Title: The diversity of arthropods in homes across the United States as determined by environmental DNA analyses.

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Abstract
We spend most of our lives inside homes, surrounded by arthropods that impact our property as pests and our health as disease vectors and producers of sensitizing allergens. Despite their relevance to human health and well-being, we know relatively little about the arthropods that exist in our homes and the factors structuring their diversity. Since previous work has been limited in scale by the costs and time associated with collecting arthropods and the subsequent morphological identification we used a DNA-based method for investigating the arthropod diversity in homes via high-throughput marker gene sequencing of home dust. Settled dust samples were collected by citizen scientists from both inside and outside more than 700 homes across the United States, yielding the first continental-scale estimates of arthropod diversity associated with our residences. We were able to document food webs and previously unknown geographic distributions of diverse arthropods—from allergen producers to invasive species and nuisance pests. Home characteristics, including the presence of basements, home occupants, and surrounding land-use, were more useful than climate parameters at predicting arthropod diversity in homes. These non-invasive, scalable tools and resultant findings not only provide the first continental-scale maps of household arthropod diversity, our analyses also provide valuable baseline information on arthropod allergen exposures and the distributions of invasive pests inside homes.

Introduction
As soon as humans started building homes, they began to provide, inadvertently, shelter to other animals, including scavengers, but also parasites and associated pathogens. Arthropods likely represent the majority of these animal species (Bertone et al. 2016; Leong et al. 2016). The most conspicuous arthropods in houses tend to be those that are unwanted, either because they damage our homes (e.g., termites), and products (e.g. pantry pests), are viewed as unsightly (e.g., house centipedes), irritate our pets (e.g. fleas), or negatively affect our health. Some arthropods disperse pathogens and antibiotic resistant microorganisms through homes and similar built environments as they move from feces, sewage, carrion, and waste to our food, beds, and bodies (Graczyk et al. 2001; Olsen et al. 2001; Ahmad et al. 2011; Zurek & Ghosh 2014). Other arthropods, including dust mites and cockroaches, affect us more indirectly, by

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shredding sensitizing allergens that can lead to allergic rhinitis, allergic asthma, and atopic eczema (Arlian 2002; Calderón et al. 2015), with the frequency and severity of these disorders having increased in the United States in recent decades (Bousquet et al. 2001; Moorman et al. 2007). Our general understanding of the types of arthropods we are exposed to inside our homes, and their geographic distributions, remain limited despite the potential impacts of these arthropods on our health and the overall quality of our living spaces.

Most studies investigating indoor arthropods have focused exclusively on a select few pest species that are most conspicuous, such as cockroaches and ants (Smith 1965; Schal & Hamilton 1990), or those species that are most damaging to building materials, such as termites (Lax & Osbrink 2003). Yet, recent in-depth studies of arthropods in houses in a single city, Raleigh, North Carolina suggests that these relatively well-studied species represent a miniscule fraction of household arthropod diversity (Bertone et al. 2016; Leong et al. 2016). Bertone et al. (2016) found species from an average of 62 arthropod families per home, with many hundreds of morphospecies identified across the fifty homes surveyed. How these arthropods might be distributed within or among cities and geographic regions remains largely undetermined. Since approximately 56% of United States’ households use insecticides (Grube et al. 2011), arthropods are almost certainly common in many homes, but the full extent of this arthropod diversity across the United States remains unknown. One reason these knowledge gaps persist is that surveying arthropod diversity using standard methods is difficult. Collecting and accurately identifying the broad range of arthropod taxa that can be found in homes is costly, labor-intensive, highly invasive, and logistically challenging. For example, just to survey arthropod diversity in 50 homes in North Carolina required six person hours per home to first collect samples, and then at least two hours per home by trained entomologists to morphologically identify the arthropods via microscopy. Even with such exhaustive techniques, some species remain impossible to identify because of their life stage, sex, or because of sample degradation.

Recently, DNA sequencing-based methods have been developed with the potential to circumvent some of the limitations of traditional morphology-based identifications and, in doing so, accelerate the process of surveying arthropod diversity from environmental samples. Typically these DNA-based methods rely on PCR amplification and high-throughput sequencing of a fragment of the cytochrome c oxidase subunit 1 gene (Cox1/Co1/COI) using arthropod-specific primers that should amplify this gene region from nearly all arthropods (Hebert et al.)
The targeted gene region is advantageous as it is sufficiently variable to permit identification of most arthropods down to the taxonomic level of genus or species (Meusnier et al. 2008). Moreover, the targeted gene region is sufficiently short to permit analyses of arthropod diversity from environmental samples even if the arthropod DNA has been degraded. Such techniques have been used previously to quantify arthropod diversity in taxonomically unresolved communities (Burgar et al. 2014), animal diets (Bohmann et al. 2011), artificially mixed communities (Yu et al. 2012), and tree canopies (Gibson et al. 2014), but, to our knowledge, such molecular techniques have not previously been used to survey arthropod diversity inside homes.

We used such a DNA-based approach to rapidly assess arthropod diversity in settled home dust collected from inside and outside ~730 homes across the continental United States. Our goals were to characterize the diversity of arthropods found inside homes, determine which arthropods are repeatedly found inside homes versus those which are transient, outdoor taxa, and identify those factors that predict the continental-scale distributions of individual taxa (such as dust mites). Specifically, we investigated some of the building, environmental, and occupant factors that predict what homes harbor a greater diversity of arthropods. We also assembled some of the first maps of arthropods detected in homes across the United States to describe how the distributions of specific taxa of medical, economic, or ecological interest vary across regions.

**Materials and Methods**

**Sampling**

Dust samples were collected as part of the Wild Life of Our Homes citizen science project, using methods described previously (Barberán et al. 2015b; a). Briefly, citizen scientists used sterile, dry swabs to sample the small ledge of the top of the door trim from the main living area inside homes and the door trim on the outside of the main entrance to the home. Swabs were then shipped to Boulder, CO and stored at -20 °C until further processed. These samples were collected as part of a previous investigation on the biogeography of bacteria and fungi in the United States, where participants completed an online questionnaire about home occupants, design, and habits wherein. All participants were provided a written informed consent form approved by NC State University's Human Subjects Research Committee (Approval No. 2177), as well as a sampling kit and instructions for sampling their home. Our study included 1462 of
these settled dust samples collected from every state in the continental United States, with 732 collected from inside homes and 730 from outside homes (SI Appendix, Fig. S1). We identified characteristics of the sample geographic location and associated household information using occupant surveys and location information as described in more detail previously (Barberán et al. 2015b).

**Molecular Analyses**

DNA was extracted from swabs using MoBio Powersoil-htp kits, using the modified protocol described previously (Fierer et al. 2008). We amplified a ~157 bp region of the mitochondrial cytochrome c oxidase subunit I gene using arthropod-specific primers (Zeale et al. 2011; as used for high-throughput sequencing such as Bohmann et al. 2011) (SI Appendix, Table S1). Primers were modified to permit multiplex sequencing on the Illumina MiSeq platform as both the forward and reverse primers included the appropriate Illumina adapters and the reverse primers included a unique error-correcting 12-basepair barcode for each dust swab sample per sequencing run (Hamady et al. 2008). The 25 µL PCR cocktail included 3 µL gDNA, 12.5 µL Master Mix (Promega Biotech Co.), and 1 µM each of the forward and reverse primers. Cycling parameters were modified from Pinot et al. (2014) (94 °C for five min, followed by 45 cycles of 94 °C 30 s, 45 °C 45 s, 72 °C 45 s and a final extension at 72 °C for 10 min). Duplicate PCRs for each sample and the associated controls were pooled, cleaned, and normalized using SequaPrep Normalization Plates (Thermo Fisher Scientific). Samples were pooled and sequenced on three Illumina MiSeq runs using 2x250 chemistry at the University of Colorado BioFrontiers Institute Next-Gen Sequencing Core Facility. In order to increase sequencing run quality with such an AT-rich amplicon, we increased the standard amount of PhiX used during the sequencing runs from 15% to 30% as suggested by the manufacturer.

**Bioinformatic Analyses**

We generated approximately 43 millions sequences. Sequences were demultiplexed using a custom python script 'prep_fastq_for_uparse.py' (https://github.com/leffj/helper-code-for-uparse) and the reverse reads were used for downstream analyses due to higher quality scores. Sequences were filtered and OTU (operational taxonomic unit) picking was performed using the UPARSE pipeline (USEARCH v.7) (Edgar 2013). Briefly, sequences were initially quality filtered in UPARSE (Edgar 2013), sequences were trimmed to the expected amplicon length (158 bp) with those sequences removed that were of insufficient quality (maxee value <1.5), below the
minimum expected amplicon length (90 bp), or singletons (i.e. unique sequences that appeared only once across the entire dataset). Sequences were clustered de novo at 99% similarity. Taxonomy assignments were performed in QIIME (Caporaso et al. 2010), using the hierarchical naïve bayesian classifier from the Ribosomal Database Project at 99% similarity and with a confidence of 0.5 (Wang et al. 2007), re-trained with a custom reference database curated from the Bar Code of Life Database (v3) (Ratnasingham & Hebert 2007). Due to the variable homology of the Co1 gene across taxa, we agree with others (Porter et al. 2014) that it is critical to use such a confidence-based hierarchical taxonomy classifier when assigning identity to these sequences. As expected, the differential variability in the sequenced Co1 gene region, as well as potential sequencing errors, led to multiple OTUs per genus for certain taxa, so all unique OTUs were collapsed into their respective genera for downstream analyses (see SI Appendix, Fig. S2 for more information on the sequence processing pipeline).

We validated this molecular and bioinformatics pipeline by processing and sequencing the targeted Co1 gene region from a collection of 58 arthropod taxa that were morphologically identified by a trained entomologist (SI Appendix, Table S2). Based on this validation, identified genera in our sample set with singleton sequences were further removed and sequences were transformed into presence-only data. Three taxa were identified in a control sample of our sequencing run, and were therefore conservatively removed from the analysis (These taxa made up less than 0.4% of that identified taxa). Sequences identified as originating from marine and aquatic arthropods (e.g., crabs, shrimp, and crayfish) were removed from downstream analyses (except for mapping in SI Appendix, Fig. S3), as they likely originated from pet or human food. Given that arthropods can vary dramatically in size and DNA can be shed through frass (feces), sequence abundances are not directly proportional to biomass and all data were therefore analyzed using presence/absence metrics alone (Elbrecht & Leese 2015; Barnes & Turner 2016).

Data Analyses

All analyses were carried out in the R environment (R Core Team 2013) (See SI Appendix, Fig. S1 for sample sizes used in different analyses.)

To determine how the results from our DNA-based survey compare to results obtained with surveys conducted using traditional sampling and morphology-based identification methods, we compared our findings with those of a recent study that investigated the arthropods of 50 homes in Raleigh, North Carolina (Bertone et al. 2016). For these analyses, we focused just on
those indoor dust samples collected from 87 North Carolina houses included in this study. The Bertone et al. (2016) survey involved exhaustive sampling of all rooms in each of the 50 houses within close proximity to the city of Raleigh, N.C., where ours involved door trim swabs from a non-overlapping set of 87 homes collected within the same state.

To investigate what factors might be driving differences in diversity, we ran generalized linear models (GLM) with Poisson errors. Variables were selected using forward and backward stepwise model selection by the Akaike information criterion (AIC) as implemented in the R function stepAIC (Venables & Ripley 2002). Three variables (urbanization, presence of basement and presence of pets) explained 9.3% of the variability of indoor arthropod diversity. To determine if the greater diversity trend with pet ownership was consistent across broader taxonomic groups, we compared diversity across the factor of interest within the top five orders in homes (Lepidoptera, Diptera, Hemiptera, Coleoptera, and Araneae) with false discovery rate (fdr) corrections. All variables (outdoor and indoor variables; see ref (Barberán et al. 2015b) for the complete list of variables explored) explained 15.4% of the variability of indoor arthropod diversity.

To determine what arthropods were preferentially found in indoor versus outdoor home samples, we determined what taxa had significant fidelity to each of these locations. This was calculated using Pearson's phi coefficient of association for each genus and each habitat using the multipatt function (De Cáceres et al. 2010) in the R package indicspecies (De Cáceres & Legendre 2009). We adjusted the p values for multiple comparisons using an fdr correction. For taxa significantly correlated with indoor or outdoor environments, we visualized the correlation strengths using a heat map.

For select taxonomic groups of interest, we mapped detection in indoor samples over the continental United States.

**Results and Discussion**

Based on swab collections of settled dust and the arthropod DNA found therein, we noted the presence of more than 600 unique arthropod genera inside the 732 homes included in this study. Across all homes, we detected 28 orders of terrestrial arthropods. The taxa identified range from those that are well-studied inhabitants of the home (e.g., dust mites, cockroaches,
and carpet beetles) to taxa that are not commonly considered to be household inhabitants (e.g., aphids and certain parasitic wasps) (SI Appendix, Table S3). Most of these unexpected taxa, however, appear to be associated with homes as highlighted by the fact that many of these same unexpected taxa were also found in homes where arthropod diversity was surveyed using a more traditional approach (Bertone et al. 2016).

Most arthropod taxa were restricted in their distributions and only detected in a few homes (Fig. 1). Surprisingly, the genera that were most ubiquitous (i.e. detected in at least 10% of those homes with arthropods) included mosquitoes (Aedes spp.) and plant-associated aphids (Aphis spp.), along with those taxa more commonly associated with the home environment such as dust mites (Dermatophagoides spp.), carpet beetles (Anthrenus spp.), and Indian meal moths (Plodia spp.). More broadly, taxa from the orders Lepidoptera (moths and butterflies), Hemiptera (true bugs), and Diptera (true flies) were found in a majority of indoor samples (Fig. 1). These three orders also represent the majority of the arthropod diversity detected across the homes (SI Appendix, Fig. S4). Many of the taxa that were frequently detected inside homes include light-bodied, flying arthropod species (Figs. 1, 2). The abundance and diversity of such taxa may reflect the extent to which insects are attracted to, and subsequently trapped inside, homes, either by home lighting (Somers-Yeates et al. 2013), elevated carbon dioxide inside homes (van Loon et al. 2015), or household organic waste (Karimifar et al. 2011).

Comparisons of DNA-based methods to more traditional survey methods

Despite the known limitations of DNA-based surveys—articulated well elsewhere (e.g. 14, 18)—many of the dominant taxa identified with our DNA-based approach were also identified using more traditional survey methods (SI Appendix, Figs. S5, S6). Our results compared favorably, despite sampling different homes within North Carolina, and using very different sampling approaches. The two methods captured many of the same families and orders of arthropods. Moreover, the most diverse orders identified were similar in the two studies, (SI Appendix, Fig. S7). Though many of the arthropod families identified from morphology-based collections were also found in the molecular survey, there were some exceptions (SI Appendix, Table S4). We did not detect certain taxa such as ants (Formicidae), termites (Isoptera) or book lice (Liposcelididae) at the genus level (SI Appendix, Table S4) likely due to known primer biases (Clarke et al. 2014; Elbrecht & Leese 2015), limitations of the reference databases, or inherent limitations in sampling dust. Conversely, the DNA-based survey was better able to identify the
presence of moths and butterflies (Lepidoptera) as we were not limited by requiring intact specimens, identifiable parts of Lepidoptera (e.g. scales), or certain life stages (SI Appendix, Fig. S7).

Our approach has some limitations when compared to traditional approaches. For one, it does not yield collections of specimens, or species-level resolution, necessary for additional studies of taxonomy or life history. Nor is it easy to link the proportional abundance of taxa in our samples to true abundances in homes. But our DNA-based survey approach also has a number of key advantages over more standard arthropod diversity assessment methods. Our sampling was not an invasive procedure requiring trained individuals to spend hours combing through a home to conduct intensive surveys. Rather, dust samples could be collected by the citizen scientist volunteers—requiring mere minutes per home. A swab of settled dust from door trim revealed arthropod diversity in the home. Furthermore, our DNA-based approach could identify arthropods from their feces alone or remnant, miniscule parts in dust. Thus, despite the known limitations of using environmental DNA to survey arthropod diversity (see (Gibson et al. 2014), and the fact that we were using a single dust sample to characterize arthropod diversity found throughout a home, our study allows us to describe arthropod diversity across an unprecedented number of homes located across the United States, thus yielding our first insight into the continental-scale diversity and distributions of arthropods inside homes.

Factors predicting arthropod diversity in homes

Arthropod diversity in dust varied among homes. We were able to detect more than 40 arthropod genera in some individual homes with 72% of the homes having fewer than 5 detectable genera. We expected the diversity of arthropods found indoors to be positively associated with outdoor temperature (Dunn et al. 2009) and precipitation (Jenkins et al. 2011), as these factors are known to influence arthropod diversity outdoors. However, we found that outdoor climatic conditions were not reliable predictors of indoor arthropod diversity. Thus, given the relative homogeneity of environmental conditions inside homes and the presence of many arthropods living inside homes, it is difficult to use the large body of literature focused on outdoor arthropod diversity patterns to infer indoor arthropod diversity patterns.

We did find significant effects of surrounding land use type on indoor arthropod diversity, with rural homes harboring, on average, 50% more arthropod diversity than suburban or urban homes (Fig. 3) (p< 0.001, Wilcoxon test). Likewise, the diversity of arthropods in homes was
predictable from other home characteristics. Specifically, we found elevated arthropod diversity in homes with a basement (Fig. 3) ($p<0.05$, Wilcoxon test), and in homes with either dogs or cats (Fig. 3) ($p<0.001$, Wilcoxon test). The increased arthropod diversity found in homes with basements could be explained by the additional, cave-like habitat a basement offers, harboring uniquely-adapted taxa not found elsewhere in homes (Epps et al. 2014; Bertone et al. 2016). The presence of more diverse arthropod assemblages—across taxonomic orders—in homes with pets may result from the pets acting as dispersal agents—either dispersing arthropods directly associated with their bodies into homes (e.g. fleas and ticks) or by pets inadvertently bringing arthropods, or arthropod DNA, indoors from the outside (SI Appendix, Fig. S8). These arthropod diversity patterns parallel those reported previously for microbial diversity; the presence of dogs and cats also increases the diversity of bacteria and fungi found inside homes (Barberán et al. 2015a). When you welcome a pet into your home, you are also introducing a diversity of other organisms—and their sensitizing allergens—an observation that may explain why children raised in homes with dogs or cats frequently have reduced susceptibilities to allergenic disorders (Ownby 2002).

**Indoor versus outdoor arthropod assemblages**

The dust samples collected from door trim inside homes consistently harbored more arthropod diversity than did outdoor samples collected from exterior door trim; an observation consistent with patterns observed for both home bacterial and fungal communities (Barberán et al. 2015a) (Fig. 3) ($p<0.001$, Wilcoxon test, paired). For those 477 homes for which we had samples from both inside and outside the home (permitting direct comparisons), we found 2.5 times more arthropod genera, on average, inside the home than outside the home. This may be due to a greater level of DNA degradation in outdoor samples. In addition, outdoor dust samples have DNA from arthropods found outside the home, while those dust samples collected from inside homes have arthropod DNA both from those organisms that originated outdoors and those arthropods actually living, and originating from, the interior of the home itself.

We distinguished which of these arthropods are transitory visitors dispersed into the home from outdoors, and which are likely synanthropic (living out their life cycles with humans and originating from inside homes) by comparing the presence of taxa found inside versus outside homes. We identified 22 taxa (out of 703 indoor and outdoor arthropod taxa detected) that were more likely to be identified from inside homes versus outside (SI Appendix, Fig. S9).
‘indoor’ taxa span a wide range of arthropod classes from arachnids (mites) to Insecta (moths, flies, fleas, cockroaches, wasps, etc.), and Collembola (springtails). Taxa more likely to be found indoors could be more common because they are truly synanthropic, or because they preferentially accumulate inside homes. Dust mites (*Dermatophagoides* spp.), carpet beetles (*Anthrenus* spp.), Indian meal moths (*Plodia* spp.), fruit/vinegar flies (*Drosophila* spp.), and fleas (*Ctenocephalides* spp.) are common indoor inhabitants that we expected to find inside homes, as their diets are strongly associated with resources found in many homes (Bertone *et al.* 2016). While fleas feed on us and our pets, other taxa innocuously spend their lives consuming human skin cells or skin products (e.g., sebum), textiles, microbes, and each other. We also found some taxa to be far more abundant indoors than outdoors which we did not expect to be living inside homes, including diverse herbivores: aphids (*Aphididae*), owlet moths (*Noctuidae*), planthoppers (*Delphacidae*), and parasitoid wasps (*Rhopalosomatidae* and *Braconidae*). We were more likely to find evidence of aphids in home dust than we were cockroaches, despite the latter receiving more attention as a pest and source of allergens. It may be that these diverse, often tiny, flying insects are lured into homes by unintentional attractants generated by humans in built spaces, or via passive collection (as in Malaise traps). Additionally, human occupants may be directly transporting some of these arthropods into the home as hitchhikers on ornamental plants.

Given the high frequency of aphids detected inside homes, we investigated this group in more detail to determine if known aphid parasitoids and predators tended to co-occur with their prey. If they do co-occur, it would suggest that we can reconstruct known biological interactions from the arthropod DNA left in the collected dust samples. We found that parasitoid wasps (*Braconidae: Aphidiinae*) tended to co-occur with aphids (*Aphididae*) more than expected by chance. In other words when we find homes with aphid parasitoids, we are far more likely than expected by chance to also find aphid DNA in the same sample (p < 0.001, Chi square) (*SI Appendix*, Fig. S10). Likewise, when we looked for DNA from known predators of aphids (e.g. *Coccinella* spp., *Coleomegilla* spp., *Harmonia* spp., *Hippodamia* spp., *Hyperaspis* spp., *Scymnus* spp., and *Aphidoletes* spp.) they were also more likely to be found in homes with aphid DNA (p < 0.001, Chi square) (*SI Appendix*, Fig. S10).
The biogeography of key arthropod taxa

Some of the more frequently identified arthropods from indoor dust samples are considered to be of critical importance to human health, e.g. allergenic dust mites and cockroaches, or are perceived as nuisance pests of homes, e.g. pestiferous ladybugs, boxelder bugs, and invasive cockroaches. For these taxa, we used our analyses of the indoor dust samples to identify where these taxa are distributed throughout the United States (Fig. 4, see SI Appendix, Fig. S11 for additional maps).

No continental scale maps exist for most of the 600+ taxa we identified in this study. However, because of the prevalence and significant economic costs associated with dust mite allergies, considerable effort has gone into building maps of predicted dust mite distributions (Portnoy et al. 2013). We found that dust mites were more likely to be detected in homes located in more humid regions of the United States (Fig. 4), patterns that overlap well with the predicted distributions of dust mites reported in Portnoy et al. (2013). Thus, even though we surveyed a small amount of settled dust from door trim, we were still able to reconstruct the expected geographic distributions of dust mites. This is particularly surprising, as dust mite allergens are not typically found on such hard surfaces (Portnoy et al. 2013). As many arthropod taxa shed potentially sensitizing allergens, maps of dust mites and other potentially allergenic arthropods—including cockroaches, aphids, ladybugs, and even various shellfish (Fig. 4, SI Appendix, Figs. S3, S10, S11)—provide baseline information on the distributions of known, or likely, allergenic arthropods across the United States (Gaillard 1950; Esch et al. 2001; Arlian 2002). Such information could be integrated with data on antigen sensitivities to better understand allergenic arthropod exposures across the United States and their potential health ramifications.

Our geographic distribution maps also reflect the local outdoor ecology of other arthropod taxa. For example, the detected northern distribution of ladybugs (Harmonia spp.) in homes is consistent with this allergenic pests' cold intolerance, which leads them to seek shelter in homes during the winter (Goetz 2009) (Fig. 4). Similarly, the nuisance pest, boxelder bugs (Boisea spp.) also overwinter in homes (Cranshaw 2011). As expected from other surveys of house arthropods, boxelder bugs were detected more often in homes that were generally at higher latitudes, and were within the range of their main host plant boxelder trees (Acer negundo L.) (Digital representation of “Atlas of United States Trees” by Elbert L. Little, Jr. 1999; Cranshaw 2011) (Fig. 4).
Many of the taxa we found indoors are not native to the United States, imported as biocontrol agents (e.g., ladybugs like *Harmonia* spp.), or accidentally introduced, such as certain mosquito species (*Stegomyia* spp.). The value of our maps in visualizing the ranges of emerging invasive arthropod taxa is demonstrated in the distributions of the Turkestan cockroach (*Shelfordella* spp.) (Fig. 4). This cockroach is an invasive species known to inhabit the south and southwestern United States, but predicted to spread north and east due to its availability for internet purchase as ‘feeder’ cockroaches (Kim & Rust 2013). We detected this species as far north as Massachusetts (Fig. 4) and in doing so reveal that this cockroach has already spread, undetected.

**Conclusion**

We show that non-invasive DNA-based surveys of settled dust can reveal many of the arthropods present inside homes. As a result, we were able to conduct the first continental-scale survey of arthropods in our homes. Indoor arthropod diversity is, to some degree, predictable from home design, degree of urbanization, and the presence of pets, while the distributions of individual taxa often reflect the outdoor environment. These findings demonstrate the value of dust-based arthropod analyses for reconstructing arthropod food webs in homes and mapping the distributions of important arthropod taxa, including emerging and known arthropod allergens and invasive pest species. Moreover, the results presented here provide baseline information on the arthropods found inside our homes at the continental scale and demonstrate the utility of this DNA-based approach for future work focused on documenting how changes in climate, housing patterns, or building design may alter the arthropods we come into contact with as we go about our daily lives.

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Tables and Figures

Figure 1. Ubiquity of arthropods detected in settled dust samples collected from inside homes. (A) Genera detected. (B) Orders detected in at least 5% of homes. Y-axes indicate the percentage of homes (out of 651 homes with arthropods detected) where those arthropods were detected.

Figure 2. Proportional arthropod diversity—number of unique taxa per order—detected across all indoor samples with each box representing a unique arthropod order and the size and color of the box scaled to reflect the number of unique taxa within that order.

Figure 3. Significant predictors of arthropod diversity inside home dust. (A) Indoor versus outdoor samples (p<0.001, Wilcoxon test, paired). (B) Indoor samples dust in relation to whether the home was in a rural versus a suburban or urban area (p<0.001, Wilcoxon test). (C) Indoor dust samples in homes with or without basements (p<0.05, Wilcoxon test) or (D) with or without pets (p<0.001, Wilcoxon test).

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Figure 4. Distributions of select allergenic, pestiferous and invasive arthropods in indoor dust. (A) Predicted allergenic dust mite (Dermatophagoides spp.) range versus detected range, (B) alien ladybugs (Harmonia spp.), (C) invasive cockroach (Shelfordella spp.), (D) boxelder bugs (Boisea spp.) versus the range of its primary host plant, boxelder trees (Acer negundo L.).
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