Development and field evaluation of the sentinel mosquito arbovirus capture kit (SMACK)

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Abstract

Background: Although sentinel animals are used successfully throughout the world to monitor arbovirus activity, ethical considerations and cross-reactions in serological assays highlight the importance of developing viable alternatives. Here we outline the development of a passive sentinel mosquito arbovirus capture kit (SMACK) that allows for the detection of arboviruses on honey-baited nucleic acid preservation cards (Flinders Technology Associates; FTA®) and has a similar trap efficacy as standard light traps in our trials.

Methods: The trap efficacy of the SMACK was assessed against Centers for Disease Control and Prevention (CDC) miniature light traps (standard and ultraviolet) and the Encephalitis Vector Survey (EVS) trap in a series of Latin square field trials conducted in North Queensland, Australia. The ability of the SMACK to serve as a sentinel arbovirus surveillance tool was assessed in comparison to Passive Box Traps (PBT) during the 2014 wet season in the Cairns, Australia region and individually in the remote Northern Peninsula Area (NPA) of Australia during the 2015 wet season.

Results: The SMACK caught comparable numbers of mosquitoes to both CDC light traps (mean capture ratio 0.86:1) and consistently outperformed the EVS trap (mean capture ratio 2.28:1) when CO2 was supplied by either a gas cylinder (500 ml/min) or dry ice (1 kg). During the 2014 arbovirus survey, the SMACK captured significantly more mosquitoes than the PBT, and 2 and 1 FTA® cards were positive for Ross River virus and Barmah Forest virus, respectively, while no arboviruses were detected from PBTs. Arbovirus activity was detected at all three surveillance sites during the NPA survey in 2015 and ca. 27% of FTA® cards tested positive for either Murray Valley encephalitis virus (2 detections), West Nile virus (Kunjin subtype; 13 detections), or both viruses on two occasions.

Conclusions: These results demonstrate that the SMACK is a versatile, simple, and effective passive arbovirus surveillance tool that may also be used as a traditional overnight mosquito trap and has the potential to become a practical substitute for sentinel animal programs.

Keywords: Arboviruses, Sentinel animals, Mosquitoes, Passive Traps, Surveillance

Background

Sentinel animals have long been used throughout the world to monitor arbovirus activity and have been employed in Australia since the late 1960s [1–3]. Many Australian states/jurisdictions continue to use sentinel chickens to detect Murray Valley encephalitis virus (MVEV) and West Nile virus (Kunjin subtype; WNV_KUNJ) while sentinel pigs have been employed to monitor Japanese encephalitis virus in the Torres Strait and the Cape York Peninsula (JEV) [4, 5]. Despite the ability of sentinel animals to detect arbovirus activity, there are many difficulties with their use. For instance, there are ethical and logistical implications associated with using the animals, as well as challenges in detecting closely related viruses due to cross-reactions in serological assays and a limited ability to only target viruses that infect the selected sentinel animal or those that are transmitted by vectors that feed on the sentinel animals [6, 7]. Given these...
limitations, a strategy has been developed using CO$_2$-baited mosquito traps that house sugar-soaked nucleic preservation cards [6]. While an infected mosquito probes during sugar-feeding virus is expectorated onto the cards, which are subsequently tested for the presence of viral RNA using molecular assays.

To overcome issues with powered traps, such as component malfunction and requirement for electricity to power trap fans and lights, a CO$_2$-baited passive (non-battery powered) box trap (PBT) was developed by Ritchie et al. [8]. The PBTs were used to house the honey-soaked cards in a field trial in northern Australia, where multiple arboviruses were detected [9]. Based on the the McPhail fly trap [10] the PBT does not rely on battery-powered fans to capture host-seeking females but instead simply utilises the attractiveness of CO$_2$ and the passive retention of captured mosquitoes within a translucent plastic crate. Despite the utility of the PBT as a cheap and efficient passive surveillance device, it may underperform when compared against standard light traps [11] and there are still design principles in need of improvement for long-term deployment. For instance, field trials suggest that keeping mosquitoes alive for several days post capture can increase detection of viruses on the cards [12].

In the current paper, we describe the development and field evaluation of the Sentinel Mosquito Arbovirus Capture Kit (SMACK). The SMACK consists of a CO$_2$-baited passive trap that improves upon the original PBT design, particularly modifications to enhance mosquito collection efficacy, mosquito survival post capture, and increased sugar-feeding on honey-baited nucleic acid preservation cards. We then assess the efficacy of the SMACK as a mosquito surveillance device against three standard battery-powered light traps. Finally, we assessed the efficacy of the SMACK for use as a sentinel arbovirus surveillance tool against an unmodified PBT during the 2014 wet season and when deployed as a surveillance tool in the remote Northern Peninsula Area (NPA) of Australia during the 2015 wet season.

**Methods**

**General trap design**

A 20 L translucent plastic storage box (29 × 37, 27 cm deep) (Icon Plastics, Victoria, AU) with a clip-on lid was chosen as the main body of the SMACK, as this size was previously shown to outperform smaller PBT designs [8] (Fig. 1a). Flinders Technology Associates (FTA*) cards (Whatman International Ltd, Maidstone, UK) were placed in 70 ml collection jars in which the bottom had been cut so that the FTA* card was exposed to the mosquitoes. To reduce the desiccation of the honey solution on the FTA* cards, the collection jars contained sponge material (DTA Australia, Victoria, AU) that was soaked in 50 % honey (diluted in distilled water) immediately before the FTA* card was added. This method also allows the cards to be easily inserted and removed from the trap without having to access the cards from inside the trap by anesthetizing captured mosquitoes. The PVC ventilation pipe on top of the PBT described by Ritchie et al. [8] was omitted in the SMACK design and instead a 10 mm hole was created to enable rubber gas tubing to be inserted through the top of the trap. A 12 x 25 mm air stone (Aqua Nova, Petras Fisheries Pty Ltd, Sydney, AU) was attached...
to the end of the tubing inside the trap to disperse CO₂ over the sieve. A water reservoir was attached to the inside of the trap and consisted of a 500 ml plastic container (Tellfresh®, Victoria, AU) in which a 3 x 10 cm opening was cut into the lid. The reservoir was filled with distilled water and a chamois sponge (Slurpex, Reedman Agencies, North Sydney, AU) was placed through the opening. A removable 18 cm diameter sieve (2 mm aperture metal mesh) with a 5 cm diameter opening was chosen as the optimal entry configuration. We outline the selection of this entry configuration over that used in the PBT [8] and a smaller mesh sieve below.

**Trap entry comparisons**

Three different passive trap entry configurations were compared based on the number of mosquitoes collected per trap night. We used the standard 10 cm hard PVC spigot used in the original PBT, and fine mesh sieves (2 mm aperture metal mesh) of 10 and 18 cm diameters. The mesh sieves were positioned such that the bottom of each faced the inside of the trap and a 5 cm diameter opening was cut in the middle to create an entry point. The field study incorporated a 3 x 3 Latin square experimental design and was conducted in a mixed *Melaleuca* and mangrove swamp adjacent to the Smithfield Waste Disposal Facility near Cairns, Australia (−16.826613°, 145.707065°). All traps were placed approximately 50 m from each other at three different sampling points. The traps were operated for 12 hr each night from 18:00–06:00 with CO₂ gas supplied at a rate of 500 ml/min using a customized gas regulator (Cortis, unpublished) and timer (Pope #1010371, Toro Australia Pty Ltd) setup (Fig. 1b). All traps were rotated to the next position after each collection to reduce sampling point specific differences. Collected mosquitoes were killed in a freezer and morphologically identified in the laboratory [13]. Two morphologically similar species, *Culex annulirostris* (Skuse) and *Culex sitiens* (Wiedemann), were grouped together as the *Cx. sitiens* subgroup because morphological differentiation is difficult in cases where key diagnostic features are damaged [14]. Differences in the mean number of mosquitoes collected were compared by analysis of variance (ANOVA) on log (n + 1) transformed abundance data. The effect of entry type and trap location (i.e. Latin square number) on recorded community composition (relative abundance of each species) was analysed by permutational multivariate analysis of variance (PerMANOVA) models of Bray-Curtis dissimilarities [15].

**Comparison to standard mosquito light traps**

Following the determination of the entry to be used in the SMACK we conducted Latin square trials to assess the efficacy of the SMACK as a mosquito surveillance device compared to a CDC model 512 miniature light trap (CDC, John W Hock; http://johnwhock.com), CDC model 912 Miniature Downdraft Blacklight (CDC + UV, John W. Hock), and an Encephalitis Vector Survey (EVS) light trap (Australian Entomological Supplies; http://www.entosupplies.com.au). Two full 4x4 Latin square trials were conducted, one in which CO₂ was only supplied using 1 kg dry ice and one in which CO₂ was only supplied through a 10 kg compressed gas cylinder at 500 ml/min (Fig. 1b). When baiting passive traps with dry ice, it is critical that the CO₂ gas hose is attached to the top of the insulated cooler (Fig. 1b). Our experience indicates that hoses attached to the bottom of the cooler can become clogged from ice forming from water that condenses within the hose/cooler junction. In addition to recording the total number of mosquitoes collected by each trap type, we also recorded the total number of non-target insects, or by-catch, collected for each trap type. Mosquitoes were identified to species, whereas non-target insects were identified to order. The nightly trapping regime, location of the study, and statistical analyses were the same as those outlined in the entry type comparisons above.

**Mosquito survival study**

To assess the benefits of the addition of the water reservoir and chamois in the SMACK on mosquito survival (longevity) we analysed daily survivorship between SMACKs containing water reservoirs and unmodified PBTs that do not contain such reservoirs. Two traps of each type were stocked with ca. 200 field collected mosquitoes, primarily a mixture of *Aedes vigilax* (Skuse), *Cx. annulirostris*, and *Verrallina funerea* (Theobald). The mosquitoes were collected the previous evening and were returned to the laboratory where they were then distributed amongst the traps. The traps were hung outside the Mosquito Research Facility on the James Cook University (Cairns) campus beneath a 99% shade cloth. Each trap contained temperature and humidity loggers (iButtonLink LLC, Whitewater, WI, USA) and were suspended a distance of 1 m from the ground. An additional pair of data loggers was placed underneath a table adjacent to the hanging traps to monitor ambient air temperature and humidity. Temperature and humidity recordings were taken every 15 min for the length of the study. The number of dead mosquitoes was recorded each day for a period of 14 d. Differences in daily survivorship, as determined by Kaplan-Meier survivorship curves, between the SMACKs and PBTs were determined by the log-rank test [16].

**Mosquito sugar-feeding rate**

We used food colouring added to the honey [6] prior to application on the FTA® cards to quantify the sugar-
feeding rate of field-collected mosquitoes housed in SMACKs for a period of 3 d. Two SMACKs were stocked with ca. 500 female mosquitoes, primarily a mixture of *Ae. vigilax*, *Cx. annulirostris*, and *Ve. funerea*, which were collected the previous evening in the field. Each morning for a period of 3 days different coloured (day 1, blue; day 2, yellow; day 3, red) honey-soaked FTA® cards were introduced into each trap. The following morning a sub-sample of 100 mosquitoes was aspirated from each trap and the sugar-feeding status of each female was determined by observing the abdomen through a stereo microscope. If the female had sugar-fed the colour of the dye on the FTA® card was easily observed within the abdomen (Fig. 2a). After determining the feeding status of each female they were removed from the study. For each consecutive day, the number of secondary sugar-meals was determined based on the mixing of the two different coloured dyes (e.g. blue and yellow = green). On the third day, the number of third sugar-feedings was determined by the mixing of the three different dyes (i.e. purple/brown).

Virus detection in SMACK vs PBT

The ability of the SMACK to detect endemic arboviruses in a field setting was assessed against an unmodified PBT during a pilot trial in May, 2014. The study was conducted in the mixed *Melaleuca* and mangrove swamp in which the trap comparison studies were performed and in a tropical rainforest site (−16.818193°, 145.680896°) adjacent to the James Cook University, Cairns, Australia, campus. One of each trap type was set at each location and the traps were set a distance of 50 m apart within each site. Since this study began prior to the optimization of the chamois water reservoir, the pilot SMACK was set with two 8 x 13 cm thin sponge panels that were moistened with water then fixed to the trap walls with a screw. Traps were operated on a weekly collection schedule during which CO₂ gas was released nightly from 18:00–06:00 at 500 ml/min. Each trap was set with two honey-soaked FTA® cards. At the end of each weekly collection period the FTA® cards were collected and replaced, the sponges rewetted, and captured mosquitoes removed. Captured mosquitoes were returned to the laboratory and morphologically identified [13]. Collected FTA® cards were individually wrapped in Glad Snap Lock Mini® bags (Clorox Australia Pty Ltd, Padstow, NSW), labeled and transported to the Centre for Infectious Diseases and Microbiology Laboratory Services (CIDMLS), Institute for Clinical Pathology and Medical Research (ICPMR), Westmead Hospital, where they were stored at −80 °C until processed for virus detection. The cards were screened for the presence of Ross River virus (RRV) and Barmah Forest virus (BFV) RNA, the two most common and widespread arboviruses in Australia [17], as well as WNV_KUN and MVE by a nested real-time RT-PCR using EvaGreen (Biotium, Hayward, CA) following published protocols [12]. Positive RRV and BFV samples were confirmed by sequencing the product of a section of non-structural protein P4 (174 bp and 219 bp, respectively for RRV and BFV) and comparing generated sequences to GenBank accession numbers [GQ433354.1; RRV] and [AF339488.1; BFV].

Sentinel arbovirus surveillance study

A longitudinal study was conducted in the remote Northern Peninsula Area (NPA) of northern Australia to assess the ability of the SMACK to serve as a sentinel mosquito arbovirus detection system under natural field...
conditions. The NPA area was chosen based on its remoteness and historic use of sentinel pigs to monitor Japanese encephalitis virus (JEV) activity [18, 19]. The field sites included the NPA waste disposal site located in Bamaga (−10.893184°, 142.395665°), a cattle yard in Seisia (−10.852551°, 142.370513°), and a piggery 3 km from Injinoo (−10.891992°, 142.353362°). A single SMACK was set at each site during the first week of February 2015, and operated on a fortnightly collection schedule until May 11th 2015. Each trap contained two honey-soaked FTA® cards and CO\(_{2}\) gas was supplied through a 10 kg gas cylinder and regulated to operate for 12 hr each night (18:00–06:00) at a rate of 500 ml/min. At the end of each fortnightly collection period, the traps were reset by discarding dead mosquitoes, refilling the water reservoir, and replacing the honey-soaked FTA® cards with new cards. Collected FTA® cards were wrapped in Parafilm M® (Bemis NA, Neenah, WI), labelled, and posted to QHFSS laboratories for analysis. FTA® cards were processed for MVEV, WNV\(_{KUN}\) and JEV virus detection following established protocols [6, 9].

Results

Trap entry comparisons

Significant differences \((F_{2,6} = 11.4, P = 0.02)\) were observed among the entry types in the number of mosquitoes collected per trap night (Fig. 3a). Overall, the 18 cm diameter (mean ± SE mosquitoes captured = 1781 ± 165) mesh sieve outperformed the PVC-Spigot (980 ± 121) of the PBT and the smaller 10 cm sieve (1307 ± 23). No significant difference \((F_{2,4} = 4.5, P = 0.91)\) in the composition of mosquito collections (i.e. the proportion of each collection total belonging to a particular genus) was observed among the entry types (Fig. 3b). Based on these results, the 18 cm diameter sieve was chosen as the entry configuration for the SMACK.

Comparison to standard mosquito light traps

The SMACK collected comparable numbers as both CDC trap designs and the EVS trap when CO\(_{2}\) was supplied either with dry ice or via compressed gas (Fig. 4a, Table 1). Species richness (ca. mean of 8 species/trap for each trial) and community composition (Fig. 4b) was comparable among all trap types when CO\(_{2}\) was supplied from dry ice or via compressed gas. The light traps, especially the CDC + UV captured significantly \((P < 0.001)\) more non-target insects (Fig. 4c) than the SMACK and EVS traps. Throughout the 8 trappings, the SMACK collected no non-target insects. The majority of non-target insects belonged to the orders Lepidoptera, Diptera, Coleoptera, and Hymenoptera (Fig. 4d).

Mosquito survival study

The addition of the chamois sponge and water reservoir resulted in significant increases in daily relative humidity \((t_{12} = 2.68, P = 0.01)\), with a mean (± SE) daily increase of 7.0 ± 1.3 % compared to the control PBT, which did not contain the water reservoir (Table 2). No difference in the relative humidity recorded inside the PBT and ambient relative humidity was observed. No significant difference in daily high temperatures recorded within the traps was observed, as well as when compared against the ambient daily high temperature recorded each day. The increased humidity in the SMACK corresponded to a significant \((X^2 = 28.1, P < 0.001)\) increase in the daily survival probability of captured mosquitoes (Fig. 5). Overall, 93.3 ± 1.9 % of mosquitoes in control traps were
dead after 2 days, whereas in traps with sponge, 47.0 ± 7.6% and 33.8 ± 6.5% were alive after 7 and 14 days, respectively.

Mosquito sugar-feeding rate
The sugar-feeding study revealed that a mean (± SE) of 80.0 ± 6.1% of field-collected mosquitoes were sugar-fed on any given day (Fig. 2b) and that 29.2 ± 11.5% had sugar-fed at least twice between days 2 and 3. Further, 8.5 ± 2.9% of field-collected mosquitoes had sugar-fed a minimum of three times by the end of day 3. No significant difference ($F_{1,3} = 1.3, P = 0.89$) in sugar-feeding rates among the species used in the study was observed.

Virus detection in SMACK vs. PBT
The pilot SMACK captured significantly more mosquitoes than the PBT. A mean (± SD) of 1298 ± 1381 and 555 ± 892 mosquitoes were captured in seven weekly paired trappings of the SMACK and the PBT, respectively ($t_{6} = 2.02, P = 0.046$). A total of 12 FTA* cards each for the SMACK and PBT were tested for the presence of BFV and RRV. Two FTA* card pools (2 cards/pool) collected from SMACKs were positive, both of which originated from the tropical rainforest habitat. One pool tested positive for RRV and the other pool tested positive for BFV and RRV. In contrast, no FTA* card pools were positive from the unmodified PBT.

Sentinel arbovirus surveillance study
A total of 48 individual FTA* cards, 16 from each trap site, were tested for the presence of MVEV, JEV and WNV$_{KUN}$ (Table 3). Overall, 13 FTA* cards were positive, comprising 13 cards positive for WNV$_{KUN}$ and 2 cards positive for both WNV$_{KUN}$ and MVEV. Arbovirus detections were greatest during the month of March (7 positive cards) and at the NPA waste disposal site (5 positive cards resulting in 7 virus detections). WNV$_{KUN}$ was the most widely distributed arbovirus (detected multiple times at all three collection sites), whereas MVEV had a limited distribution (detected only at the NPA waste disposal site) and was only detected during the month of March.

Discussion
Effective disease surveillance forms a vital component of any program aimed at reducing the impact of arboviruses on human and animal health. However, the logistics of monitoring arbovirus activity in remote locations using
Table 1 Mean (±SE) species abundance for each trap type when CO\textsubscript{2} was supplied by a gas cylinder at 500 ml/min or with 1 kg of dry ice pellets

| Species                        | Sentinel Arbovirus Capture Kit (SMACK) | CDC model 512 miniature light trap (incandescent) | CDC model 912 miniature light trap (ultraviolet) | Encephalitis Vector Survey trap |
|--------------------------------|----------------------------------------|---------------------------------------------------|--------------------------------------------------|--------------------------------|
|                               | Cylinder Dry Ice | Cylinder Dry Ice | Cylinder Dry Ice | Cylinder Dry Ice | Cylinder Dry Ice |
| Aedes albosculetellus          | 2 (3) 0          | 0                  | 0                  | 0                  | 0                  |
| Aedes alternans                | 1 (1) 10 (10)    | 11 (7)             | 0                  | 4 (4)              | 31 (19)            | 3 (2) 7 (8)        |
| Aedes kochi                    | 181 (131) 74 (49) | 62 (41)            | 61 (27)            | 57 (23)            | 106 (59)           | 92 (37) 63 (46)   |
| Aedes notoscriptus             | 20 (17) 13 (8)   | 15 (9)             | 8 (5)              | 11 (4)             | 0                  | 13 (2) 2 (3)      |
| Aedes palmarum                | 0 9 (6)          | 0                  | 0                  | 0                  | 0                  | 0                  |
| Aedes tremulus                 | 0 0              | 4 (5)              | 0                  | 0                  | 0                  | 0                  |
| Aedes vigilax                  | 198 (43) 1597 (362) | 82 (31)          | 1941 (295)         | 336 (105)          | 3198 (906)         | 53 (16) 998 (431) |
| Anopheles bancroftii           | 0 0              | 2 (2)              | 0                  | 0                  | 9 (9)              | 0                  3 (3) |
| Anopheles farauti sensu lato   | 97 (60) 131 (11) | 58 (24)            | 93 (29)            | 84 (41)            | 158 (46)           | 9 (3) 27 (14)     |
| Coquillettidia crassipes       | 0 0              | 0                  | 0                  | 4 (4)              | 0                  | 0                  |
| Coquillettidia xanthogaster    | 0 0              | 0                  | 0                  | 0                  | 0                  | 1 (2)              |
| Culex cubiul                   | 0 0              | 0                  | 0                  | 6 (6)              | 0                  | 12 (5)             |
| Culex gelidus                  | 1 (1) 0          | 0                  | 0                  | 0                  | 27 (0)             | 0                  |
| Culex hilli                    | 0 50 (8)         | 11 (5)             | 31 (14)            | 0                  | 37 (21)            | 2 (1) 26 (9)      |
| Culex pulius                   | 3 (2) 1          | 0                  | 1 (2)              | 0                  | 16 (4)             | 4 (5) 1 (5)       |
| Culex sitiens subgroup          | 1229 (524) 686 (146) | 1176 (513)        | 639 (123)          | 1372 (514)         | 862 (239)          | 425 (193) 221 (102) |
| Mansonia septempunctata        | 0 8 (5)          | 0                  | 0                  | 0                  | 1 (2)              | 0                  |
| Mansonia uniformis             | 0 13 (8)         | 6 (7)              | 2 (2)              | 0                  | 7 (5)              | 0                  2 (3) |
| Tripteroides magnesianus       | 9 (9) 0          | 0                  | 0                  | 0                  | 0                  | 0                  |
| Uranotaenia sp.                | 2 (3) 0          | 0                  | 4 (5)              | 6 (4)              | 7 (7)              | 0                  |
| Verrallina carmenti            | 99 (44) 42 (26)  | 85 (75)            | 56 (33)            | 134 (58)           | 41 (30)            | 34 (15) 8 (7)     |
| Verrallina funerea             | 316 (99) 125 (28) | 192 (39)          | 158 (58)           | 407 (115)          | 78 (38)            | 184 (52) 74 (27)  |
| Verrallina lineata             | 0 6 (6)          | 1 (2)              | 0                  | 0                  | 0                  | 0                  |

Table 2 Summary of daily relative humidity and temperature recordings observed during the mosquito survivorship (longevity) study.

| Relative Humidity (%) | Ambient Outside | Inside PBT | Inside SMACK |
|-----------------------|-----------------|------------|-------------|
| Mean (±SE)            | 81.0 (2.4)      | 80.3 (2.5) | 88.0 (1.4)  |
| Record Low            | 53.3            | 53.6       | 68.9        |
| Record High           | 99.6            | 99.2       | 99.1        |

| Temperature (°C)      | Ambient Outside | Inside PBT | Inside SMACK |
|-----------------------|-----------------|------------|-------------|
| Mean (±SE)            | 27.3 (0.41)     | 27.9 (0.45) | 27.7 (0.44) |
| Record Low            | 24.2            | 24.3       | 24.2        |
| Record High           | 31.5            | 32.4       | 32.1        |

Summary of daily relative humidity and temperature recordings observed during the mosquito survivorship (longevity) study. The SMACK contained a water reservoir from which moisture was released from a chamois sponge (Slurpex, Reedman Agencies, North Sydney, AU). PBTs did not contain a water reservoir system. Temperature and humidity recordings were taken every 15 min for the length of the study (14 d).
standard battery-powered mosquito traps is often problematic, while cross-reactions in serological assays can greatly reduce the specificity of sentinel animal serology surveys [4]. In the current study, we highlight the development of a sentinel mosquito arbovirus capture kit (SMACK) that does not require battery-power, can compete with standard mosquito traps, maximizes mosquito longevity after collection, and is successful at monitoring arbovirus activity in remote locations.

The enhanced survivorship of field-collected mosquitoes housed in SMACKs compared to those housed in PBTs allow for a maximum number of individual feeding events to occur on FTA® cards. Our results reveal that over a period of 3 days >80% of field-collected mosquitoes will acquire at least a single sugar-meal, and approximately 20–25% will have sugar-fed at least twice over the same time period. These results suggest that although mosquitoes do crowd onto honey-baited FTA® cards, crowding does not inhibit sugar feeding. Consequently, previous protocols advising the use of insecticides (added to the honey) to reduce crowding on honey-baited FTA® cards [8] should be amended to maximize the number of feeding events occurring, and subsequently the amount of virus being expectorated upon the cards.

The detection of four different arboviruses, including the two most important arboviruses in Australia, RRV and MVEV, as well as the detection of multiple viruses on a single FTA® card (WNV_{KUN}, MVEV), demonstrates that the SMACK is capable of adequately sampling the local vector population to monitor arboviruses vectored by different mosquito species with varying ecologies. While the Cx. sitiens subgroup will include both Cx. sitiens and Cx. annulirostris, which we did not separate in the current study, earlier studies in which large numbers of Cx. annulirostris were collected at the same sites [8] and other areas where Cx. sitiens does not occur [9] indicate that passive box traps are effective at capturing Cx. annulirostris [8]. Furthermore, although each detected arbovirus is primarily vectored by Cx. annulirostris, arguably the most medically important mosquito in Australia [20, 21], other vectors of RRV and BFV, such as Ae. vigilax [22, 23], accounted for a large proportion of captured mosquitoes. The detection of WNV_{KUN} in 13/48 FTA® cards from SMACKs set at three sites suggests that the method is sensitive for flavivirus detection in remote areas. Further, the detection of MVEV on two FTA® cards from the SMACK set at Bamaga represents the fourth detection of MVEV using passive traps fitted with FTA® cards [24]. These results, in combination with previous studies which suggest sugar-based systems were more sensitive at detecting arbovirus activity than currently monitored sentinel animals [9, 25], highlight the potential of sugar-based systems to complement and possibly serve as a substitute for sentinel animal programs. However, such comparisons are still limited and full parallel trials comparing the sensitivity, utility and cost of the SMACK, as well as other sugar-based arbovirus surveillance systems, to sentinel animals are needed before any operational changes are made.

In addition to its utility as a long-term arbovirus surveillance device, the SMACK was equally effective as the CDC miniature light traps (average capture ratio 0.86: 1), and more effective than the EVS trap (capture ratio 2.28: 1), at monitoring local mosquito populations when operated on a nightly collection schedule. Further, no
difference in the composition of mosquito collections was observed between the SMACK and light traps indicating that there was little bias in terms of the species collected. Additionally, in contrast to the CDC light traps, the SMACK did not collect any non-target insects during field testing resulting in decreased processing times relative to the CDC light traps. These results, combined with its lack of reliance on battery or mains power (household/city power), make the SMACK a potential inexpensive substitute for traditional mosquito traps that are deployed for a single night. We note that variations in environmental conditions and differences in the responses of individual mosquito species can dramatically influence individual trap efficacy [26, 27]. This was evident in field comparisons of the PBT and EVS trap undertaken in different regions of Australia in which the EVS trap substantially outperformed the PBT [11]. Accordingly, potential SMACK users should initially run this trap in parallel with existing systems to check for relative sensitivities at collecting mosquitoes and costs of operation before changes to existing operational protocols are made. Potential users should consider other practicalities such as cost per unit and size when considering using the SMACK as an overnight mosquito trap. For instance, the SMACK will be available commercially for ca. 80.00 USD (bioquip.com), which makes it the cheapest of the traps tested (CDC, $106.00 USD, johnwhock.com; CDC+ UV, $169.00 USD, johnwhock.com; EVS, $96.95 USD, bioquip.com), while the larger size of the SMACK may be cumbersome during large scale overnight surveillance operations when laboratory and transport space is limited.

**Conclusions**

These results demonstrate that the SMACK has the potential to be a versatile, simple, and highly sensitive arbovirus surveillance tool that may also be used as a traditional overnight mosquito trap. The versatility of the SMACK enables it to be used to complement existing sentinel animal programs and, importantly, serve as a viable substitute when the use of sentinel animals is not feasible. There is also the potential to detect additional vector-borne pathogens transmitted by other hematophagous arthropods using the SMACK. For instance, sugar-baited FTA™ cards have been used to detect Schmallenberg virus in the expectorate of *Culicoides* biting midges [28], which are traditionally surveyed using CO₂-baited traps [29, 30]. Finally, the efficacy and simplicity of the SMACK make it suitable for use in developing countries in which the need for cheap, simple, and efficient arbovirus surveillance tools is often greatest.

**Competing interests**

The authors declare that they have no competing interests.

**Authors’ contributions**

BJJ: participated in trap design and field testing, performed the statistical analysis and drafted the manuscript. TK: conducted arbovirus surveillance field work in the Northern Peninsula Area, Queensland. SHM, AVDH, JLM, and CT: conducted the laboratory arbovirus detection assays and helped draft the manuscript. GC: contributed to trap design, field testing, and developed CO₂ regulator system. KF: trap design and manufacturing. MT: trap design, field testing and participated in the arbovirus surveillance study in Cairns. SD: participated in the arbovirus surveillance study in Cairns and helped draft the manuscript. SAR: conceived of the trap design and the study, participated in field testing, conducted the arbovirus surveillance study in Cairns and helped draft the manuscript. All authors read and approved the final manuscript.

**Acknowledgements**

We would like to thank the Molecular Diagnostics team at Queensland Forensic and Scientific Services. We would also like to thank Chris Paton for his assistance with the trap and entry type comparison studies.

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**Received**: 10 August 2015 **Accepted**: 28 September 2015

**Published online**: 06 October 2015

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