Evidence for regional aeolian transport of freshwater micrometazoans in arid regions

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Scientific Significance Statement
Ephemeral desert water bodies are known for their diverse biota including bacteria, algae, protists, fungi, and micrometazoans. When these water bodies dry, many organisms survive as resting stages (propagules), which may hatch when water returns, or may disperse. However, it is unclear whether propagules of micrometazoans can disperse by wind across regional scales (≥ 100s km) to colonize isolated basins. Our study provides the first evidence that micrometazoan propagules can survive desert dust storms—and that regional-scale windstorms can help them disperse to new habitats. Accordingly, transport by wind likely plays an important role in regional dispersal and potential colonization of aquatic invertebrates in drylands.

Abstract
While separated by large expanses of dry terrain unsuitable for aquatic biota, aridland waters possess high biodiversity. How aquatic micrometazoans disperse to, and colonize, these isolated ephemeral habitats are not well understood. We used a multi-faceted approach including wind tunnel and rehydration experiments, and next-generation sequencing to assess potential movement of diapausing propagules of aquatic invertebrates by anemochory across regional scales (102–105 km). Wind tunnel experiments using dry playa sediments with added micrometazoan propagules demonstrated that after entrainment by saltation and downwind transport, propagules could be subsequently rehydrated and were viable. Further, rehydration of fallen natural dust yielded micrometazoans, including rotifers, gastrotrichs, microcrustaceans, and nematodes. Using conserved

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Author Contribution Statement: JAR collected dust samples from UTEP and Hueco Tanks, analyzed and rehydrated all dust samples, co-designed and implemented the wind tunnel experiments, created HYSPLIT trajectories and GIS image, and helped draft and edit the manuscript. JEM performed sequencing and community analyses including preparation of figures, and helped write those portions of the manuscript. RSVP provided use of wind tunnel and ancillary materials, co-designed wind tunnel experiments, interpreted findings, and edited the manuscript. M-YL provided input into the original experimental design and bioinformatics support, and edited the manuscript. RLW contributed to the conceptual model, aided in the design of the rehydration experiments, helped to draft and edit the manuscript, and developed the artwork. TEG helped conceptualize and design the overall study, developed protocols for and oversaw the dust sampling and particle size analysis protocols, obtained dust samples, helped interpret the findings, and helped draft and edit the manuscript. E JW helped conceptualize and design the overall study including the model, collected dust samples, prepared samples for next-generation sequencing, helped interpret the findings, and helped in drafting and editing the manuscript.

Data Availability Statement: All the data used to analyze the data for this paper are available at the UTEP Bioinformatics Data Repository at http://datarepo.bioinformatics.utep.edu/getdata?acc=ACIEJDV41U1ZNS1.

Additional Supporting Information may be found in the online version of this article.

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Desert aquatic habitats differ from temperate systems in many respects; most obvious is a limited hydroperiod (Williams 2006). Indeed ability to withstand prolonged dryness with large variations in temperature and ultraviolet radiation is critical for the biota of these habitats (Jocque et al. 2010). Yet desert aquatic life is adapted to this duality: flourishing when basins are filled, but withstanding inevitable drought of uncertain duration. Persistence is accomplished through life history adaptations including production of small, drought-resistant stages that quickly re-animate and develop rapidly when suitable conditions return (Brock et al. 2003; Walsh et al. 2014). These stages also can facilitate dispersal (Vanschoenwinkel et al. 2008).

In summarizing the roles of major ecological processes responsible for the distribution of biota, McGill (2010) noted that for dispersal, data at intermediate spatial scales are lacking ($\sim 10^2$–$10^5$ m). This holds true for our understanding of aquatic biota movement among hydrologically disconnected landscapes. Intermediate scales are particularly important in drylands where regional monsoons govern timing and duration of rehydration of temporary waterbodies (Scuderi et al. 2010). These rains fill basins, which upon drying render the land into an expanse of polygonal shaped mud cracks or saline crusts (Fig. 1). Winds blowing across these landforms entrain dust (silt- and clay-sized grains, diameter < 50 $\mu$m) and sand (diameter > 50 $\mu$m) (Field et al. 2010). Researchers have estimated that on a global, annual basis > 1000 Tg of soil is emitted from the ground into the atmosphere as dust, with approximately one-third derived from ephemeral aquatic systems (Ginoux et al. 2012). Along with dust, resting stages of micrometazoans can be transported; however, it is unclear whether this is strictly a local phenomenon or whether such transport occurs across hundreds of kilometers.

The “Everything is everywhere” hypothesis posits that most aquatic microbiota (~ 1–2 mm) are cosmopolitan because they have small propagules that disperse easily by anemochory, hydrochorry, and zoochory (Fontaneto 2011; Viana et al. 2013). Research supports this view for bacteria (Yamaguchi et al. 2012), fungi and protists (Barberan et al. 2015), and phytoplankton (Incagnone et al. 2015), but little is known regarding micrometazoans (Fontaneto 2011). Thus we know a good deal about the dispersal of small aquatic biota at local and global scales, including both temperate and cold habitats (Havel and Shurin 2004; Fontaneto et al. 2006; Nkem et al. 2006). We also recognize the importance of biotic networks (i.e., co-dispersal via zoochory) that enhance dispersal (Tesson et al. 2016). If propagule stages are small enough and sufficiently resistant, they should be able to disperse with sediments during storm events, and survive (Finlay 2002).

In contrast, the extent to which dispersal mechanisms operate on regional scales in dryland systems is relatively unknown. In the U.S. desert southwest hydrochory is restricted to endorheic flows or to rivers that carry biota through aridlands, potentially dispersing it into local floodplain basins during times of exceptional floods (Kobayashi et al. 2015). While the potential for zoochory among aridland basins is clear, it is primarily limited to wet seasons (Sánchez et al. 2012; Viana et al. 2013; Valls et al. 2017). In aridlands, aquatic system isolation and stochastic rain events followed by prolonged dryness and subsequent wind dispersal may be important forces shaping species distribution, extirpation, interrupted gene flow, ecological specialization, speciation, and endemism (Ricklefs 2008; Collins et al. 2014; Hubert et al. 2015).

Previous research has established anemochory of aquatic micrometazoans on local scales (~100 km): i.e., branchiopod (Graham and Wirth 2008) and Artemia cysts (Parekh et al. 2014) were recovered from wind tunnel sediments, and dispersing stages of > 15 invertebrate taxa were captured in an isolated rocky outcrop (Vanschoenwinkel et al. 2009). Further, Vanschoenwinkel et al. (2011) documented zooplankton dispersal ≤ 140 km downwind across those outcrops. Important factors in anemochory include wind speed and direction (Horváth et al. 2016), as well as propagule morphology and sediment grain size (Pinceel et al. 2016). Mesocosm experiments also indirectly support local dispersal (Cáceres and Soluk 2002; Cohen and Shurin 2003). Numerous studies have documented that biological material may be wind-dispersed over distances > ~1000 km: e.g., bacteria and fungi (Barberan et al. 2015), and pollen (Grewling et al. 2016). While these studies have examined components of propagule dispersal, ours is the first to investigate whether aquatic micrometazoans can become entrained in regional-scale, aeolian events, be transported hundreds of kilometers, and remain viable.

As depicted in our conceptual model (Fig. 1), we propose that geophysical processes of aeolian transport in deserts operate to disperse micrometazoans and biological processes determine community structure (Field et al. 2010; Heino et al. 2015). Our model can be described as follows. Regional water sources contain propagules of locally defined, but regionally diverse species assemblages (Sources). This γ-diversity is then filtered by dispersal through wind corridors...
that entrain dormant propagules along with dust (Anemochory). Propagules move varying distances based on size, density, and other properties that affect their aerodynamics, as well as the vagaries of the wind (Jenkins et al. 2007). Thus we should expect to find that distance decay plays a role in dispersal of micrometazoans and that geographic features of the land can influence dispersal rates (Incagnone et al. 2015). Propagules landing in suitable habitats may emerge from dormancy during wet phases (Biotic processes) (Pinceel et al. 2013) and depending on community dynamics may become established (De Meester et al. 2002).

Scattered across the drylands of southwestern U.S.A. and northern Mexico are countless, highly disconnected, ephemeral aquatic habitats that collectively form a hotspot of biodiversity (Olson and Dinerstein 2002). This area also often experiences large, regional-scale dust storms that preferentially

Fig. 1. A model of aeolian dispersal of micrometazoan propagules and subsequent community development in aridland aquatic systems. Dispersal of dust and propagules (Brown dots) begins with physical processes (Brown arrows) that act as filters that sort all particles: (1) Sources (of particulate materials), (2) Anemochory (wind dispersal). For propagules, additional filters (Blue) govern potential for colonization: (3) ability enter into a desiccated state (propagule production), (4) resistance of propagules to stresses associated with drying and prolonged drought, and (5) resilience, the ability of the propagules to hatch depending on edaphic conditions. While life history adaptations of new arrivals or those already present (Dashed lines) may sanction persistence and hatching, continuance is not assured. Community development and propagule replenishment are biological processes. Collectively these processes work in concert shaping the $\alpha$, $\beta$, and $\gamma$-diversity of active aquatic communities. Terms used describing aeolian and biotic processes are given in Supporting Information Document 4.
emanate from these habitats when they are dry (Lee et al. 2009; Baddock et al. 2011). Our research question was: Can micrometazoans be dispersed via wind over regional scales (10^2–10^3 km) and survive transport to colonize distant aquatic habitats? Thus, we examined five elements necessary for aeolian transport. Each of which should contribute to the dispersal of micrometazoans in aridland aquatic ecosystems as described in our conceptual model (Fig. 1). (1) To determine possible sources of dust landing at a Chihuahuan Desert collecting site on the campus of The University of Texas at El Paso (UTEP), we back-calculated trajectories of 13 dust-bearing, wind events (Sources). (2) To assess whether the size of falling sediments overlapped sizes of micrometazoan propagules, we analyzed the particle size distribution of the dust falling from those events (Anemochory). (3) To determine whether these propagules are entrainable during dust storms, we conducted wind tunnel experiments that mimic dust emissions from the sediment-propagule banks of dry playas (Sources; Anemochory). (4) To investigate propagule viability after transport, we conducted rehydration experiments for wind tunnel-entrained dust and fallen dust in the Chihuahuan desert (Hatching). (5) To more fully characterize the taxonomic distribution of propagules from selected windstorms, we conducted next-generation sequencing (NGS) on fallen dust. This method captures taxa that do not respond to hatching cues provided in rehydration experiments.

**Methods**

**Collection and processing of dust**

Falling dust was collected from windstorms (2002–2016) using one of the following types of passive collection traps: marble dust collectors (MDCO), modified Wilson and Cooke (MWAC) samplers, and Big Spring Number Eight (BSNE) samplers (Goossens and Offer 2000). Samples were collected using MDCOs from two sites in El Paso County, Texas: 10 samples from UTEP Biology Building Rooftop (B), and two from Hueco Tanks State Park and Historic Site (HT) (Rivas et al. 2018; Supporting Information Document S1, Supporting Information Table S1). Additional samples from the arid southwest U.S.A. were collected: one from Jornada Basin LTER (LJ), NM, and two from White Sands Missile Range (WSMR), NM (using MWAC) and two from Yellow Lake playa (YL), TX (using BSNE). Samples were collected November through May, when regional-scale dust storms prevail (Novlan et al. 2007). We measured particle size by laser diffraction (Sperazza et al. 2004) using ~ 0.3 g of each sample with a Malvern Mastersizer 2000 (Malvern Instruments, UK). Particle size was also determined for sediments captured in each section of the wind tunnel (0.5 g). The dry method was used to preserve the initial particle size of transported sediment.

**Wind trajectories**

Using the NOAA HYSPLIT model (Stein et al. 2015) we determined potential sources of transported materials using back-calculated flow trajectories of wind events for dustfall samples collected at UTEP. This method determines origins and transport pathways of air masses based on the latitude and longitude coordinates of starting or ending points. All parameters were set to the default settings. These included a total run time of 24 h (encompassing the duration of the dust event), 24 trajectories (generating a mean trajectory), and a height of 500 m above ground level (representing dust arriving at the receptor site in the atmospheric boundary layer, from where it could fall out into the collector).

**Wind tunnel experiments**

To test entrainment of playa invertebrate diapausing stages (five common freshwater species: *Eulimnadia texana*, *Triops longicaudatus*, *Streptochalps* sp., *Daphnia magna*, *Brachionus calyciflorus*; two representative brackish water species: *Artemia salina*, *B. plicatilis*) by saltation (energetic sandblasting: the dominant method of dust emission (Shao 2008)) aeolian transport was simulated in a laboratory, suction-type wind tunnel at the USDA-ARS Big Spring Field Station (Van Pelt et al. 2009). Details of experiments are provided in Supporting Information Document S2, Supporting Information Table S2. Successful transport of propagules in these simulations was demonstrated by counting propagules deposited in each of three downwind sections of the wind tunnel. These are a transfer section (dispersal of a few meters), a settling chamber (10–100s of meters), and the filter section (up to 10^6 m). Subsamples of sediment deposited in each section were rehydrated in an appropriate medium (see Rehydration experiments: Wind tunnel, below) and monitored for hatching. Number of propagules recovered in each section was calculated as sum of hatchlings plus any unhatched propagules remaining after rehydration.

**Rehydration experiments**

Subsamples of dust from (a) dust collectors and (b) wind tunnel experiments were rehydrated as follows.

(a) Collectors: A total of 47 samples were rehydrated from dust collected from UTEP Biology Building Rooftop (B), Hueco Tanks State Park and Historic Site (HT), White Sands Missile Range (WSMR), Yellow Lake (YL), and Jornada Basin LTER (LJ). Subsamples (1–3 g) were rehydrated in 250 mL of sterile MBL medium (Stemberger 1981). They were incubated at 25°C (two subsamples) and 12°C (one subsample) under a 12 : 12 (Light : Dark) photocycle. Additional details are given in the Supporting Information Document S1. They were examined under a dissecting scope for emerging invertebrates daily until no new taxa were found for three successive observations, and finally after one additional month. Micrometazoans were identified, photographed, and preserved as vouchers (Supporting Information Document S1, Supporting Information Fig. S1).

(b) Wind tunnel: Sieved wind tunnel sediments were divided into subsamples: three from the transfer section, five from the settling chamber, and one from the filter section.
About 2.0 g of sediment were rehydrated (3–5 subsamples) in 75 mL of an appropriate medium (Supporting Information Document S2). All cultures were maintained at room temperature, under constant illumination, and observed daily for 3 weeks. The filter section represented a single collection point, but the sample was divided into two subsamples for rehydration. As a control, five subsamples of 3.0 g of abrader sand was rehydrated with MBL and incubated under the same conditions.

NGS and community analysis

Total DNA was extracted from dust samples ($n = 17$; $\sim 0.25$ g each) using a PowerSoil kit (MoBio, Carlsbad, California) following the manufacturer's protocol. DNA was submitted to MRDNA labs (Shallowater, Texas) for 18S tag-encoded FLX-Titanium amplicon pyrosequencing using SSU_F04/SSU_R22 primer sets. Replicate samples of two dust events (UTEP [BD14: Biology Building Roof Top, Dec 2014; HT [HTMY14: Hueco Tanks State Park and Historic Site, May 2014]) were sequenced, giving a total of 19.

We analyzed sequencing reads using QIIME (v.1.9.1; Caporaso et al. 2010) and clustered at 97% sequence identity to delineate operational taxonomic units (OTUs). OTUs were then taxonomically assigned using BLAST (Altschul et al. 1990) against the Silva v.128 reference (Yilmaz et al. 2014). Fungal sequences were excluded in downstream analysis. Abundance barcharts based on remaining OTU assignments were constructed to illustrate patterns of diversity among samples. We also conducted Principal Coordinate Analysis (PCoA) to assess similarity of OTUs. (Additional details are given in the Supporting Information Document S3).

Results

Mean particle size of sediment samples for each dust event ranged from: Hueco Tanks (HT), 100–112 $\mu$m; UTEP (B), 81–295 $\mu$m (10–35% of collected material represented dust $<50$ $\mu$m; Fig. 2; Supporting Information Document S1, Supporting Information Table S1); White Sands Missile Range (WSMR), 178–200 $\mu$m; and Yellow Lake (YL), 6–12 $\mu$m. The substantially smaller particle size at Yellow Lake (YL) represented aggregates of clay and fine silt-sized, lacustrine mud particles loosely bound by salts. Note that the size range of micrometazoan propagules falls within the size range of dispersing dust we collected (Fig. 2).

NOAA HYSPLIT back-trajectories for samples collected at UTEP, demonstrated that dust was transported from multiple directions, predominately from the southwest ($n = 7$), with two from the south, one from the east and three from the north (Fig. 3). All of these windstorms crossed extensive regions of the Chihuahuan Desert encompassing ephemeral aquatic systems located in the Sierra Madre Occidental and the Rio Grande Valley.

Wind tunnel experiments

As expected mean particle size decreased in the three downstream sections of the wind tunnel: transfer section, settling chamber, and filter section (462 (SD = ± 6.3%)), 184 (SD = ± 5.5%), and 52 (SD = ± 5.2%) $\mu$m, respectively. Rehydration of sediments from each section indicated that propagules were transported throughout (Supporting Information Document S2, Supporting Information Table S2) and that some individuals of all taxa were viable. For example, in the first experiment > 18,000 fairy shrimp propagules of the original $\sim 300,000$ were recovered in the transfer section, of which $\sim 18%$ of were viable. In the settling chamber almost 19,000 fairy shrimp propagules were recovered, of which $\sim 38%$ were viable. Finally in the filter section > 6000 propagules were recovered and 0.6% were viable. Rehydration of the abrader sand yielded no organisms.

Dustfall rehydration experiments

Rehydrations yielded representatives of several broad taxonomic groups: algae (all, except BD14-2), ciliates (Hueco Tanks State Park and Historic Site (HT), UTEP Biology Building Roof Top (B), White Sands Missile Range (WS), Yellow Lake (YL)), gastrotrichs (UTEP Biology Building Roof Top (B)), nematodes (UTEP Biology Building Roof Top (B), Hueco Tanks State Park and Historic Site (HT)), ostracods (Yellow Lake (YL)), and monogonont rotifers Cephalodella sterea (UTEP Biology Building Roof Top (B)), Proales cf. similis (Yellow Lake (YL)), Ptygura beauchampi (UTEP Biology Building Roof Top (B)) and bdelloid rotifers Adineta vaga, Philodina tranquilla (UTEP Biology Building Roof Top (B) and Hueco Tanks State Park and Historic Site (HT)). Bdelloid rotifers occurred in 21% of rehydrated samples, while monogonont rotifers and nematodes were found
in 6%, and gastrotrichs, ostracods, and fairy shrimp were found in 2%.

**NGS and community analysis**

Eukaryotic-specific primers (SSU) recovered 34,086 reads and 3,327 OTUs after fungi were removed (Supporting Information Document S3). As seen in the barchart, sequencing replicates had similar taxonomic assemblages but at different proportions (Fig. 4A: HTMy13-1, 13-2 [Hueco Tanks May 2013, replicates 1 and 2]; BD14-1, 14-2 [UTEP Biology Building Rooftop, December 2014, replicates 1 and 2]). This is also evident in the Hueco Tanks (HTMY13) sequencing replicates shown in the PCoA plot (Fig. 4B). Typically, and as seen here, abundant taxa mask the diversity of the rare forms. For instance, BMy14b (UTEP Biology Building Rooftop, May 2014) had 54 metazoan OTUs, ~82% of reads were ciliates. The assigned taxonomy identified rotifers, nematodes, ostracods, and gastrotrichs; rehydration experiments yield a similar suite of taxa (Table 1). PCoA plots show differences in taxonomic composition among sampling locations, with BD14-2 (UTEP Biology Building Rooftop, December 2014, replicate 2) having a unique taxonomic composition.
Focusing on the UTEP samples, taxonomic assemblages can be differentiated by wind direction. Species assemblages in dust collected from wind events originating from the southwest can be isolated from those originating from other directions by a plane across the three axes.

Discussion

We confirm that propagules of micrometazoans are entrained during Chihuahuan desert wind events and that anemochory provides a mechanism for their dispersal over regional scales. In addition, some propagules retain viability through the entire dispersal process. Our conclusions are based on a unique approach that weds geological and biological methods. These techniques can be used to further explore community assembly in ephemeral aquatic systems.

The modeled back trajectories show that wind events in the Chihuahuan Desert deposited material in downwind collectors at UTEP. These sediments crossed dust-emitting ephemeral aquatic systems, including playas such as the Paleolake Palomas Basin, Mexico (Baddock et al. 2016) and Lake Lucero (White Sands, New Mexico) (White et al. 2015), centered 79 km and 112 km away from UTEP, respectively (Fig. 3). Other regional dust sources traversed by the incoming winds include ephemeral riverbeds, sand sheets and dunes, alluvial systems, and agricultural lands (Rivera Rivera et al. 2010; Baddock et al. 2011; Horváth et al. 2016). Easterly winds infrequently bring dust to the Chihuahuan Desert from the Great Plains, including particles derived from playas such as Yellow Lake (Sweeney et al. 2016). These trajectories match previously identified dust flow pathways into El Paso (Novlan et al. 2007).

As sediment particles move during wind events, saltation, bombardment, and collisions of large particles loft smaller sediment grains, as well as dormant stages of aquatic invertebrates into the atmosphere (Fig. 1: Physical processes). We know that factors such as propagule size, density, morphology, original habitat, and wind influence transport distance. In our work assessing regional transport, the size range of micrometazoan propagules coincided with that of windfallen dust and sand, both for rotifers (50–200 μm) and crustaceans, such as Notostraca (~ 400 μm), Anostraca (~ 270–380 μm), and Spinicaudata (~ 200 μm) (Thiery and Gasc 1991) (Fig. 2). Moreover, Ricci et al. (2003) asserted that bdelloid rotifers adhere to sand grains as they enter anhydrobiosis and others have shown that these taxa can be transported during local wind events (Havel and Shurin 2004; Vanschoenwinkel et al. 2009).

Wind tunnel experiments have long been used to study many aspects of dust emission and aeolian transport (Anderson et al. 1991), but no attempts to use them for documentation of...
anemochory of invertebrate propagules have been made until recently. Pinceel et al. (2016) demonstrated dispersal using a highly simplified wind tunnel analog, determining the threshold velocity of emission and that wind dispersal is highly correlated to propagule morphology and size. Our wind tunnel experiments more accurately recreated the mechanics of dust emission, including bombardment by saltating sands onto polygon-shaped, soil samples representative of natural playa surfaces. Collection of sediments and propagules from downstream points represented long distance dispersal (Supporting Information Documents S2). However, even if established that micrometazoan propagules are transported with dust, it remained to be shown that they retained viability throughout the highly turbulent, energetic process of saltation (Shao 2008).

Our wind tunnel experiments demonstrate emission of viable propagules (Supporting Information Documents S2). While some propagules were deposited in the transfer section of the wind tunnel, representing dispersal of only a few meters, the greatest numbers of propagules were always present in the settling chamber. Propagules transported to this section suggest dispersal of 10s–100s of m downwind. However, a sizeable number of propagules were present in the filter section, suggesting potential dispersal of up to $\sim 10^6$ m.

The exception was the absence of Daphnia ephippia in the filter section. It is possible that the unique shape of an ephippium does not permit long distance dispersal. Moreover, the dispersing propagules from all seven taxa showed viability after transport (Supporting Information Documents S2). Our wind tunnel results show a decline in number of propagules dispersed in air from the settling chamber to the filter section (farthest downstream). This decline shows that not all propagules disperse equally well.

Resurrection ecology has demonstrated that viable micrometazoan propagules can be recovered from lake sediments. We used this technique to demonstrate the potential viability of propagules found in falling dust. However similar to species recovery from sediments, we posited that micrometazoans recovered from dust are a small subset of the regional pool.

Our results confirm high taxonomic diversity of micrometazoans transported through regional dust events. While not confirming viability, NGS better captures taxonomic diversity of environmental samples (Santoferrara et al. 2016). We found hundreds of micrometazoan OTUs in fallen dust. The five samples with highest diversity (excluding fungi) were collected from the UTEP site (29–54 OTUs; Supporting Information Document S3, Supporting Information Table S3). Of these, several were assigned to taxa that were found in

### Table 1. Metazoan taxa that were found in both the environmental sequencing (Operational Taxonomic Units and [reads]) and during rehydration (R) of dust events. Full results are recorded in the Supporting Information Document S3, Supporting Information Document S4.

| Collector site | Sample | Gastrotricha | Nematoda | Ostracoda | Anostraca | Rotifera |
|---------------|--------|--------------|----------|-----------|-----------|----------|
| Hueco Tanks   | HTMy13-1 | —            | 3 [14]   | —         | —         | R        |
|               | HTMy13-2 | —            | 11 [62]  | —         | —         | R        |
|               | HTF14   | —            | R        | —         | —         | —        |
| UTEP          | BA13    | —            | 1 [6]; R | 3 [13]    | —         | R        |
|               | BMr14   | —            | —        | —         | —         | R        |
|               | BMy14b  | R            | 6 [7]; R | —         | —         | R        |
|               | BD14-1  | —            | 1 [1]    | —         | —         | —        |
|               | BA15    | —            | R        | —         | —         | —        |
|               | BMy15   | —            | 6 [16]   | —         | —         | —        |
|               | BN15    | —            | 3 [3]; R | —         | —         | 7 [34]; R|
|               | BMr16   | —            | 2 [3]    | —         | —         | 3 [5]; R |
| Jornada       | LJMr14  | —            | 5 [49]   | —         | —         | 1 [1]    |
| White Sands   | WSJ10a  | 1 [4]        | 2 [5]    | —         | —         | —        |
|               | WSJ10b  | —            | 4 [6]    | —         | —         | —        |
| Yellow Lake   | YLMr02  | —            | —        | R         | —         | —        |
|               | YLMr03  | —            | 2 [7]    | 20 [1349] | R         | R        |

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rehydration experiments and/or used as propagules in the wind tunnel experiments (Table 1). Because different portions of the samples were used, it is not surprising that there is incomplete correspondence between taxa recovered by the two methods. Dust is emitted from a variety of sources and winds mix it as it crosses different landforms. Moreover, routes and speeds of storms vary, giving each event a unique signature of entrained materials.

Although we have filled in some gaps in our understanding of regional dispersal of aquatic micrometazoans by wind, there are several remaining knowledge gaps. Chief among these are the following. (1) Regional differences and interannual variation in aeolian transport of aquatic propagules. (2) Detailed examination of emission and dispersal dynamics, and survival of dispersing propagules. (3) The role of propagule aerodynamics and morphology in emission and transport by wind. (4) Traits that determine survival during deflation, transport, and colonization. (5) How soils and landform characteristics influence deflation of propagules during windstorms. While much remains to be known, here we demonstrated that micrometazoan propagules remain viable after entrainment and dispersal over long distances. Thus anemochory performs an important role in regional dispersal and potential colonization of some micrometazoans in dryland aquatic habitats.

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