The extraordinary diversity of beetles (order Coleoptera; >400,000 species) has been attributed chiefly to the adaptive radiation of specialized herbivorous beetles feeding on flowering plants (angiosperms) (1–4). However, the evolution of herbivory and its impacts on beetle diversification remain poorly understood. We used large-scale genomic data to infer the phylogeny of beetles, reconstruct timing and patterns of beetle diversification, and trace the evolution of beetle genes enabling specialized herbivory. Our results shed light on the evolution of plant feeding and reveal endogenous plant cell wall-degrading enzymes (PCWDEs) as a key innovation in the adaptive radiation of beetles on plants. Nearly half of all herbivorous insect species are beetles (1), and most herbivorous beetles feed on angiosperms. Nonetheless, recent studies have failed to find a strong positive relationship between herbivory—including herbivory on angiosperms—and beetle diversification (4–7). In contrast to strictly ecological explanations, recent studies of beetle genomes (8) and beetle digestive physiology (8–12) have speculated that the diversity of herbivorous beetles may have its origins in genomic innovation, specifically the evolution of endogenous PCWDEs enabling symbiont-independent digestion of lignocellulose in plant cell walls.

**Significance**

We inferred the phylogeny and evolution of beetles using genomic data of an unprecedented scale. Moreover, we documented the diversification of plant-feeding (herbivorous) beetles, which account for nearly half of all beetle species and a similar proportion of herbivorous insects, following convergent horizontal transfers of bacterial and fungal genes enabling the digestion of lignocellulose in plant cell walls. Our findings clarify beetle phylogenetic relationships and reveal new insights into the evolution of specialized herbivory and why there are so many species of beetles. Furthermore, they underscore the intimacy and complexity of the evolutionary relationships between insects, plants, and microorganisms and show how analyses of large-scale genomic data are revealing the evolution and genomic basis of insect biodiversity.

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Data deposition: The 1KITE transcriptomes reported herein are permanently archived at the National Center for Biotechnology Information (NCBI), https://www.ncbi.nlm.nih.gov/bioproject/183205 under the Umbrella BioProject ID PRJNA 183205. Datasets S1 through S4, including the resulting alignments of phylogenomic data from beetles and their putative POWDEs, and gene trees and BLAST hits for candidate POWDEs, are archived at Zenodo, DOI: 10.5281/zenodo.3522944.

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on Earth—liberating sugars, amino acids, and other essential nutrients (SI Appendix, Table S1) (9, 10). Apart from GH families 1 and 9, which have ancient origins in animals (14), beetle-encoded PCWDEs are thought to have originated from bacteria and fungi via horizontal gene transfer (HGT) (9, 15). Gene duplication and functional diversification post-HGT has created multigene families of PCWDEs with complementary catalytic activities (16).

PCWDEs (other than GH1 and GH9) are reported from the genomes of fewer than 50 beetle species—all specialized herbivores—representing 5 of the 190 described extant beetle families (SI Appendix, Table S2). Their presence in beetle genomes is notable because it was previously thought that beetles, like many other herbivorous animals, lack the capacity for significant endogenous lignocellulose digestion (13) and therefore require symbionts (whose genomes encode PCWDEs) for successful herbivory. While PCWDEs have been shown to play essential roles in beetle–plant interactions (8, 11), their phylogenetic distribution and impacts on beetle diversification remain virtually unknown (17).

Results and Discussion
Phylogenetic Relationships. We reconstructed the phylogeny of beetles using 1,907,014 amino acid sites from 4,818 nuclear genes obtained via RNA sequencing (RNA-Seq) or genome skimming (SI Appendix, Methods, Figs. S1–S14). Apart from GH family 1, we recovered a synapomorphy of the 2 families (33, 35). Upper Permian (Changhsingian Stage, ca. 254 to 252 Ma, China) wood borings were recently attributed to beetles (36, 37), indicating an early origin of wood-boring habits in Coleoptera.

Our estimates for beetle net diversification rates were not exceptional, consistent with previous studies (3–5). Furthermore, significant increases or decreases in net diversification rate were recovered for most of the same higher-level taxa of beetles as previous studies (Fig. 2) (3–5). The timing of branching events in the beetle phylogeny was in general agreement with ref. 4 and ref. 38 for deep nodes but somewhat older than most other (earlier) studies (e.g., refs. 3 and 5). However, node ages within series Cucujiformia, which contains most herbivorous beetle species, were mostly much younger than in ref. 38, similar to the results of ref. 4.

We recovered evidence for adaptive radiations of beetles into a variety of ecological niches. For example, predaceous ground beetles (Carabidae) and aquatic diving beetles (Dytiscidae and related) experienced rapid radiations in the early Jurassic increases in diversification rate. Early splits in herbivorous Buprestoidea (metallic wood-boring beetles) and Phytophaga, and herbivorous/saprophagous scarabs (Scarabaeidae) occurred in the Jurassic and early Cretaceous, which is also when core angiosperms diversified (39, 40). These same beetle groups subsequently underwent crown diversification coeval with the rise of angiosperms to ecological dominance during the mid-Cretaceous (41) (Figs. 1 and 2 and SI Appendix, Figs. S11–S14). One-fourth of all rate increases in the beetle phylogeny were associated with herbivory (Fig. 2)—which is more than any other factor (Fig. 3).

Other groups of beetles that are species-rich in the modern fauna and associated with living plants or plant litter, including detritivores, saprophages, and mycophages, also diversified during the middle to late Mesozoic (e.g., rove beetles [Staphylinidae], scarabs [Scarabaeidae], and darkling beetles [Tenebrionidae]), consistent with the view that ecologically diverse groups of modern beetles coevolved with angiosperms and in the novel habitats that angiosperms created (3). Within Phytophaga, the weevil families Caridae and Cimbridae experienced significant decreases in diversification rate during the Mesoico (Fig. 2). These groups are thought to have ancestral associations with gymnosperms (specifically, conifers) (2), in contrast to the angiosperm associations of most extant Phytophaga.

Beetle groups that are today associated with gymnosperm pollen, including some early Cucujoida and Phytophaga (Boganiidae: Paracucujinae, Cimbridae, most Nemonychidae, and Erotylidae: Archostemata, and Myxophaga (Methods and SI Appendix, Figs. S8 and S9).
Fig. 1. A dated phylogeny of beetles showing the distribution of putative PCWDEs, inferred from 4,818 nuclear genes (Methods and SI Appendix). Branches in suborder Polyphaga are color-coded by superfamily. Numbers indicate nodes constrained by fossil priors (SI Appendix, Table S5). Filled squares indicate the presence of putative PCWDEs and GH32 invertases (color-coded by gene family) based on analyses of whole-genome (asterisk) or RNA-Seq data. GH1 and GH9 have known ancient origins in metazoans (14, 47) and were expected to occur in most species. Asterisks denote results from the analysis of WGS (versus RNA-Seq) data. Numbers of homologs are indicated in each box when previously published (SI Appendix, Table S1). Note that Rhinorhipus was added after the initial analyses were completed based on a new ML tree search, which recovered the same topology. Bootstrapping was not conducted due to computational constraints. However, its placement was the same in the S21-taxon tree, where it had 100% ML bootstrap support. All higher taxa shown to illustrate morphological diversity (but not all species) were sampled. Cupes image courtesy of Matthew Bertone (North Carolina State University, Raleigh, NC). All other photos courtesy of Udo Schmidt (photographer).
Pharaxonothinae), appeared by the late Jurassic, well before bees and butterflies (21, 42). They were likely among the first insect pollinators of gymnosperms and early angiosperms. Furthermore, our results are consistent with fossil evidence in suggesting that pollinivory was a transitional state between detritivory, mycophagy, and saprophagy (Cucujodea) and specialized herbivory (Phytophaga) (43–45) (Fig. 1 and SI Appendix). The apparent prevalence of transitions in Coleoptera from generalized diets, such as detritivory and saprophagy, toward more specialized diets, such as mycophagy and herbivory, is consistent with the high rate of such transitions across insects (6).

**Comparative Genomics of Beetle-Encoded PCWDEs.** We studied putative PCWDEs encoded in 154 transcriptomes or genomes corresponding to the 147 taxa in Fig. 1 (SI Appendix, Figs. S15–S27, and Tables S1, S2, and S8). We also studied GH32 invertases, which catalyze the conversion of sucrose—the primary form of photoassimilated carbon in plant vascular tissues—to glucose and fructose. Like PCWDEs, invertases have played a potentially important role in the evolution of specialized herbivory (e.g., see ref. 46).

GH1 and GH9 have ancient origins in animals (14, 47) and were nearly ubiquitous in our study. The other gene families we studied were found almost exclusively in Buprestoidea and Phytophaga, which encode an expansive, remarkably similar array of PCWDEs (Figs. 1 and 3). Buprestoidea and Phytophaga are the most species-rich and most specialized radiations of herbivorous beetles (1). Their feeding habits collectively include chewing, mining, and boring of virtually all kinds of plant tissues (living or dead) and plant taxa.

Outside of Buprestoidea and Phytophaga, 10 beetle species scattered widely across the phylogeny had matches to 1 PCWDE gene family (other than GH1 and GH9), and 2 beetle species each matched 3 gene families (Figs. 1 and 3). These included Bostrichidae (Xylobiops, GH32), Cleridae (Thanasisinus, GH48), Elateridae (Melanotus, GH32), Lycidae (Porrostoma, GH32), Melyridae (Anthoconus, GH32), Micromalthidae (Micromalthus, GH10), Oedemeridae (Oedema, GH10), Ptiliidae (Acrotrichis,
GH45), Ptinidae (Ptinus, GH28, GH32, and GH45), and Zopheridae (Pycnomenus, GH10, GH28, and GH45). In contrast, Buprestoidea and Phytophaga had species with matches from up to 7 families of PCWDEs, often with multiple apparent homologs from each family. Independent losses and reacquisitions of PCWDEs are known in Phytophaga (10) and were observed in this study.

Overall, we documented putative endogenous PCWDEs (including GH32 invertases, excluding GH1 and GH9) from 22 families of beetles. Previous to this study, they were known from only 5 beetle families (Figs. 1 and 3 and SI Appendix, Tables S1 and S2, and Fig. S27). Within Buprestoidea, we report GH10, GH45, GH48, and CES, in addition to previously reported genes. Also within Phytophaga, we document PCWDEs from the families Attelabidae, Belidae, Brentidae, Caridae, Megalopodidae, and Nemonychidae, in addition to the families of Phytophaga from which these genes have been previously reported. Thus, we significantly expand knowledge of the phylogenetic distribution of putative PCWDEs encoded in the genomes of Coleoptera, while at the same time establishing that they are particularly diverse in the 2 lineages which these genes have been previously reported. Thus, we significantly expand knowledge of the phylogenetic distribution of putative PCWDEs encoded in the genomes of Coleoptera, while at the same time establishing that they are particularly diverse in the 2 lineages which these genes have been previously reported.

Microbial Donors of Beetle PCWDEs via HGT. The inferred last common ancestors and potential donors of beetle PCWDEs were bacteria and fungi, including taxonomic groups that are today quintessential degraders of lignocellulose and other complex polysaccharides in plant and soil detritus (Fig. 3, SI Appendix, Figs. S15–S26, and Datasets S1–S3) (48). Some of these groups of bacteria and fungi are also found in beetle guts (49). Beetle-derived PCWDEs nonetheless formed well-supported clades distinct from microbial taxa in our phylogenies. Moreover, they were largely placed within the same clades as their homologs derived from other insect genomes and transcriptomes, including those derived from high-quality draft genomes. Within these clades, some gene families contained clusters of closely related sequences from the same beetle higher taxa or species, consistent with lineage-specific gene duplications post-HGT.

Physical incorporation of genes encoding PCWDEs into the genome of one or more beetle species has been documented for Buprestoidea-derived GH28, GH32, GH43, GH44, and PL4 and for Phytophaga-derived GH5-2, GH5-8, GH10, GH28, GH32, GH45, GH48, and CES (16, 50, 51) (SI Appendix, Table S1). Enzyme product functionality (metabolic integration) has been demonstrated for Buprestoidea-derived GH43, GH44, and PL4 and for Phytophaga-derived GH5-2, GH5-8, GH10, GH28, GH32, GH45, and CES (16, 50, 51). For GH5-2, GH28, GH45, and CES, a similar gene has been independently horizontally transferred to a plant-feeding organism outside of Insecta (16).

Evidence from the available high-quality draft genomes of Buprestoidea and Phytophaga further indicates that these genes are encoded in beetle genomes and are not the result of contamination (8, 17). The emerald ash borer beetle (Agrilus planipennis; Buprestoidea) genome encodes GH28, GH32, GH43, GH44, and PL4 (4) (each represented by multiple copies in the genome; Fig. 1), all of which have been PCR-amplified from adult A. planipennis elytra and legs—tissues not known to contain symbionts.
(50, 51). These genes are also frequently arranged in tandem arrays on scaffolds containing other beetle genes. For example, 3 of the 5 *A. planipennis* GH43 genes reside in the same genomic scaffold (8, 51). Four of the 5 GH43 genes have single exons (51), which is unusual, since almost all of the GH genes we studied in beetle genomes contain multiple exons. Furthermore, the 5 gene families encoding *A. planipennis* PCWDEs are almost exclusively expressed in the larval midgut (50, 51). Together with the presence of N-terminal secretory signal peptides in the putative proteins, this suggests that these enzymes are secreted to facilitate plant cell wall digestion (51).

Similar observations have been made using the genomes of herbivorous Phytophaga. For example, the Asian longhorned beetle (*Anoplophora glabripennis*; Phytophaga: Chrysomeloidea) genome contains multiple copies of GH5, GH28, GH45, and GH48 and a single copy of GH32 (8, 52), and the mountain pine beetle (*Dendroctonus ponderosae*; Phytophaga: Curculionoidea) genome contains GH28, GH32, GH45, GH48, CE8, and PL4, all of which are multicyclic in the genome (10, 52–55). Many of the GH family genes from *A. glabripennis* and *D. ponderosae* have been PCR-amplified from beetle tissues not known to contain microbial symbionts (10, 53–55). Similarly, many of these genes have been functionally characterized in one or both species using complementary DNAs generated from RNA samples obtained from individual beetles different from the ones used for genome sequencing (8, 10, 55). Furthermore, the *A. glabripennis* and *D. ponderosae* PCWDEs (like those encoded in the *A. planipennis* genome) are frequently arranged in tandem arrays on genomic scaffolds containing other beetle genes (8, 54).

**Convergent Evolution of Beetle PCWDEs.** The appearance of similar expansive arrays of PCWDEs in Buprestoidae and Phytophaga, separated by over 250 Ma of evolution, appears to result from convergent evolution via HGT, rather than vertical transmission from a common ancestor (Figs. 1 and 3). However, the mechanisms behind these HGT events remain obscure. Gene families in common (excluding GH1 and GH9) and thus candidates for convergence included GH10, GH28, GH32, GH43, GH45, GH48, CE8, and PL4; only GH5 and GH44 cellulases were not shared, perhaps reflecting slightly different strategies for PCW digestion. PCWDEs were absent from the near relatives of Buprestoidae and Phytophaga and most other Coleoptera. Furthermore, we recovered separate clades of genes corresponding to PCWDEs from Buprestoidae and Phytophaga.

Among the Phytophaga, the diversity of gene families encoding PCWDEs and the number of gene family members both appear to increase in the phylogeny from root to tips (Fig. 1). This suggests the stepwise evolution of symbiotic-independent mechanisms for plant cell wall degradation from symbiotic-dependent ones, although other scenarios are possible. Amplification and functional divergence of PCWDEs post-HGT (8) may have facilitated the evolution of increasingly specialized plant-feeding habits in Buprestoidae and Phytophaga, including the exploitation of woody tissues and pectin-rich young leaves and stems, seeds, and fruits (2, 19, 56). The existence of multiple copies of these genes in most beetle genomes may reduce constraints on their functional evolution, facilitating substrate diversification (activity toward additional/different plant cell wall polysaccharides) post-HGT. GH48 genes, which were found in most Buprestoidae and Phytophaga, may also help degrade fungal chitin (8, 57) and were likely relevant to the repeated radiation of specialized fungus-feeding habits in Phytophaga (2).

**Adaptive Radiation of Herbivorous Beetles Post-HGT.** Remarkably, in both Buprestoidae and Phytophaga, the origins of PCWDEs are phylogenetically and temporally linked to significant increases in net diversification rate—1) along the stem of Buprestoidae and 2) along the stem of Phytophaga—indicative of adaptive radiations (58) (Figs. 2 and 3). This suggests that PCWDEs enabled Buprestoidae and Phytophaga entry into new adaptive zones, notably including penetration of the woody plant barrier, without needing to obtain and maintain symbionts for PCW degradation. Pectinases and invertases likely promoted feeding on cambium and sapwood, fruits, leaves, and seeds, while (hemi)cellulases may have promoted feeding on wood (59). Endogenous PCWDEs were therefore likely key to the evolution of specialized plant-feeding habits, such as leaf mining and seed, stem, and wood boring, which also required dealing with novel plant allelochemicals, nutritional and defensive barriers, and recalitrant plant biopolymers (57).

Phytophaga began to diversify ~50 Ma earlier than Buprestoidae, perhaps giving the former an evolutionary advantage and accounting for the lesser taxonomic and ecological diversity of extant Buprestoidae—typically, the first organisms to enter an adaptive zone have an advantage (60). Furthermore, while the wood-feeding habits of Buprestidae are similar to those of certain Phytophaga, the Buprestoidae and Phytophaga also exhibit significant differences in life history, behavior, and trophic habits, with potential implications for their abilities to transition between ecological adaptive zones and their resulting diversification rates. More in-depth taxon sampling is needed to further elucidate timing and patterns of gene gain, loss, and amplification, especially in Buprestoidae. Additionally, large-scale functional genomic studies are needed to characterize the roles of the candidate genes we have identified in herbivory and diet specialization among Coleoptera, that is, beyond the relatively few beetle species and genes that have been studied to date. Moreover, additional genes are known to play roles in plant cell wall degradation (and detoxification of plant allelochemicals) but remain little studied in beetles (8, 14).

**Ecological Opportunity and Evolutionary Innovation at the Beetle–Plant Interface.** Buprestoidae and Phytophaga exhibit a similar pattern of increasing host tissue specificity temporally and phylogenetically linked to the evolution of endogenous PCWDEs. We propose that the specialized but versatile trophic apparatus and evolving metabolic repertoire of Buprestoidae and Phytophaga helped them adapt to and track the increasing diversity of angiosperms during the Mesozoic, despite an escalation in the potency and variety of angiosperm chemical defenses (61) and diversification of angiosperm plant cell walls (62). Moreover, facultative symbionts that code for some of the same PCWDEs as beetle genomes (57) likely increased digestive efficiency. We speculate that ecological constraints proposed to render host specialists more susceptible to extinction, such as prolonged larval development and small population and range sizes, were ameliorated by the increasing efficiency of plant biomass assimilation afforded by endogenous PCWDEs, and by amplification and functional divergence of PCWDEs post-HGT. Moreover, this may have facilitated an even greater degree of host specificity, resulting in the evolution of increasingly more varied and specialized plant-feeding habits (63).

Early associations of Phytophaga with gymnosperms, while perhaps initially limited to cone and pollen feeding, thus expanded over time to include virtually all kinds of plant tissues and taxa. Transitions within Buprestoidae from external root feeding to specialized wood boring and leaf mining (56) may also be interpretable in this framework. Mycophagy and feeding on fungus-infested plant tissues were potentially important transitional states to specialized feeding on wood in Phytophaga and their ancestors (3, 64, 65). Phytophaga likely also transitioned from mutualistic interactions (pollenivory/pollination) to antagonistic ones (specialized herbivory on leaves and stems) with concomitant ecological impacts. The phylogenetic placement of beetle taxa that are today associated with gymnosperm cones or pollen, for example Boganiiidae: Paracucujinae and Erotylidae:
Phylogenetic Inference. We reconstructed the phylogeny of Coleoptera using phylogenomic data from 135 species representing 90 families, plus 11 outgroups (SI Appendix, Fig. S2 and Table S3). We used ML inference implemented in the program RAxML v8.2.10 (69) for phylogeny reconstruction. We then conducted a Bayesian relaxed clock divergence time analysis in MCCMCTree (70) with fossil constraints applied to 18 nodes in the phylogeny (Fig. 1 and SI Appendix, Table S5). Our study was designed to address concerns of taxon and locus diversity, outgroups, fossil selection, and age constraints that have been proposed to impact other molecular phylogenetic studies of beetles (e.g., refs. 3-5 and 38).

Diversification Rates. Net diversification rates and the temporal and phylogenetic locations of diversification rate shifts were estimated using a near-comprehensive family-level time tree generated for the same 146 species as above, plus Rhinorhipus and 374 additional species from ref. 4. (521 total species in 140 beetle families; hereafter, the 89-gene tree; SI Appendix, Table S6 and Fig. S10). This tree was subsequently expanded to include 188/190 extant beetle families (SI Appendix, Supplementary Information Text), followed by analysis in MCCMCTree with age constraints applied to 22 nodes (SI Appendix, Figs. S8 and S11 and Table S5). Only the beetle families Joridae and Cossiniiellidae, each with one extant species known only from their original type series, were missing from these analyses. Net diversification rates were estimated using MEDUSA in the Geiger package (71).

PCWDEs. We searched the whole-genome sequencing (WGS) and transcriptome data for the presence of GH genes encoding PCWDEs (SI Appendix, Table S8). GH genes encode enzymes that catalyze the hydrolysis of glycosidic bonds in oligo- and disaccharide sugars and play essential roles in degrading lignocellulose in plant cell walls and other plant polysaccharides. Some also play a role in the detoxification of plant allelochemicals (72). We also searched for pectin methyltransferases belonging to carbohydrate esterase family 8 (CE8), which catalyze the deestickering of pectin into pectate and methanol, and pectin lyases belonging to polysaccharide lyase family 4 (PL4), which cleave alpha-1,4 glycosidic bonds between 1-rhamnose and α-galacturonic acids in pectin (SI Appendix, Table S8). Pectinolytic enzymes facilitate feeding on the pectin-rich primary cell walls of sapwood and cambium, leaves, fruits, and seeds, while (hem)cellulolytic enzymes are essential for feeding on tissues rich in secondary plant cell walls, including heartwood. Finally, we searched for GH32 invertases, which catalyze the conversion of sucrose to glucose and fructose (46) (SI Appendix, Table S8).

The phylogenetic distribution of these gene families in Coleoptera was determined by identifying protein family domains in our WGS and transcriptome data (SI Appendix, Table S3) using PfamScan (73). We then used BLASTp (74) to refine these results. Subsequently, we gathered homologs for each gene of interest, and after alignment, implemented ML phylogenetic analyses using IQ-TREE (75). We explored the impact of endogenous PCWDEs on beetle diversification rates by comparing the phylogenetic location of significant increases in beetle net diversification rate with the phylogenetic distribution of gene families encoding putative PCWDEs in the 4,818-gene phylogeny. Whole-genome or RNA-Seq data were available for every terminal in this tree. The 1KITE transcriptomes reported herein are permanently archived at under the Umbrella BioProject ID PRJNA 1832105. Datasets S1 through S4 are archived at the Zenodo Digital Repository at 10.5281/zenodo.3522944 (78).

Data Use Statement. Data on genetic material contained in this paper are published for noncommercial use only. Utilization by third parties for purposes other than noncommercial scientific research may infringe the conditions under which the genetic resources were originally accessed, and should not be undertaken without obtaining consent from the corresponding author of the paper and/or obtaining permission from the original provider of the genetic material.

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