Dispersal is a critical parameter for successful pest control measures as it determines the rate of movement across target control areas and influences the risk of human exposure. We used a fine-scale spatial population genomic approach to investigate the dispersal ecology and population structure of Aedes notoscriptus, an important disease transmitting mosquito at the Mornington Peninsula, Australia. We sampled and reared A. notoscriptus eggs at two time points from 170 traps up to 5 km apart and generated genomic data from 240 individuals. We also produced a draft genome assembly from a laboratory colony established from mosquitoes sampled near the study area. We found low genetic structure (Fst) and high coancestry throughout the study region. Using genetic data to identify close kin dyads, we found that mosquitoes had moved distances of >1 km within a generation, which is further than previously recorded. A spatial autocorrelation analysis of genetic distances indicated genetic similarity at >1 km separation, a tenfold higher distance than for a comparable population of Ae. aegypti, from Cairns, Australia. These findings point to high mobility of A. notoscriptus, highlighting challenges of localised intervention strategies. Further sampling within the same area 6 and 12 months after initial sampling showed that egg-counts were relatively consistent across time, and that spatial variation in egg-counts covaried with spatial variation in Wright’s neighbourhood size (NS). As NS increases linearly with population density, egg-counts may be useful for estimating relative density in Ae. notoscriptus. The results highlight the importance of acquiring species-specific data when planning control measures.

Heredity (2023) 130:99–108; https://doi.org/10.1038/s41437-022-00584-4

INTRODUCTION

For insects that transmit disease, dispersal is a critical ecological characteristic to consider when planning management strategies. Dispersal will influence the movement of disease vectors into and out of controlled areas as well as the extent to which human, animal, or plant populations will be exposed to disease. Intrinsic and extrinsic factors such as dispersal barriers (natural or anthropogenetic) (Goldberg and Lande 2015; Schmidt et al. 2022), urbanisation (Johnson and Munshi-South 2015), and habitat fragmentation (Doak et al. 1992) can all influence insect dispersal. Once dispersal patterns are understood, they can be used to enhance the effectiveness of insect pest control strategies. For example, the larval dispersal pattern of the fruit orchard pest Operophtera brumata is used to indicate when farmers should implement control measures to mitigate the damage to plants while decreasing the use of insecticides (Edland 1983; Jeger 1999). Dispersal studies focusing on the primary vector of dengue, Aedes aegypti, have shown how its dispersal influences the spread and distribution of dengue, which has in turn shaped dengue control measures around the globe (Liew and Curtis 2004; Harrington et al. 2005; Carvajal et al. 2020).

Mosquitoes of the genus Aedes are important vectors of human and animal diseases. Their control traditionally involves insecticides (McCarroll et al. 2000), the removal of larval habitats, and the deployment of traps to reduce population sizes (Juarez et al. 2021). Novel interventions are now also applied, including the release of mosquitoes carrying novel Wolbachia bacteria (Hoffmann et al. 2011) or genetically modified mosquitoes (Harris et al. 2012) to reduce or manipulate populations. All these strategies rely on knowledge about the dispersal of the target species, which will influence the effectiveness and long-term success of control programs. For instance, in Ae. aegypti, Wolbachia stability and invasion into surrounding areas depends on mosquito movement across a local area (Schmidt et al. 2018).

Numerous methods have been deployed to estimate dispersal in Aedes. Mark-Release-Recapture (MRR) studies are commonly used (Reiter et al. 1995; Honório et al. 2003) but can come with drawbacks if applied to small organisms like mosquitoes. They can be labour intensive, require various ethical approvals, recapture rates can be low, and mosquitoes can be impacted by the marking method. Recent approaches try to overcome those difficulties by expanding MRR approaches to incorporate genetic inferences of close kin, functioning on the basis that an individual’s genotype can be considered a “recapture” of the genotypes of its relatives. This close kin mark-recapture (CKMR) framework has mainly been used to investigate abundance in large populations and to assess migration between populations (Palsøll et al. 2010). Recently, CKMR has been expanded to investigate dispersal, which it does...
by assigning dyads to kinship categories across multiple orders of kinship and then using the spatial distribution of these kin to reveal past dispersal over fine temporal scales (Schmidt et al. 2018; 2021; 2022, Combs et al. 2018; Fountain et al. 2018; Jasper et al. 2019; Trense et al. 2021; Aguillon et al. 2017). These recent studies reveal the power of novel genetic approaches to estimate individual dispersal in small organisms like Aedes mosquitoes. They also demonstrate an opportunity to use the genetic data acquired for additional analyses that go beyond dispersal, such as investigations of population structure and dynamics (e.g., neighbourhood size (Jasper et al. 2019) and levels of gene flow (Combs et al. 2018)).

Here, we use spatial population genomics to investigate the population structure and fine-scale dispersal of *Ae. notoscriptus* (the Australian backyard mosquito), a container breeding mosquito native to mainland Australia and Tasmania and invasive in the Torres Strait Islands, New Zealand, New Guinea, New Caledonia, Indonesia (Dobrotworsky 1965; Lee et al. 1987; Sunahara and Mogi 2004) and more recently in California, USA (Metzger et al. 2021). This species transmits arboviruses (e.g., Ross River virus and Barmah Forest virus) and dog heartworm (Russell and Geary 1992; Doggett and Russell 1997; Watson and Kay 1999) and is also suspected to be a significant vector of *Mycobacterium ulcerans*, the bacterium that causes Buruli ulcer (BU) (Wallace et al. 2021; Mee et al. unpublished data). Recent molecular studies have indicated several genetically different but morphologically similar populations of *Ae. notoscriptus* distributed across Australia (Endersby et al. 2013). In Victoria, two main clades occur in sympathy, and it is yet to be determined if different clades of *Ae. notoscriptus* differ in ecological characteristics such as habitat, feeding, and mating preferences, which can influence dispersal and the species’ ability to transmit diseases. Though this mosquito is an important vector, threatening human and animal health, little research focuses on its population dynamics in the field, which makes risk calculations and planning of interventions difficult. We present a draft reference genome for *Ae. notoscriptus* and use a genomic approach to investigate individual-level dispersal and spatial genetic structure of *Ae. notoscriptus* at the Mornington Peninsula, where BU cases have clustered in recent years (https://www.mornpen.vic.gov.au/Community-Services/Health-Wellbeing/Health-Safety/Buruli-Ulcer). We also calculate spatially-varying estimates of Wright’s neighbourhood size (NS: Wright 1946) throughout the study area, and compare these to egg counts to determine whether egg count data sufficiently reflect local *Ae. notoscriptus* abundance.

**MATERIAL AND METHODS**

**Reference genome assembly of *Aedes notoscriptus***

**DNA extraction and sequencing.** We extracted DNA of *Ae. notoscriptus* from a laboratory culture sampled initially in Cheltenham, Victoria, Australia, in 2018. DNA extraction was carried out using Qiagen genomic tip 20/g following manufacturer’s recommendations. An Illumina PCR-free library was then constructed using IDT Lotus kits and a fragment size of 250 bp. The library was then sequenced on Illumina HiSeqX 150 bp PE to obtain 136.22 Gb raw sequence data.

**Genome assembly.** The raw paired-end data was filtered to remove adapter sequences using TrimGalore v0.6.6 (Krueger 2021) resulting in 135 Gb of clean data. This was then assembled using MoSuRCA v3.4.0 (Zimin et al. 2013) and SparseAssembler v1 (Ye et al. 2012) to obtain high fidelity contigs, which were then merged using flye v2.7 (Kolmogorov et al. 2019). This merged assembly was then deduplicated using purge_dups v1.2.4 (Guan et al. 2020) to create the draft assembly for analysis spanning 1.55 Gb with an N50 of 3893 bp.

**Fine scale population structure and dispersal**

**Sampling of *Ae. notoscriptus***. In February 2019, we sampled *Ae. notoscriptus* from four sites of similar area in the Mornington Peninsula: Sorrento (‘north-west’), Blairgowrie (‘central’) and Rye (‘north-east’ and ‘south-east’) (Fig. 1). Sampling used 340 oviposition traps (~90 traps per sampling site) which were placed randomly within each site. The average distance between traps was 1046 m (ranging from 16 to 5480 m).

Each trap consisted of a black plastic bucket half-filled with water and containing several alfalfa pellets to attract gravid *Ae. notoscriptus* (Ritchie 2001). A ~15 cm long strip of red felt extending into the water provided an oviposition substrate. We collected felts after 7 and again after 14 days, and partially dried them to encourage hatching. Three days after collection, we hatched eggs in 500 mL reverse osmosis (RO) water containing 2–3 TetraMin tropical fish food tablets (Tetra, Melle, Germany). If no larvae hatched, felts were re-dried for three days and the hatching process repeated. We replaced water and food as appropriate. Emerging virgin adults were transferred into absolute ethanol and stored at −20 °C until DNA extraction. We selected 240 individual mosquitoes for DNA sequencing, covering 170 different trap locations. To maximise spatial distribution, we prioritised samples collected from different traps over samples from different weeks. Care was taken to only include one individual per felt to avoid the detection of close kin sampled from the same felt.

**Fig. 1 Sampling sites at the Mornington Peninsula.** ‘north-west’, ‘central’, ‘south-east’ and ‘north-east’ sites from left to right indicated by red rectangles. Transparent black circles indicate trap locations in each sampling site. Trap placement is visualised in more detail in Fig. 4.
DNA extraction and library preparation. Using keys from Dobrotworsky (1965) we morphologically identified Ae. notoscriptus mosquitoes and extracted DNA from individuals using either Qiagen DNasey Blood & Tissue Kits (Qiagen, Hilden, Germany) or Roche High Pure PCR Template Preparation Kits (Roche Molecular Systems, Inc, Pleasanton, CA, USA) following the manufacturers’ instructions. We prepared double-digested restriction site-associated DNA (rcsDNA) libraries (Ihle and Urban 2007) and followed the method for the Ae. aegypti (Rašić et al. 2014). We started with an initial digestion of 10 ng of genomic DNA, ten units each of MluCl and NalII restriction enzymes, NEB CutSmart buffer (New England Biolabs, Beverly, MA, USA), and water. Digestions were run for 3 h at 37 °C with no heat kill step, and the products were cleaned with paramagnetic beads. Modified Illumina Plink and MiniPlink (Purcell et al. 2007) were ligated onto cleaned Illumina adapters overnight at 16 °C with 1000 units of T4 ligase (New England Biolabs, Beverly, MA, USA), followed by a 10-min heat-deactivation step at 65 °C. We performed size selection using a Pippin-Prep 2% gel cassette (Sage Sciences, Beverly, MA) to retain DNA fragments of 350–450 bp.

The size selected libraries were amplified by PCR, using 1 μL of size-selected DNA, 5 μL of Phusion High Fidelity 2× Master mix (New England Biolabs, Beverly MA, USA) and 2 μL of 10 μM standard Illumina P1 and P2 primers. These were run for 12 PCR cycles, then cleaned and concentrated using 0.8x paramagnetic beads. Each ddRAD library contained 24 mosquitoes, and each was sequenced on a single sequencing lane using 150 bp chemistry. Libraries were sequenced paired-end at GeneWiz, Inc (Suzhou, China) using a HiSeq 4000 (Illumina, California, USA).

Data processing. We used the Process_radtags program in Stacks v2.0 (Catchen et al. 2013) to demultiplex sequence reads. Low-quality reads were discarded using a 15 bp sliding window if the average Phred score dropped below 20. We used Bowtie v2.0 (Langmead and Salzberg 2012) to align reads to the Ae. notoscriptus reference genome assembly (described in 2.1), using –very-sensitive alignment settings. We filtered all alignments to paired reads that aligned concordantly, requiring the two paired reads to align to the same contig to avoid multi-mapping using Samtools (Danecek et al. 2021). Stacks_Ref_map program was used to build Stacks catalogs, from which genotypes were called at a 0.05 significance level and –min-mapq 15 to filter any remaining multi-mapped reads. We generated VCF files for the catalog with the Stacks program Populations (Catchen et al. 2013). Single nucleotide polymorphisms (SNPs) were required to be genotyped in ≥90% of mosquitoes, with a minor allele count of ≥3 (R 0.90 –mac 3 –nvaf). Beagle v4.1 (Browning and Browning 2016) was used to impute and phase the dataset in a 50,000 bp sliding window within 3000 bp overlap. Finally, vcftools thinned SNPs so that no two SNPs are at 500 bp distance to each other (~thind 500). After filtering, we retained 11,091 SNPs.

Genetic diversity and local population structure. Populations was used to calculate pairwise Fst values between the four sampling sites. We tested for isolation by distance between all samples and within each of the four sampling sites using the mantel.randtest function in the ade4 package of R (Dray and Dufour 2007). We also tested for isolation by distance in the dataset separating pairs of individuals identified as close kin, as close kin can drive isolation by distance patterns at sufficiently fine scales (Aguillon et al. 2017). The simple Mantel tests analysed pairwise genetic distance matrices and the natural logarithm of Haversine pairwise geographic distance, employing 9999 permutations and Bonferroni correction to assess statistical significance. Roussieu’s a (Roussieu 2000) provided genetic distances, calculated in SPAGEDi (Hardy and Vekemans 2002). Additionally, we used the pairwise genetic distance and geographical distance matrices to measure spatial autocorrelation using the mgram function in the R package ecocat (Goslee and Urban 2007) to build correlograms.

To contextualise these spatial genetic structure results, we compared results for Ae. notoscriptus with an Ae. aegypti population from Cairns, Australia, sampled in 2015 using similar protocols and with a similar spatial distribution of traps (Schmidt et al. 2018). This Ae. aegypti dataset contained both individuals carrying a Wolbachia infection (wMel) from recent releases in the area and wildtype individuals without the infection (WT). We analysed these separately to avoid bias. We down sampled each Ae. aegypti dataset only to include one sample per trap to achieve maximum comparability to the Ae. notoscriptus dataset. Additionally, we included the mgram analysis on two sets of the Ae. notoscriptus dataset separated into the ‘north-west’ and ‘central’ as well as ‘north-east’ and ‘south-east’ samples sites to archive a more comparable trap distance distribution to the Ae. aegypti datasets.

To further investigate the local population structure of Ae. notoscriptus and coancestry between individuals of the different sampling sites, we used the program fineRADstructure (Malinsky et al. 2018), which we ran with default settings.

Local dispersal estimates of individuals. We investigated the association between pairwise kinship and distance to infer specific dispersal of the population. We used the very-sensitive alignment settings. We filtered all alignments to paired reads that aligned concordantly, requiring the two paired reads to align to the same contig to avoid multi-mapping using Samtools (Danecek et al. 2021). We generated VCF files for the catalog with the Stacks program Populations (Catchen et al. 2013). Single nucleotide polymorphisms (SNPs) were required to be genotyped in ≥90% of mosquitoes, with a minor allele count of ≥3 (R 0.90 –mac 3 –nvaf). Beagle v4.1 (Browning and Browning 2016) was used to impute and phase the dataset in a 50,000 bp sliding window within 3000 bp overlap. Finally, vcftools thinned SNPs so that no two SNPs are at 500 bp distance to each other (~thind 500). After filtering, we retained 11,091 SNPs.

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was chosen as providing the best fit. Results returned by the Kriging model were cross-validated using the `xvalid` function. We visualised results through the `image` function. In addition, we estimated NS for the entire dataset, using the inverse of the regression slope of pairwise individual genetic distance (Rousset’s $a$) against the natural logarithm of geographical distance (Rousset 2000). We omitted pairs < 100 m apart as the linear relationship of genetic and geographical distance may break down at distances within the dispersal estimate $\sigma$.

**RESULTS**

**Population structure and dispersal**

*Genetic diversity and local population structure*. Pairwise $F_{st}$ estimates were lowest between the ‘north-east’ and ‘south-east’ sites and highest between the ‘north-west’ and ‘south-east’ sites (Table 1; Fig. 1). All $F_{st}$ estimates were very low.

The fineRADstructure plot (Fig. 2) showed no clear clustering of coancestry between mosquito pairs collected from the same sampling sites, indicating gene flow between all sites.

We found isolation by distance in the positive relationship between genetic distance and the natural logarithm of geographical distance when calculated on the entire *Ae. notoscriptus* dataset (Bonferroni-corrected $P$-value $= 0.02$, $r = 0.05$, mean distance between trap pairs $= 2045$ m) (Fig. S3). However, when related individuals were removed from the dataset the relationship weakened ($P$-value $= 0.07$, $r = 0.09$), indicating that isolation by distance at the scale of the whole sampling area is mostly driven by closely related individuals. No isolation by distance was detected when tested within each sampling site, where the mean distances between trap pairs were $522$ m, suggesting no clear genetic structure for *Ae. notoscriptus* at this scale.

We found positive and significant spatial autocorrelation among *Ae. notoscriptus* samples around the range of $1300$ m ($r = 0.02$, $p = 0.005$) as well as at $3700$ m ($r = 0.03$, $p = 0.02$) (Fig. S5). At distances ranging from $4700$ m onwards, *Ae. notoscriptus* showed significantly negative autocorrelation ($r = −0.05$, $p < 0.001$) where individual mosquitoes were effectively less related than they would be at random. By comparison, the population of *Ae. aegypti* from Cairns showed a signature of spatial structure that was

| Site          | North-west | Central | South-east | North-east |
|---------------|------------|---------|------------|------------|
| North-west    | 0.00163    | 0.00165 | 0.0013     |            |
| Central       | 0.0008     | 0.0008  |            |            |
| South-east    |            |         | 0.0005     |            |

All standard errors $= 0.00000$. Site locations are visualised in Fig. 1.

Fig. 2  FineRadstruture coancestry map and tree. The left-hand side panel indicates genotype sampling site (‘north-west’ in magenta; ‘central’ in green; ‘south-east’ in light blue; ‘north-east’ in grey). The central panel shows co-ancestry between genotypes, with light yellow indicating low co-ancestry, and darker yellows, oranges and reds indicating progressively higher co-ancestry.
Fig. 3 Spatial autocorrelation (left) and density of trap distances (right) of Aedes notoscriptus ‘north-west’ + ‘central’. A Aedes notoscriptus ‘north-east’ + ‘south-east’ B WT (wild type = uninfected) Aedes aegypti C and wMel infected Aedes aegypti D Mantel correlograms on the left depict the association between genetic distance and geographical distance among pairs of the same distance class of 100 m. Error bars show 95% confidence intervals and were calculated using 999 bootstrap replicates. Significant associations (α = 0.05) are shown as filled circles. Histograms graphs on the right describe the pairwise distances between traps of the three datasets. Ae. aegypti data is from Schmidt et al. (2018).
positive and significant in the first distance class of 100 m (WT = wild type: \( r = 0.08, p = 0.02 \); wMel = Wolbachia infected: \( r = 0.05, p = 0.001 \)), then decreased sharply, dropping below zero beyond 500 m (Fig. 3B, C). In Ae. aegypti, significant negatively autocorrelated values were observed from around 1100 m (WT: \( r = -0.104, p = 0.005 \); wMel: \( r = -0.05, p = 0.01 \)). The range of trap distances for WT Ae. aegypti was 35 m to 3280 m and 36 m to 3057 m for the wMel infected population (Fig. 3B, C).

When separated into two datasets (as described in 2.2.4), Ae. notoscriptus does not show any significant negative autocorrelation (Fig. 3), indicating that the larger trap distances in the full Ae. notoscriptus dataset does not bias the original analysis towards larger distances.

**Local dispersal estimates of individuals.** We identified three putative full-sibling and 8 half-sibling pairs using PC-Relate. We also designated 8 pairs with \( k > 0.07 \) as putative first cousins. Jasper et al. (2019) estimated that if the local population is constant, twice as many cousins than sibling pairs can be assumed as an average of two offspring from any one individual will themselves have offspring. However, here we present cousin pairs that we could confidently distinguish from unrelated individuals, meaning that cousin pairs are likely underrepresented.

Full-siblings were separated by a mean distance of 466 m (median = 179 m) and exhibited a maximum observed distance of 1267 m. We found a mean separation distance for half-siblings of 1296 m (median = 340 m, max = 5173 m), and 2778 m (median = 2825 m, max = 4664 m) for first cousins (Fig. 4).

We detected two full-sibling pairs within a range of 24 m to 179 m while the third full-sibling pair was found 1154 m apart (Fig. 4). Most half-sibling pairs were detected within 0 m and 700 m, and several pairs were found between 1247 m and 2739 m (Fig. 4). We found pairs of first cousins at a mean separation distance of 1400 m, with the largest distance being 5186 m (Fig. 4).

**Graphical analyses of egg counts and neighbourhood size**

**Egg counts.** While ordinary kriging was performed on cube-transformed egg counts, numbers per trap are visualised after back transformation to present observed egg counts per trap (Fig. 5A, B). The plots show a pattern of increased egg counts from ‘east’ to the ‘north-west’ sites. The estimated prediction error after cross validation indicates that on average, an error of \( \pm 1.5 \) eggs can be expected at any given location. Patterns of spatial variation in egg counts were consistent across the two time points (Nov 2019 and Feb 2020), with higher egg counts being observed in the ‘north-west’ compared to the ‘north-east’ and ‘south-east’.

**Neighbourhood size.** The NS for each trap location estimated using sGD is shown as coloured circles in Fig. 6, which also shows the Kriging predictions of NS throughout the area, calculated in geoR. The map shows lower NS (190–230) in the ‘south-east’ and ‘south-west’ sites, with an increase in NS in the ‘central’ (240–265) and ‘north-west’ (260–285) sites. The estimated prediction error for ordinary kriging after cross validation indicates that, on average, an error of \( \pm 1.04 \) of mean NS can be expected at any given location. Spatial variation in NS was roughly consistent with egg counts (Figs. 5, 6), where higher values were observed in the ‘north-west’ and lower values in the ‘south-east’. NS estimates produced by sGD were smaller than those estimated from the inverse of the regression slope, with an estimated NS = 383 mosquitoes (95% CI 287–572) across the study area.

**DISCUSSION**

Recent studies applying spatial genomics methods to Aedes mosquitoes have recorded how Ae. aegypti disperse within and between buildings in urban Malaysia (Jasper et al. 2019) and have identified dispersal barriers in Ae. aegypti from Saudi Arabia (Schmidt et al. 2022). This study used population genomic approaches to investigate fine-scale dispersal in an Australian mosquito, Ae. notoscriptus, at the Mornington Peninsula, Australia, where the species is strongly suspected to be involved in transmitting *M. ulcerans* to humans. This represents the first application of genomic methods to Ae. notoscriptus, and we report both a draft genome assembly for this species and results describing how genome-wide genetic variation is spatially distributed in the Mornington Peninsula population. Our findings suggest Ae. notoscriptus can move greater distances than previously reported and appears to be a stronger disperser than Ae. aegypti. Additionally, we found that spatial genetic variation covaried with variation in egg counts across two seasons, evidence that ovitrap data may be a useful proxy for local Ae. notoscriptus density.

The locations of the full-sibling pairs that we detected in our kinship analysis can be interpreted as direct movement of a female mosquito that moved between the two traps during
oviposition. Though we found two of the three full-sibling pairs at distances of 24 m and 179 m, one full-sibling pair was detected >1 km apart (Fig. 4). This distance (i.e., 1400 m) is 5-fold larger than the farthest distance reported by MRR (i.e., 238 m) (Watson et al. 2000; Trewin et al. 2019). In these studies, the maximum distance of traps from the release point was 275 m, so dispersal over greater distances than this would not have been detected. Conducting MRR studies over greater distances becomes far more labour-intensive, which makes close-kin methods such as in this study a valuable alternative for assessing dispersal at these scales. While we reported an average separation distance of 440 m for full-siblings, it is important to note that trap placement (i.e., distances between traps) and sample selection (i.e., one individual per trap) could have prevented us detecting some smaller separation distances, which would bias dispersal estimates upwards (Jasper et al. 2022).

We detected most half-sibling pairs within a range of 0 m (found in the same trap, one week apart) to around 700 m, while some half-sibling pairs were found multiple kilometres apart (1247 m and 2739 m; Fig. 4), highlighting common mid-range dispersal (i.e., mean separation distance = 832 m). We found most first cousin pairs at a mean separation distance of 1400 m with the biggest distances being >5 km, which indicates that it takes approximately two generations for Ae. notoscriptus to disperse through the entire study area. Overall, the results of our kinship analysis present evidence for a high dispersal activity of Ae. notoscriptus at the Mornington Peninsula, also supported by the low Fst values (Table 1) between sampling sites. Likewise, the lack of clustering in the coancestry analysis provided by finRAdstructure (Fig. 2) indicates a lack of genetic structure among sites, which reflects high gene flow between different sites, which also points towards high dispersal capacity of Ae. notoscriptus in the study area.

Studies of dispersal and population structure in Aedes mosquitoes have previously mainly focused on the dengue vector Ae. aegypti (e.g., Muir and Kay 1998; Harrington et al. 2005; Rasheed et al. 2013; Schmidt et al. 2022; Kotsakiozi et al. 2018; Jasper et al. 2019). This species’ reportedly limited dispersal has been used as a proxy for the dispersal of related container breeding mosquito species. Past MRR studies investigating
dispersal in adult *Ae. notoscriptus* in urban environments of Queensland present contradictory results. Watson et al. (2000) describe similarly limited dispersal of *Ae. notoscriptus* compared to previous dispersal studies in *Ae. aegypti*, while Trewin et al. (2019) directly compared the two species in an MRR study and found that *Ae. notoscriptus* disperses further than *Ae. aegypti*. Both studies were conducted in urban areas of Queensland, with recapture trap distances ranging from 25 m to 265 m (Trewin et al. 2019) and 275 m (Watson et al. 2000). However, both studies indicate that, *Ae. notoscriptus* seems less restricted by barriers such as roads compared to *Ae. aegypti*. While our data are not suitable for directly estimating dispersal distance in *Ae. notoscriptus*, as we did not have a large enough set of close kin dyads (Jasper et al. 2022) or an estimate of density (Rousset 2000), if *Ae. notoscriptus* dispersal is greater than *Ae. aegypti* this would suggest it moves greater than ~200 m per generation (Guerra et al. 2014; Schmidt et al. 2022).

To further investigate differences in population structure and dispersal of the two species, we compared the results of our spatial autocorrelation analysis to two comparable datasets of *Ae. aegypti* (see 2.2.4). Our analysis showed genetic dissimilarities between *Ae. notoscriptus* were only significant at greater than 4 km separation, while *Ae. aegypti* had significant dissimilarities from around 1 km, which suggests greater dispersal activity in *Ae. notoscriptus* (Fig. S6) in comparable urban environments within Australia. Spatial autocorrelation in both the wildtype and wMel infected *Ae. aegypti* revealed evidence of a strong pattern of localised genetic structure, pointing towards limited dispersal in this species. Because *Ae. notoscriptus* showed a strong positive Mantel correlation at more than tenfold larger distances than the observed correlation shown in *Ae. aegypti*, dispersal seems less restricted in *Ae. notoscriptus* and dispersal of distances >500 m may commonly occur (Fig. 3). Isolation by distance detected in both *Ae. aegypti* datasets strengthens this interpretation, while we detected no isolation by distance in *Ae. notoscriptus* at similar spatial scales (i.e., within sampling sites). We detected isolation by distance in *Ae. notoscriptus* on the scale of the entire dataset (i.e., across all sampling sites) and calculated a neighbourhood size (NS: Wright 1946) of 287–572, which is consistent with NS calculated in *Ae. aegypti* (Jasper et al. 2019) and *Ae. albopictus* (Schmidt et al. 2021) when calculated using the inverse of the regression slope. NS calculated by sGD incorporated spatial variation in NS throughout the area and returned neighbourhood sizes somewhat lower.

The observed differences in dispersal behaviour of different *Aedes* species could be explained by differences in flight abilities, feeding and breeding preferences, and the likelihood of being passively moved (Verdonschot and Besse-Lototskaya 2014). While both species have adapted to breed predominantly in artificial containers, there are differences in other critical ecological factors between *Ae. aegypti* and *Ae. notoscriptus*. While *Ae. notoscriptus* does seek human hosts for blood-feeding, this species reportedly feeds on other animals such as dogs, birds, horses, possums, and fruit bats (Kay et al. 2008), while *Ae. aegypti* is highly anthropophilic (Harrington et al. 2001). This difference in host preferences may contribute to the higher dispersal activity observed in *Ae. notoscriptus*, as this species is not reliant on human blood meals to breed successfully, and hence can move further from human proximity. Additionally, MRR studies that investigated *Ae. notoscriptus* dispersal found exceptionally low male recapture rates compared to other *Aedes* species (Watson et al. 2000; Trewin et al. 2019), which could mean that male *Ae. notoscriptus* respond to different cues than other *Aedes* mosquitoes, which can influence dispersal. However, the described ecological and behavioural features of *Ae. notoscriptus* could likely differ between different clades of this species, leading to contrasting results of studies investigating traits in different populations and locations and should therefore be interpreted with caution.

Our results and analyses indicate a high dispersal ability of *Ae. notoscriptus*, which is vital to consider when planning control strategies. Trewin et al. (2019) argue that while *Ae. aegypti* may be controlled on the level of small blocks, the control of a highly dispersive container breeder such as *Ae. notoscriptus* will likely require much bigger areas, which is expected to be expensive and labour intensive. A pilot mosquito control study conducted in early 2021 aimed to reduce the population size of *Ae. notoscriptus* at the Mornington Peninsula non-chemically to further investigate the role of the species in the transmission of Buruli ulcer (https://www.health.vic.gov.au/infectious-diseases/behaving-buruli-in-victoria; Mee et al. unpublished data). The study employed gravid traps for four weeks to remove female mosquitoes and their offspring from the environment. The trial was evaluated by egg counts collected before and after the intervention and resulted in no measurable reduction in numbers of *Ae. notoscriptus* eggs. The success of the intervention could have been affected by several factors, including low uptake of gravid traps by residents, but was also likely affected by the size of the controlled zones. The

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Fig. 6 Ordinary Kriging of Neighbourhood size (NS) throughout the study area at the Mornington Peninsula. Neighbourhood size was calculated for each trap location using sGD in R. Trap locations are plotted as circles. Colours indicate NS. Distribution of NS numbers kriging is based on shown in the top-right panel. NS was calculated from genetic data from individuals collected in February 2019.
intervention zones (~250 x 350 m) were sized under the reasonable assumption that Ae. notoscriptus dispersal in urban areas is similar in scale to Ae. aegypti. However, our results highlight that generalising ecological traits between related species can be problematic when planning species-specific interventions. The dispersal estimates we discussed above indicate that it is likely that the dispersal of Ae. notoscriptus exceeded the size of controlled zones. As a result, zones were likely to be invaded by mosquitoes from surrounding areas while control measures were in place, which could have compromised the success of lowering mosquito numbers. Controlled zones were also likely to be re-invaded quickly after the trial, given the generation time of Ae. notoscriptus of approximately one month.

Finally, the consistent egg counts observed over both time points suggests that egg counts may provide a useful indicator of local abundance. However, local variation in egg numbers was substantial, highlighting the importance of trapping at high densities when assessing local population density with egg numbers. As applied in this study, oviposition traps represent a useful tool for this application as they are cheap, readily available, and easy to set up. We also found a trend between egg count variation and spatially explicit estimations of NS, with egg numbers and NS predictions showing lower values in the ‘north-east’ and ‘south-east’ sampling sites with increasing values through the ‘central’ to the ‘north-west’ sites at both time points (i.e., November 2019 and February 2020) (Figs. 5, 6). Though NS calculations showed much less variation throughout the area than egg counts, this comparison indicates that higher egg counts correlate with larger NS and, therefore, density of mosquitoes. Our comparison of egg counts to mosquito densities matches previous studies that present a positive relationship between egg numbers collected by oviposition traps and numbers of adult mosquitoes collected in adult traps (Tantowijoyo et al. 2016; Feria-Arroyo et al. 2020).

CONCLUSIONS

This study uses recently developed genomic approaches to provide evidence for high dispersal abilities of Ae. notoscriptus at the Mornington Peninsula, Australia. Dispersal in this population seems to take place over greater scales than in the Cairns population of the congeneric Ae. aegypti, which highlights the importance of acquiring species-specific and even population-specific ecological information when planning management strategies. We showed the challenge of generalising between related species when planning interventions. Finally, we provide evidence that spatial variation in egg counts collected by oviposition traps can be linked to spatial variation in neighbourhood size and, thus also, density and may be a valid means of estimating relative densities and the impact of control measures.

DATA AVAILABILITY

Data has been archived at Dryad https://doi.org/10.5061/dryad.05qft8f60. Aeodes notoscriptus genome available at https://doi.org/10.25391/9zs9w-hw85.

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ACKNOWLEDGEMENTS
We thank Nancy Endersby-Harshman and Qiong Yang for laboratory assistance and Moshe Jasper and Nancy Endersby-Harshman for helpful discussions.

AUTHOR CONTRIBUTIONS
TLS, AAV and VP designed the study. TLS, SEL and PTM led field collections. VP did molecular work and data analysis with input from TLS. IRV developed genome. VP wrote the manuscript with TLS and AAV, with input from all authors.

FUNDING
This research was funded by a University of Melbourne Early Career Researcher grant awarded to TLS and by the NHMRC Partnership project 1196396. VP was financially supported by the Australian Government Research Training Program Scholarship. TLS was supported by Program Grant 1132412. AAV was supported by the NHMRC grant 1118640.

COMPETING INTERESTS
The authors declare no competing interests.

ADDITIONAL INFORMATION
Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41437-022-00584-4.
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