window (8 days maximum; see Table 1). Our calculations also accounted for initial baiting time and costs (people work to deploy and bait the traps but no rats are captured). Table 1 presents grid characteristics, and the number of people, hours and splits required to completely cover each grid as part of one trapping occasion.

Code and data for the CMR analysis and simulations are available at https://github.com/TCornulier/DuronRodentControl.

Results

Rat population densities and home ranges inferred from spatially-explicit capture recapture

The best SECR model had 29.81% relative support and included the effects of group of individuals and session on $g_o$, and group of individuals on $\sigma$ (Table 2). The
probability of capture $g_0$ was significantly lower for adult male *R. rattus* and higher for juvenile *R. rattus* than for *R. exulans*, whereas there was no significant difference in $g_0$ between adult female *R. rattus* and *R. exulans* (Table 3). Moreover, $g_0$ was significantly lower for CMR sessions 2 (December 2014) and 6 (September 2015) than for session 1 (September 2014; Table 3). Parameter $\sigma$ was significantly lower for juvenile *R. rattus* than *R. exulans*, and higher for adult male *R. rattus* than *R. exulans* (Table 3, 4). Rat densities varied greatly across CMR sessions, from 9.48 ± 1.84 ind.ha$^{-1}$ in July 2015 to 33.61 ± 9.00 ind.ha$^{-1}$ in November–December 2014 (Fig. 2). *R. rattus* adult densities were higher for females than for males during the three first capture sessions and similar during the three last ones (see Suppl. material 1). *R. rattus* juvenile densities markedly increased from CMR session 1 (September 2014: 0.27 ± 0.27 ind.ha$^{-1}$) to sessions 2 and 3 (December 2014: 8.74 ± 2.74 ind.ha$^{-1}$; March 2015: 9.94 ± 1.68 ind.ha$^{-1}$) and dropped again to low levels (< 4 ind.ha$^{-1}$) from session 4 to 6 (May to September 2015) (see also Suppl. material 1). No *R. exulans* was captured during the first two CMR trapping sessions. *R. exulans* were first live-trapped during session 3 (March 2015) and their densities remained stable over the three following sessions (Fig. 2; see also Suppl. material 1). Home range sizes (HR$_{95}$ and HR$_{50}$) were on average three times as large for *R. rattus* adult males as for *R. rattus* adult females (Table 4).

**Effects of removal trapping on rat population density and structure**

While nine trapping days were required to approach a near zero capture rate during removal session 1 (10 rats were captured at day 9 out of 209 traps), this rate was achieved after only two or three trapping days during the subsequent removal sessions. Rat density at the start of our removal experiment (D$_{before}$: May 2015) was estimated at 32.1 ind.ha$^{-1}$ (Fig. 2; see also Suppl. material 2). When comparing to the densities

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**Table 2.** Selection of the 10 best spatially explicit capture recapture (SECR) models. Model selection for effects of individual groups (indgr), session, group of session (sessgr), age, sex and learned response (b) on the probability of capture at the center of the home range ($g_0$) and on the scale parameter ($\sigma$). Note that although the effect of rat removal appeared in the second best model, this variable did not have any significant effect on sigma ($b = -0.21$, SE = 0.16, LCI = -0.52, UCI = 0.10).

| $g_0$ | $\sigma$ | N parameters | Log likelihood | AICc | % Weight |
|-------|----------|--------------|----------------|------|----------|
| indgr + session | indgr | 13 | -2046.78 | 4120.64 | 29.81% |
| indgr + session | indgr + removal | 14 | -2045.91 | 4121.06 | 24.20% |
| age + session | indgr | 11 | -2049.55 | 4121.87 | 16.09% |
| indgr + session | indgr + sessgr | 14 | -2046.77 | 4122.77 | 10.28% |
| age + sex + session | indgr | 12 | -2049.18 | 4123.27 | 8.01% |
| age + session | indgr + sessgr | 12 | -2049.55 | 4124.01 | 5.51% |
| b + indgr + session | indgr + sessgr | 15 | -2046.50 | 4124.43 | 4.48% |
| b + species + age + session | indgr + sessgr | 14 | -2049.44 | 4128.12 | 0.7% |
| age + session | sex | 9 | -2060.66 | 4139.84 | 0 |
| b + indgr + sessgr | indgr + sessgr | 11 | -2060.19 | 4143.14 | 0 |
before each removal session ($D_{\text{before}}$: 32.11–7.78 inds.ha$^{-1}$), lethal trapping achieved a 80–93% reduction in rat density after each of the four subsequent sessions ($D_{\text{after}}$: 0.56–2.56 ind.ha$^{-1}$) and allowed maintaining low numbers throughout a trapping period of seven months (Fig. 2, see Suppl. material 2). Densities of the two rat species remained constant from July to November 2015 (Fig. 2). A higher $R.\text{exulans}:R.\text{rattus}$ ratio was observed in the removal area as compared to the CMR area in May and September 2015 (Table 5).
Insights on reinvasion processes

During the last four trapping sessions (May to November 2015), contrasted patterns of rat juvenile proportions were observed between the CMR area, juveniles were recorded in highest numbers in May and July and then dropped to zero in September, while in the removal area, juveniles were recorded in lowest numbers in May, July and September before reaching highest numbers in November. Sex-ratios were balanced during May and July, but biased towards males both in September in the CMR area and in November in the removal area (Table 5). Heavier adult *R. rattus* and *R. exulans* were generally found in the removal area than in the CMR area (Table 5). The proportion of males with well-developed scrotal sacs increased earlier in the removal area (75% in July; Table 5).

In the removal area, average distances of trapped rats to the edge of the trapping grid (Dist \_rats \_\_\_r) fluctuated greatly during the four removal sessions. In July, six weeks after removal trapping had been initiated, most rats were captured near the edge of the removal trapping grid (Dist \_rats \_\_\_r ± SD = 11.98 ± 10.34 m), whereas in May, September and November, rat captures were distributed more evenly within the removal grid (Dist \_rats \_\_\_r = 19.98 ± 7.55 m; 31.81 ± 8.04 m; 22.79 ± 2.24 m, respectively; see also Suppl. material 3). A total of 27 individuals were captured in the CMR area at least once before and once after the beginning of the removal experiment. The barycenters of

| Table 4. Estimation of σ (m), 95% kernel home range (HR \(_{95}\), ha) and home range core area (HR \(_{50}\), ha). These estimations were performed for the four groups of individuals: *R. exulans*, *R. rattus* juveniles, *R. rattus* adult males, *R. rattus* adult females. σ was estimated from the following SECR model: \(g_0 \sim \text{indgr.} + \text{session, } \sigma \sim \text{indgr.} \). HR \(_{95}\) = \(\pi x (2.45 \times \sigma)^2\) and HR \(_{50}\) = \(\pi x (1.18 \times \sigma)^2\). |
|---|---|---|---|---|
| Group of individuals | \(\sigma\) | HR \(_{95}\) | HR \(_{50}\) |
| | mean | range | mean | Range | mean | range |
| *R. exulans* | 14.87 | 12.22–18.10 | 0.42 | 0.28–0.62 | 0.09 | 0.06–0.14 |
| *R. rattus* juveniles | 12.46 | 10.72–14.48 | 0.29 | 0.22–0.40 | 0.07 | 0.05–0.09 |
| *R. rattus* adult males | 19.97 | 17.31–23.04 | 0.75 | 0.57–1.00 | 0.17 | 0.13–0.23 |
| *R. rattus* adult females | 11.05 | 9.75–12.51 | 0.23 | 0.18–0.30 | 0.05 | 0.04–0.68 |

| Table 5. Rat population characteristics in both the rat removal and CMR trapping grids. These characteristics are given for the last four sessions of capture. Rr ad : *Rattus rattus* adults; Re ad : *R. exulans* adults. |
|---|---|---|---|---|---|---|
| Date | Trapping method | Number of rats trapped | Re : Rr ratio | Juvenile : Adult ratio | Male : Female ratio | Mean weight (± SD, g) | Mean weight \(Rr\) | Adult males with scrotal sac (%) |
| | | | | | | | Re ad. (± SD, g) | |
| 21–26 May 2015 | CMR | 74 | 0.24 | 0.35 | 0.95 | 158.51 ± 26.02 | 57.92 ± 5.59 | 2.86 |
| 28 May–05 June 2015 | Removal | 266 | 0.48 | 0.13 | 0.95 | 168.07 ± 30.04 | 62.24 ± 7.58 | 9.02 |
| 04–10 July 2015 | CMR | 36 | 0.50 | 0.42 | 1.00 | 147.00 ± 25.67 | 59.08 ± 7.80 | 22.22 |
| 11–15 July 2015 | Removal | 59 | 0.51 | 0.03 | 1.03 | 173.11 ± 39.64 | 66.97 ± 11.37 | 75.00 |
| 02–09 Sept. 2015 | CMR | 26 | 0.23 | 0 | 1.36 | 173.16 ± 34.34 | 67.83 ± 10.52 | 73.33 |
| 08–12 Sept. 2015 | Removal | 57 | 0.34 | 0.14 | 0.97 | 175.13 ± 29.98 | 79.82 ± 10.55 | 86.21 |
| 28 Oct.–01 Nov. 2015 | Removal | 65 | 0.47 | 0.32 | 1.62 | 177.26 ± 27.15 | 85.42 ± 9.41 | 76.32 |
Figure 3. Estimated barycenters of home range centers (HRC) of rats recaptured in the CMR grid. Only HRC of rats recaptured at least once in the CMR grid ($n = 27$), before (HRC$_{before}$) and after (HRC$_{after}$) rat removal had been initiated are presented here. Grey and black dots represent individual rat HRC barycenters before and after removal, respectively, while green and red filled circles represent the average HRC barycenters across individuals before and after removal, respectively. Twenty-seven other individuals first captured in the CMR area were recaptured in snap-traps in the removal area (red lines and black stars).
their home range center coordinates did not significantly shift towards the adjacent removal area in response to rat removal (X-axis barycenter coordinates, $Z = 198, p = .84$; Y-axis barycenter coordinates, $Z = 257, p = .11$) (Fig. 3). Furthermore, although rat control efforts appeared in the second best SECR model, this variable did not have any significant effect on $\sigma$ ($\beta = -0.21, SE = 0.16, 95\%CI = [-0.52–0.10]$) (Table 2).

A total of 27 rats equipped with PIT-tags in the CMR area were recaptured in the removal area over the course of the study (Fig. 3; see Suppl. material 4): 6 *R. exulans* (3 females: 1 adult and 2 juveniles, and 3 adult males) and 21 *R. rattus* (12 females: 8 adults and 4 juveniles, and 9 adult males). On average, adult male *R. exulans* and juvenile female *R. rattus* were the individual categories that travelled the longest distances, with respectively 138.30 ± 79.28 m and 145.01 ± 69.12 m travelled. Nineteen individuals travelled between 50 and 100 m and only six travelled > 100 m before being recaptured in the removal area. The longest distance was travelled by a *R. exulans* adult male (203.78 m in one day, from the CMR to the removal area).

**Simulated scenarios of control effort**

To simulate scenarios of removal efficiency relative to trapping session duration and project expenditure, we used the average value of parameters $g_0$ (0.09) and $\sigma$ (14.6 m) specifically estimated for our wire cage type of rodent live-trap, based on the best SECR model. With this simulation exercise and set of parameter values, we intend to demonstrate the usefulness of the proposed approach to compare the cost-effectiveness of plausible rodent management scenarios; however the approach could be easily adapted to other study systems, once key parameters are estimated for the relevant species and trapping devices.

If we were to ignore local social and economic constraints and assume that each trapping grid could be completely covered within one day, two grid layouts would allow achieving a cumulative rat capture probability $\geq .80$ after 15 trapping days (Fig. 4A): layout 1 (15 × 15 m; 1.00) and layout 2 (25 × 25 m; .95), while three other grid layouts would achieve a cumulative capture probability $\geq .50$: layout 3 (25 × 50 m; .75), layout 4 (25 × 75 m; .54) and layout 6 (50 × 50 m; .50). In our models, splitting the 200-ha grid into smaller adjacent management units allowed accommodating for these constraints, but in the meantime this strategy faced the risk of being too costly if trapping was carried out for a long period. Interestingly, however, when plotting the cumulative capture probability against cumulative project expenditure (in euros) over a theoretical scenario of 10 persons trapping over 30 days, it became apparent that the highest gain:cost ratio was achieved by layout 1 (.66: 24 900), layout 3 (.57: 22 700) and layout 2 (.63: 25 980) (Fig. 4B). Under this scenario, none of the ten different layouts reached a cumulative capture probability of .80. Given the local social constraints of 10 persons available for 15 days and 4 daily working hours per person, layouts 3 and 4 appeared to be the most cost-efficient, with a cumulative capture probability $> .30$ for a total cost < 17 200 euros (Fig. 4B). It is to be noted, however, that as layout 1 would require a total of 8 days to complete the initial baiting, this strategy could not be implemented if trappers were only available for 15 days.
Discussion

This study demonstrates that mechanical trapping can help maintain rat densities at low levels on a 5.5-ha area despite challenging environmental, logistical and social conditions. Combining live and lethal trapping experiments over 15 months in New Caledonian rainforest habitats has provided us with essential baseline rat biological parameters to inform effective management planning in tropical montane forests. Our cost-effectiveness analysis of trapping efforts also contributes to increase the evidence base that is currently lacking for improving the efficiency of rodent control projects and provides useful practical guidelines to practitioners involved in community-based pest management (Duron et al. 2017).

Rat population densities and home ranges in a tropical montane forest of New Caledonia

Rat population biology and dynamics on tropical rainforest islands remain less well understood than in other systems. Our CMR study showed great seasonal variation in rat densities, with a marked peak of ca. 34 ind.ha$^{-1}$ during the austral summer that
dropped three times lower in winter. In another study conducted in a New Caledonian lowland rainforest, rat densities were estimated at ca. 25 ind.ha$^{-1}$ in winter (Brescia 2011). Overall, rat densities in New Caledonia appear relatively high compared to average densities recorded in an Hawaiian mesic forest – $R$. rattus: 7.1 ind.ha$^{-1}$; $R$. exulans: .3 ind.ha$^{-1}$ (Shiels 2010); or in a low altitude New Zealand temperate forest – $R$. rattus: 6.2 ind.ha$^{-1}$ (Hooker and Innes 1995). By contrast, very high black rat densities (up to 65 ind.ha$^{-1}$ in summer) have been observed in the dry tropical forests of Europa Island in the Indian Ocean (Russell et al. 2011). In our study, black rat densities fluctuated seasonally, as observed elsewhere in the tropics, probably also in response to seasonal changes in food availability (Harris and Macdonald 2007; Russell et al. 2011). By contrast, densities of Pacific rats remained lower throughout our study, ranging from 0 to 4.3 ind.ha$^{-1}$, although a peak of ca. 17 ind.ha$^{-1}$ was detected in the first removal trapping session (May-June 2015). This pattern could be explained by negative interspecific interactions (e.g. Russell et al. 2004), in particular the competition release of Pacific rats from the removal of black rats over the course of the nine-day removal session, as indicated by the steady increase in Pacific rat capture rates as Black rat capture rate decreases. Some level of competition for traps may have occurred between species but given that trap saturation was relatively moderate in our CMR grid (average successful traps per session = 57.6%), it is unlikely that competition risk solely explains the observed differences in density between species.

In our study conducted in the Mont Panié wilderness reserve, black rat home ranges varied between .2 and .7 ha according to age and sex. This is much smaller than what was found in a Hawaiian mesic forest (3.8 ha; Shiels 2010), but more similar to the dry forest habitats of Juan de Nova and Europa Islands in the Indian Ocean (from .3 to .8 ha) (Ringler et al. 2014). Rat home range size is expected to vary widely in space and time in relation to in situ rat densities and the intensity of competitive interactions, the underlying food resource availability, and the timing of rat breeding season (Russell et al. 2011; Harper and Bunbury 2015; Harper et al. 2015). In our study, home ranges did not vary significantly across seasons, despite large temporal variations in rat densities. However, it is possible that we could not detect any seasonal variation in home range sizes due to insufficient recapture data by session. Care should be taken, however, when comparing home range size estimated from spatially-explicit capture-recapture and telemetry location data due to methodological differences and/or sampling error (Nathan 2016). Although SECR-derived $\sigma$ has been shown to underestimate telemetry-based home range radius, it still remains a useful estimate of space use as it concurrently incorporates movement and rat-trapping device interaction behaviours.

**Effects of a seven-month rat removal experiment on rat population densities in the Mont Panié wilderness reserve**

Controlling rats for 5–9 consecutive nights every five to seven weeks over a 5.5-ha area (on a 20 × 20 m trapping grid) allowed reaching an initial 16-fold density decrease (from 32.1 to 2.6 rats.ha$^{-1}$), followed by a five to fourteen-fold decrease (down to
Insights on rat reinvasion processes

A reduction in rodent abundance may be followed by a rapid reinvasion, induced by enhanced immigration and/or increased breeding and survival of remaining adults and juveniles (Innes et al. 1995; Gundersen et al. 2001; King et al. 2011; Hansen et al. 2020). In New Zealand, following the eradication of eight forest fragments averaging 5.3 ha in size, black rats were shown to travel up to 600 m in a few days and reinvade cleared areas very quickly despite large areas of intervening non-preferred habitat (King et al. 2011). In a Vietnamese farming system, the populations of *Rattus argentiventer* and *R. losea* were shown to respond to pest control mainly through an increase in recruitment, immigration and survival of residents, but not through enhanced breeding productivity (Brown and Tuan 2005). A recent study conducted on black rats in Australia showed that reinvasion of controlled areas was slow and occurred both through immigration and settlement by different categories of individuals: juveniles, adults or sub-adults without permanent home ranges, and individuals occupying neighboring areas expanding their home ranges (Hansen et al. 2020).

In our study, the level of control efforts applied (5–9 trapping days every 5–7 weeks) appeared to have prevented rapid, complete reinvasion of a 5.5-ha forest area. As our removal trapping only covered the austral winter, it is possible that reinvasion...
rates would have been higher during the subsequent summer period. We found, however, some evidence of rats travelling from the CMR area to the adjacent removal area already on the first removal trapping day, and then throughout the trapping period. Moreover, six weeks after the initial knock-down of the rat population, captures were mainly recorded along the edge of the removal area. Nevertheless, it remains unclear whether these scattered movements and spatio-temporal patterns of captures were driven by immigration, were a consequence of the expansion of existing home ranges in the adjacent CMR area, or simply reflect the higher capture probability of neighboring individuals dwelling in the near vicinity of the removal area. Our analyses of rat home range barrycenters and size (sigma) did not reveal any marked shift in space use nor any enlargement of home ranges within the trappable rat population of the CMR area in response to ongoing control efforts. Our data also suggest that some reestablishment may have been driven by density-dependent enhancement of breeding: a larger proportion of reproductively active males as well as young juveniles (R. rattus < 52 g) were detected at an earlier stage in the removal area (July 2015) compared to the CMR area. It is unlikely that these sex and age distribution patterns would have been explained by seasonal differences in trappabilities within individual rat groups since sex-and age-specific capture probabilities remained relatively constant over the course of the removal trapping experiment (see Suppl. material 2). Given the low numbers of rats captured in each of the three removal trapping sessions following the initial knock down, if enhanced breeding had occurred, it probably did not contribute to a substantial or sustained recovery of the controlled population.

Dispersal is commonly observed to be male-biased and principally realized by juveniles in most rodent species, including Rattus spp. (Brown and Tuan 2005; King et al. 2011; Krebs et al. 2011). Had immigration occurred in our removal area, we did not find any evidence for male rats being involved in the recolonization process more than females: the sex ratio observed in the removal area remained fairly balanced and constant as control efforts progressed, except during the last removal trapping session, six months after the initial knock down, when males represented 60% of captures. In addition, we did not find any support for juvenile-driven dispersal: juvenile numbers were very low in the removal area during each of the four removal trapping sessions. Without further evidence on the movements and origins of the rats captured in controlled areas (inferred from e.g. genetic analyses, rhodamine-B marking experiments), it remains unclear as to what processes typically govern rat population reestablishment in such contexts.

**Rat management recommendations**

Our modelling exercise indicated that it is, in principle, realistic to control invasive rats over moderate-sized areas in challenging environments, using lethal trapping. For example, in the absence of local and economic constraints, a one-shot reduction in rat density of 93% was achievable over 200 ha on a 15 × 15 m grid in a total of 1585 work hours or 5 trapping days (and a 100% reduction in 3169 work hours or 10 trapping days) (see Fig. 4A). This represented 385 (if 93%) or 792 (if 100%) man-work-days for an approximate operational budget of 22,000–38,000 euros per trapping session, assum-
ing four working hours per trapper per day. However, within the 150 man-work-days limit imposed by local workforces, none of the ten trapping grid layouts evaluated in this study could be practically implemented at the scale predefined by local stakeholders (200-ha pilot area). Treating the entire management area as smaller adjacent units that could be trapped in successive time stages was essential for achieving complete coverage. This strategy has been successfully implemented in the Scottish Outer Hebrides for the control of non-native American mink *Mustela vison* populations over large spatial scales (Macleod et al. 2019). While our simulations showed that splitting the entire management area would in theory enhance project feasibility, the limited workforce available in the Mont Panié wilderness reserve will inevitably hinder trapping effectiveness. Indeed, with 10 persons available for 15 days at a time, none of the grid layouts would achieve a cumulative rat capture probability higher than .30, which would probably not be sufficient to produce tangible benefits to native forest biodiversity. Alternative options to maximise removal efficiency may be to either reconsider the total size of the management area (e.g. 100 ha instead of 200 ha), if solely reliant on local workforces, or involve a few committed professional trappers to maintain a constant high trapping effort, although the latter would require additional financial resources over a protracted period of time. Other options include the use of self-resetting traps, which could be particularly useful in remote areas. The creation of “mainland islands” should also be considered as an alternative option to permanent rat removal, if and when resources to set up such an ambitious program are available. Ultimately, in areas where local workforce is limited and the use of aerial baiting is not accepted by the local communities, ensuring that the scale of management is fit for purpose is key to success. This decision should be based on the fine balance between maximising the use of available resources and achieving a cumulative capture probability that is meaningful for biodiversity targets.

With this study, we demonstrate that scenario modelling, informed by trapping experiments, is a flexible tool for informing the design of cost-effective control programs of island-invasive rodent populations, under idiosyncratic local circumstances. Due to rats’ productivity and reinvasion rates, a one shot reduction in density is clearly not enough to produce tangible benefits to native biodiversity. Given the prospect of assessing optimal strategies for a sustainable rat control program, acquiring rat demographic rates (productivity, survival, dispersal movements) will be helpful to explore finer components of trapping design (e.g. number of trapping sessions per season, year and habitat types) and inform on the most cost-effective trapping regime (how often to trap, for how long and where) in the long-term. While our modelling approach was developed for rats in the Mont Panié reserve, it could be easily adapted to other systems and invasive pest species that could be controlled by lethal trapping.

**Acknowledgements**

This study was funded by the Northern Province of New Caledonia to REFCOR project (Convention n° 12C240, 14C330 and 15C154). We are grateful to Josepho Bahomal, Hélène De Méringo, Oriana Garcia-Iriarte, Raphaël Gouyet, Matthieu Mativet,
Mathilde Méheut, Martin Thibault from IRD for their help in the fieldwork, and the University of Aberdeen for hosting a visit. We thank the team of Dayu Biik NGO, Alain Couhia, Ismaël Farino, Djaèk Folger, Ronald Tein, Aldo Tiempouène, Josine Tiavouane, Silvano Wanguène for field logistics and their help in the rat capture experiments. We also thank Frederic Rigault and Jéremy Anso for their help in preparing maps and figures, Murray Efford for support with the secr analyses, and Pablo Garcia Diaz for providing comments on a draft manuscript.

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Supplementary material 1

Density of rats
Authors: Quiterie Duron, Thomas Cornulier, Eric Vidal, Edouard Bourguet, Lise Ruffino
Data type: Statistics
Explanation note: Density of rats (ind.ha-1) according to capture session, species, sex and age. Density was estimated with SECR model: \( D \sim 1, g_0 \sim \text{indgr} + \text{session}, \sigma \sim \text{indgr} \).

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Link: https://doi.org/10.3897/neobiota.63.53811.suppl1

Supplementary material 2

Rat abundance and density, rat capture probabilities
Authors: Quiterie Duron, Thomas Cornulier, Eric Vidal, Edouard Bourguet, Lise Ruffino
Data type: Statistics
Explanation note: A) Rat abundance and density (ind.ha-1) before and after rat removal for the four sessions of rat removal trapping. The total number of individuals in the removal area was estimated with the “Zippin removal” method. Densities were estimated based on the rat removal grid size plus a boundary strip of 9 ha. B) Rat capture probabilities (± SE) estimated from removal trapping with the Zippin removal method by sex, age and session.
Supplementary material 3

Mean distances (± se) of trapped rats from the edge of the removal area during the four trapping sessions
Authors: Quiterie Duron, Thomas Cornulier, Eric Vidal, Edouard Bourguet, Lise Ruffino
Data type: Graphic results
Explanation note: Mean distances (± SE) of trapped rats from the edge of the removal area during the four trapping sessions. The grey dotted line represents the mean distance of the 209 traps to the edge of control area. When histogram bars are under the grey line, rats were captured closer to the edge compared to a scenario where rat captures would be evenly distributed across the grid.
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Link: https://doi.org/10.3897/neobiota.63.53811.suppl3

Supplementary material 4

Distances (in meters) travelled between rats’ home range centers in the CMR area and their recapture in the removal area for 27 individuals
Authors: Quiterie Duron, Thomas Cornulier, Eric Vidal, Edouard Bourguet, Lise Ruffino
Data type: Statistics
Explanation note: Distances (in meters) travelled between rats’ home range centers in the CMR area and their recapture in the removal area for 27 individuals, and mean (± SE) distances travelled according to species, sex and age.
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Link: https://doi.org/10.3897/neobiota.63.53811.suppl4