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✉ For correspondence:

agrawal@cornell.edu

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Cardenolide toxin diversity impacts monarch butterfly growth and sequestration

Anurag A Agrawal^{1,2} ✉, Amy P Hastings¹, Paola Rubiano-Buitrago¹

¹Department of Ecology and Evolutionary Biology, Cornell University, Ithaca, United States • ²Department of Entomology, Cornell University, Ithaca, United States

eLife Assessment

This **important** study shows that different forms and mixtures of cardenolide toxins in tropical milkweed, especially nitrogen- and sulfur-containing types, change how monarch caterpillars eat, grow, and store these chemicals under laboratory conditions. It provides **solid** evidence to demonstrate that chemical diversity within a single group of plant toxins (cardenolides) can have combined effects on even highly specialized herbivores that are different from what one would expect from each toxin alone. However, as all experiments used leaf-disc assays with fixed "natural" toxin ratios and only one adapted herbivore species, tests on living plants, other mixture designs, and non-adapted herbivores would make the broader conclusions stronger.

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Abstract

In classic coevolutionary interactions, host plants are thought to accrue novel chemical defenses which are later countered by detoxification strategies and sometimes sequestration in specialist herbivores. We recently discovered that unusual nitrogen- and sulfur-containing (N,S-) cardenolides in some milkweed species are highly toxic, and broken down to less toxic forms which are sequestered by monarch butterflies (*Danaus plexippus*). Here we isolated and purified five dominant cardenolide toxins from the tropical milkweed, *Asclepias curassavica*, a globally abundant host plant of monarchs, and fed them to caterpillars individually or in mixture. We hypothesized that the two N,S-cardenolides in *A. curassavica* (uscharin and voruscharin) would reduce caterpillar growth and sequestration more than other abundant related cardenolides (15-Hydroxy calotropin, frugoside, calactin). Overall, cardenolide treatments caused monarchs to feed more and grow more compared to controls; nonetheless, one N,S-cardenolide (voruscharin) was not stimulatory and caused substantial reductions in growth efficiency. Consuming N,S-cardenolides caused caterpillars to sequester the lowest total amounts of cardenolides, and also reduced their efficiency of sequestration. We next tested the phytochemical diversity hypothesis, that toxin mixtures pose a substantial burden for caterpillars compared to individual compounds provided in equimolar concentrations. We prepared two types of mixtures, one containing equal concentrations of the five compounds and another "realistic mixture" where toxin concentrations reflect their natural proportions in leaves. Mixtures had a negative impact on caterpillar feeding, growth, sequestration, and sequestration efficiency compared to the average of single compounds. The equal and realistic mixtures had similar impacts on feeding and growth, but feeding on the realistic mixture resulted in the lowest sequestration. We conclude that as a result of coevolutionary interactions, even sequestering herbivores may be thwarted by highly specialized plant metabolites such as N,S-cardenolides, and that phytochemical mixtures strengthen plant defense. Toxin mixtures likely challenge detoxification and transport of plant defenses, reducing the herbivore's growth and sequestration.

Introduction

Individual plants produce thousands of so-called secondary compounds, those with no known function in primary metabolism (i.e., resource acquisition and allocation), many of which are defensive (Fraenkel 1959 [↗](#)). Although the benefits of plant defensive chemistry are well-established (Mauricio and Rausher 1997 [↗](#), Schoonhoven et al. 2005 [↗](#), Agrawal 2011 [↗](#)), why plants produce such a diversity of secondary compounds has long been a mystery (Romeo et al. 1996 [↗](#), Rasmann and Agrawal 2009 [↗](#), Speed et al. 2012 [↗](#)). Defenses from distinct chemical classes (e.g., protease inhibitors and alkaloids) can interact in myriad ways (Steppuhn and Baldwin 2007 [↗](#)), but even within a class, a remarkable amount of functional and structural diversity exists. When diversity exists within a class of compounds sharing the same mode of action (e.g., cyanogenic glycosides, cardenolides, or ellagitannins) and when deployed in the same plant tissues, a defensive benefit of deploying multiple compounds is often predicted; nonetheless, the proximate and ultimate explanation for this diversity is not well-understood (Richards et al. 2016 [↗](#), Kessler and Kalske 2018 [↗](#), Wetzel and Whitehead 2020 [↗](#)). Are individual compounds targeting different plant attackers or do mixtures act as a more effective defense than individual compounds alone?

Within a defense class, variation in the biological activity and ultimately the potency of a particular compound against herbivores may be derived from structural attributes (Berenbaum and Zangerl 1996 [↗](#), Romeo et al. 1996 [↗](#), Macel et al. 2005 [↗](#), Zaman et al. 2025 [↗](#)). For example, alkaloids are perhaps the best studied group of phytochemicals, with known biosynthetic pathways, structural modifications (e.g., glycosylation, oxidation, annulation), and differential impacts on biological targets (Bhambhani et al. 2021 [↗](#)). Such structural variation impacts the molecular complexity, physicochemical properties (e.g., polarity), and potentially specific interactions with the physiological target in an animal consumer. As a case in point, among cardenolide toxins produced by milkweeds (*Asclepias* spp., Apocynaceae), all having the same mode of action (binding to Na/K-ATPase in animal cells) (Agrawal et al. 2012b [↗](#)), hundreds of compounds have been identified, ranging from 350 to 1066 Daltons and with a broad range of polarity and structural features (Rubiano-Buitrago et al. 2025 [↗](#)). In particular, only about 5% of milkweed cardenolides have major structural modifications beyond the steroidal core, unsaturated lactone, and sugar moieties, and these have been proposed as the most complex and potent cardenolides (Rubiano-Buitrago et al. 2025 [↗](#)). Our previous work has shown >1000-fold variation in the *in vitro* functional toxicity of different cardenolides on different herbivore target enzymes (Agrawal et al. 2021 [↗](#), Agrawal et al. 2022 [↗](#)). Nonetheless, we are still looking for a predictive framework for understanding the diversity of cardenolide compounds and their ecological impacts.

In addition to individual toxins varying in their biological activity, we still understand relatively little about their combined effects when produced as mixtures (as they are typically consumed in nature). When presented together, secondary compounds may sum to a greater impact on herbivores than can be predicted by any single effect (termed a *phytochemical diversity effect*) (López-Goldar et al. 2024 [↗](#)). Experimental approaches to studying effects of phytochemical diversity within a compound class have been increasing and are often conducted from the herbivore's perspective (Berenbaum and Zangerl 1996 [↗](#), Richards et al. 2016 [↗](#), Whitehead et al. 2021 [↗](#), López-Goldar et al. 2024 [↗](#), Zaman et al. 2025 [↗](#)). We still know rather little about the impact evenness within mixtures, and whether small quantities of particular compounds may shape the net biological effect within mixtures (López-Goldar, 2024 #10996; Glassmire, 2020 #11050; Wetzel, 2020 #10869). In summary, despite the widespread occurrence of substantial phytochemical diversity within a chemical class and even in a single plant species, substantial work remains to enhance the methods, realism, and predictability such that the general consequences of phytochemical diversity can be evaluated.

Here we take an organismal approach to test effects of phytochemical diversity on the monarch butterfly (*Danaus plexippus*), with a special focus on structurally diverse cardenolide toxins. In particular, we address key challenges in advancing our understanding of phytochemical diversity

by 1) isolating the specific compounds from a single plant species, 2) administering the compounds in a realistic way to an herbivore adapted to the host plant, and 3) controlling total toxin concentrations across treatments to allow for direct comparisons without confounding toxin concentration. We have been studying the impacts of specific toxins using isolated and purified cardenolides from the foliage of tropical milkweed (*A. curassavica*), a major host plants of the monarch. In particular, unusual compounds with nitrogen and sulfur containing heterocycles (e.g., uscharin and voruscharin, hereafter N,S-cardenolides), are highly complex, non-polar, and the most toxic compounds known on the monarch Na/K-ATPase (Agrawal et al. 2021 [Agrawal et al. 2021](#), Agrawal et al. 2024b [Agrawal et al. 2024b](#)). Nonetheless, the *in vivo* impacts of these compounds on growth, sequestration, and physiological efficiencies (e.g., mass gained, or toxins stored per unit leaf consumed) have not been previously studied.

Here we start by addressing the relative impact of five dominant cardenolides from *A. curassavica* on monarch growth and sequestration individually. Accordingly, we first test the hypothesis that these N,S-cardenolides (uscharin and voruscharin, comprising up to 50% of total foliar cardenolides) reduce monarch feeding, growth and digestive efficiency compared to three other cardenolides from *A. curassavica*. We expected these compounds to be broken down to less toxic forms before sequestration (Agrawal et al. 2021 [Agrawal et al. 2021](#), Agrawal et al. 2024b [Agrawal et al. 2024b](#)), potentially reducing sequestration overall and sequestration efficiency compared to other cardenolides. Finally, we tested the phytochemical diversity hypothesis, predicting that cardenolide mixtures would reduce growth and sequestration more so than equimolar concentrations of single compounds. We address this diversity hypothesis with two types of mixtures: equal concentrations of the five compounds, versus a more realistic mixture with the compounds in proportion to their relative abundance in *A. curassavica* leaves. In sum, this work addresses the impact of a natural diversity of milkweed toxins on a native and highly adapted herbivore via impacts on digestive and defensive ecophysiology.

Results

Characterizing structural diversity of isolated cardenolides

The five dominant cardenolides of *A. curassavica* are relatively similar in molecular weight ($\approx 10\%$ diff between lowest and highest), but vary more substantially in our estimates of their chemical complexity (29%), polarity (WLOGP: 64%, retention time: 82%), and toxicity to the monarch enzyme (28-fold) (Table 1 [Table 1](#)). In particular, frugoside and calotropin are part of the proposed biochemical pathway leading to the production of the N,S-cardenolides voruscharin and uscharin, while 15-Hydroxy-calotropin is a small structural modification of calotropin (with a hydroxyl group added, Fig. 1 [Fig. 1](#)). Among the group, the N,S-cardenolides are the heaviest, most non-polar, most complex, and have the highest *in vitro* inhibitory capacity of the monarch Na/K-ATPase compared to the other three compounds (Table 1 [Table 1](#)).

Effects of single cardenolides: growth and feeding

Compared to monarchs feeding on control leaves, experimental addition of individual cardenolides increased caterpillar growth rates, and this effect was significant for frugoside, calotropin, and N,S-containing uscharin, but not the other N,S-containing cardenolide, voruscharin or for 15-hydroxy-calotropin ($F_{5,91}=3.985$, $p=0.003$, Fig. 2A [Fig. 2A](#)). Across all treatments, individual caterpillars grew more when they ate more leaf tissue, and the slopes of these relationships were the same across treatments (predicting growth: dry mass consumed $F_{1,80}=245.87$ $p<0.001$; cardenolide treatment $F_{5,80}=1.04$, $p=0.399$; interaction $F_{1,80}=0.465$, $p<0.802$, Fig S1A [Fig S1A](#)). In other words, three of the five compounds stimulated feeding, including the most potent N,S-cardenolide.

The amount of frass excreted, a component of digestion, was differentially impacted by our cardenolide treatments; in particular, when consuming N,S-containing uscharin, less frass accumulated per leaf mass consumed than for the four other cardenolides (leaf mass consumed $F_{1,79}=180.48$ $p<0.001$; treatment $F_{5,79}=2.48$, $p=0.036$; interaction $F_{1,79}=2.43$, $p=0.042$, Fig S1B [Fig S1B](#)). Nonetheless, the efficiency of conversion of digested matter (ECD, the proportion of assimilated

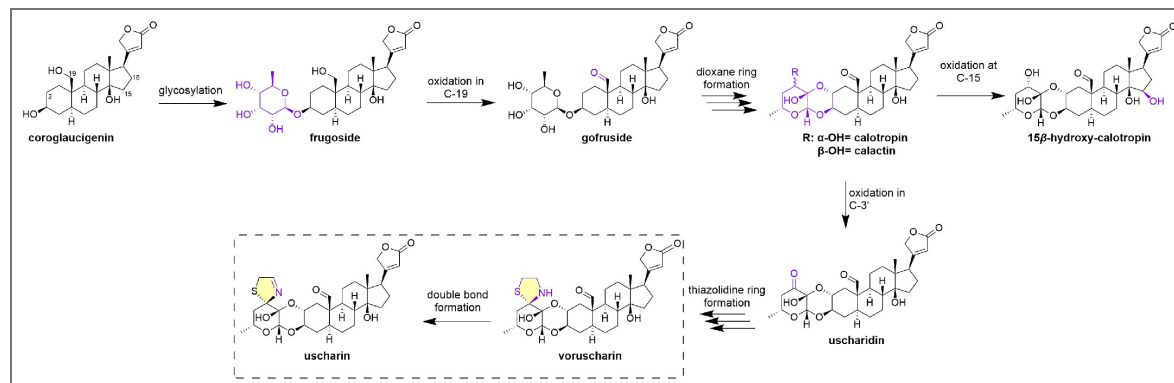
Table 1. Structural and functional features of the isolated cardenolides used in this study, comprising the dominant compounds in *Asclepias curassavica* (70% of the total leaf cardenolides).

The structural complexity values are based on dox-g (Rubiano-Buitrago et al. 2025). We also provide a non-chromatography dependent metric of polarity, WLOGP (Daina et al. 2017). IC-50 represents μM needed to inhibit the monarch Na/K-ATPase by 50% *in vitro*, data from Agrawal et al. (2021) except for 15-Hydroxy calotropin which was generated for this study (Fig. S4, Table S1). The proportion of total cardenolides in *A. curassavica* leaves is based on past work (see methods). Proportion in “real mix” was scaled up from the proportion found in leaves to sum to 1.

	15-hydroxy calotropin	Frugoside	Calactin	Uscharin	Voruscharin
Nitrogen-cardenolide	No	no	no	yes	yes
Molecular mass	548.6	536.7	532.6	587.7	589.7
Structural complexity (dox-g)	20	17	19	22	21
HPLC-DAD retention time (min.)	11.65	12.66	15.93	20.59	21.25
WLOGP	0.97	1.43	2	2.77	2.29
IC50 on Monarch enzyme (μM)	64.8	27.8	3.2	1.0	2.0
Prop. total cards in leaves	0.10	0.05	0.10	0.10	0.35
Prop. total cards in “real mix”	0.14	0.07	0.14	0.14	0.50

Figure 1. A proposed biosynthesis pathway for cardenolides of *A. curassavica* built on coroglaucigenin (Rubiano-Buitrago et al. 2025).

Modifications are indicated by purple highlighting and each arrow indicates a hypothesized step; multiple arrows indicate multiple concerted reactions without displaying all the intermediates. All compounds, except the genin, are known to occur in *A. curassavica*, however gofruside is rarely found in high quantities in the foliage (Roy et al. 2005, Rubiano-Buitrago et al. 2022).



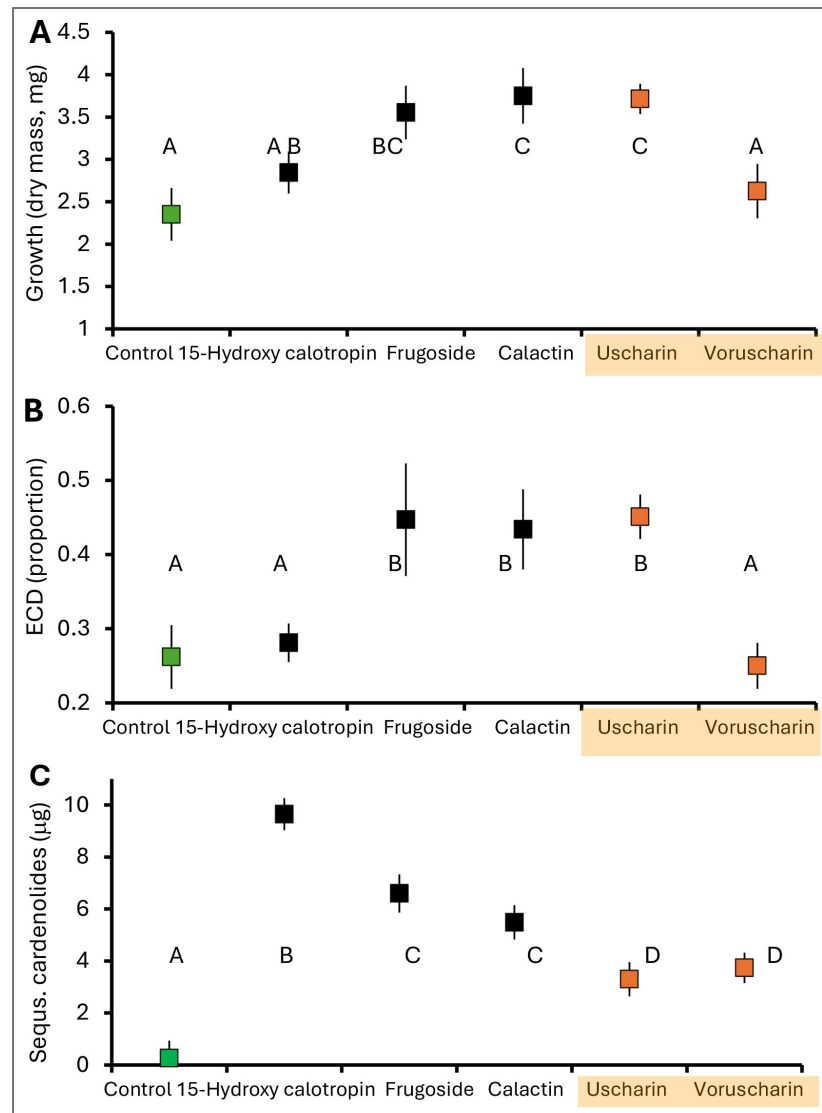


Figure 2. Monarch caterpillars are impacted by feeding on *A. incarnata* leaves painted with isolated cardenolides from *Asclepias curassavica*.

A) growth after nine days (most caterpillars in the 3rd instar), B) efficiency of conversion of digested matter, and C) total cardenolides sequestered. Shown are means $\pm$ SEs and different letters indicate a significant difference ($p < 0.05$, Fisher's LSD). The green symbol has no cardenolides added (*A. incarnata*) and the two orange symbols represent cardenolides with N,S-ring moiety.

food converted to biomass) was not impacted by uscharin; rather, the cardenolides that stimulated feeding and growth also increased conversion efficiency (up to >40%, $F_{1,85}=5.136$ $p<0.001$, Fig. 2B [↗](#)), while monarch caterpillars feeding on voruscharin and 15-Hydroxy calotropin (which showed lower growth) maintained low ECDs ($\approx 25\%$), equivalent to that of controls (Fig. 2B [↗](#)).

Effects of single cardenolides: sequestration

Our treatments with applied cardenolides to *A. incarnata* well-approximated the amount of cardenolides in leaves and sequestered by monarchs when feeding on *A. curassavica* (Fig. S2 [↗](#)). For each of the three cardenolides lacking a nitrogen moiety (15-Hydroxy calotropin, frugoside, calactin), monarchs sequestered the compound intact (>90% of the stored cardenolides were those applied to the leaves); conversely, uscharin and voruscharin were both converted to calotropin (>56% of the stored cardenolides) and calactin (>20% of the stored cardenolides). Neither uscharin nor voruscharin was detected in caterpillar bodies and only uscharin was detected in frass (comprising 20% of excreted cardenolides for caterpillars fed uscharin).

When feeding on the five compounds individually, monarch caterpillars sequestered the least total toxins when eating the two N,S cardenolides, which were converted to other compounds (Fig. 2C [↗](#)). Overall, caterpillars ingested between 63 and 83 μg of cardenolide, of which we were able to account for $\approx 15\%$ in the body and frass (Fig. 3 [↗](#)); unaccounted cardenolides may be degraded, which has been previously suggested (Seiber et al. 1980 [↗](#)). Critically, when feeding on uscharin or voruscharin, caterpillars had the lowest sequestration efficiency and excreted the least cardenolides compared to other treatments (contrast of non-N,S vs. N,S-cardenolides for proportion sequestered $F_{1,73}=64.490$, $p<0.001$; proportion excreted $F_{1,73}=76.00$, $p<0.001$, Fig. 3 [↗](#)). Thus, the N,S-cardenolides are not sequestered intact, and after conversion to non-N,S-cardenolides they are sequestered relatively poorly as well as excreted, indicating that more of these compounds are missing, and possibly degraded. Among the other cardenolides, substantial variation exists, with the monarchs sequestering twice as much 15-Hydroxy calotropin as they excreted, and the reverse for calactin, excreting twice as much as they sequestered (Fig. 3 [↗](#)). In summary, sequestration was consistently worse for the N,S-compounds, likely because detoxification is required; however the strongly disparate effects on feeding and growth were not predicted by the presence of the N,S-moiety.

Effects of mixtures

The effects of cardenolide mixtures were assessed first by contrasting the average of all single cardenolide treatments against the two mixtures for each response variable. In short, all feeding, growth, and sequestration parameters of monarch caterpillars were reduced by feeding on cardenolide mixtures (Fig. 4). For growth, the negative effect of mixtures vs. single compounds was driven primarily by reduced feeding (i.e., deterrence); in other words, when we include the amount dry mass tissue consumed in the analysis, the effect of mixtures is no longer significant in Fig. 4 [↗](#) ($p=0.173$).

Perhaps most interestingly, the efficiency of sequestration of ingested cardenolides declined from 7.5% singly to 5% in mixtures. This result controls for feeding, as we used the estimated micrograms of cardenolides ingested to calculate the efficiency. For this metric, we constructed a specific contrast for the expected values based on the proportions used in the equimolar and in the realistic mixtures (Table 1 [↗](#)). The sequestration efficiency of the equimolar mixture was reduced by 18% compared to the average of the single compounds (contrast: $p=0.036$) and was reduced by 42% in the realistic mixture compared to the expected values based on relative abundances of the five compounds ($p=0.002$).

Overall, even if not statistically significant, we note that monarch caterpillars raised on the realistic *A. curassavica* cardenolide mixture had the lowest growth and feeding, and the lowest cardenolide sequestration, excretion and sequestration efficiency compared to all single treatments, supporting a hypothesis of adaptive deployment of cardenolide mixtures by *A. curassavica* (Fig. S3 [↗](#)).

Figure 3. Monarch caterpillars differentially sequester and excrete cardenolides when eating leaves painted with isolated compounds from *Asclepias curassavica*.

Shown are means $\pm$ SEs of the proportion of total cardenolides ingested that are sequestered or excreted (bars, left axis); shown also is the amount of cardenolide ingested (green dots, right axis). 15-Hydroxy calotropin, frugoside, and calactin are sequestered intact, while uscharin and voruscharin are stored after conversion to calotropin and calactin. Orange shading indicates cardenolides with N,S-ring moiety. Data on cardenolide concentrations sequestered and excreted on a mass basis are given in Fig. S2.

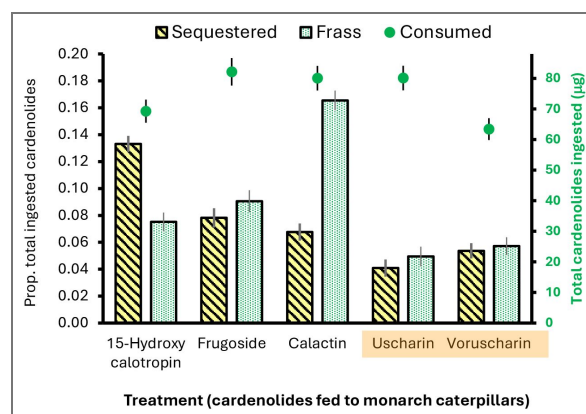
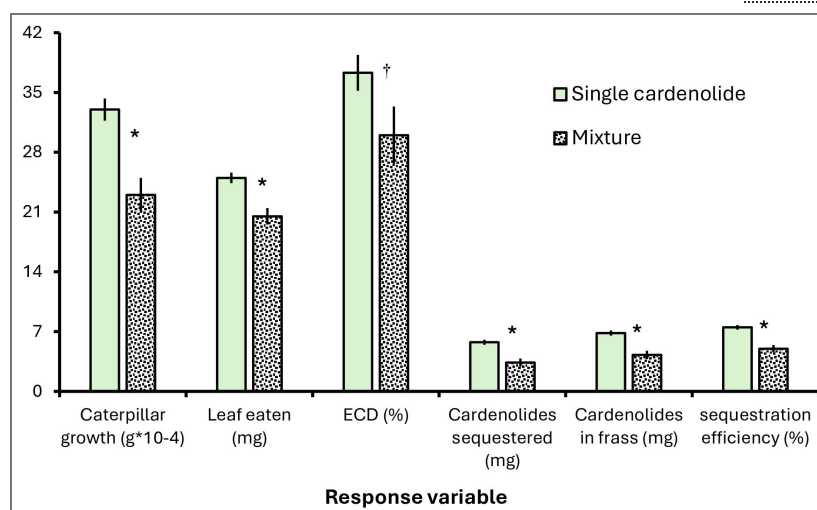


Figure 4. Monarch caterpillars differentially grow and sequester cardenolides when feeding on single isolated compounds from *Asclepias curassavica* compared to mixtures.

Shown are means $\pm$ SEs for several response variables with the units shown below the X axis. Sequestration efficiency was calculated by the mg of cardenolide ingested / mg cardenolide sequestered. Single effects are the average of the five compounds administered individually, whereas the mixture is the average of an equal mixture treatment and realistic mixture treatment. In all cases, total cardenolide concentrations were presented to caterpillars on an equimolar basis. Significance ($p < 0.01$) is shown by *, while † indicates $p = 0.07$. All treatment means are provided in Fig. S3.



Discussion

Effects of individual compounds

As is the case for most plant species, milkweeds contain complex mixtures of secondary metabolites implicated in defense against herbivores. With the advent of improved chromatographic techniques (Pandohee et al. 2023 [↗](#)), more is being discovered about such mixtures, especially on the diversity of compounds within chemical classes. With >50 cardenolide structures reported from *A. curassavica* (Roy et al. 2005 [↗](#), Li et al. 2009 [↗](#), Zhang et al. 2014 [↗](#), Rubiano-Buitrago et al. 2022 [↗](#)), we have been investigating the diversity of these compounds and their effects. These toxins are spread across plant organs, and many compounds occur in very small amounts; here we have focused on five dominant cardenolides from foliar tissues. Despite the fact that N,S-cardenolides have been known for decades, their importance in toxicity was discovered only recently (Hesse et al. 1950 [↗](#), Reichstein et al. 1968 [↗](#), Seiber et al. 1978 [↗](#), Benson et al. 1979 [↗](#), Züst et al. 2019 [↗](#)). Five years ago we reported that uscharin and voruscharin were not sequestered intact, but converted to less toxic calotropin and calactin, and imposed a cost of sequestration for monarch caterpillars (Agrawal et al. 2021 [↗](#)). Perhaps most surprisingly, in *in vitro* assays, these same compounds had relatively weak effects on a sensitive Na/K-ATPase enzyme, suggesting that they are targeted at the adapted specialists' more resistant pump (Rubiano-Buitrago et al. 2025 [↗](#)). These findings are reminiscent of classic work on coumarins, where linear furanocoumarins are tolerated by specialist black swallowtail caterpillars, but more unusual angular furanocoumarins, which are not toxic to many organisms, specifically negatively impact black swallowtails (Berenbaum and Feeny 1981 [↗](#)). Later it was found that these two types of furanocoumarins synergize to impact the specialist (Berenbaum and Zangerl 1996 [↗](#)). Despite this well-characterized example, we still know relatively little about the structural diversity of chemical defenses within a compound class and their roles against most specialist herbivores (Marquis and Koptur 2022 [↗](#)).

In the current study we confirm that N,S-cardenolides are not sequestered by monarchs, and more demonstrate that the processing of these compounds is associated with reduced sequestration efficiency and reduced cardenolide excretion compared to closely related non-N,S-compounds. One previous study fed isolated cardenolides to monarchs and reported the effects on sequestration and excretion (Seiber et al. 1980 [↗](#)). In both studies, a substantial fraction (~80%) of cardenolides was unaccounted for, with monarchs sequestering on the order of 10% of the cardenolides they consume, smaller amounts excreted, but a large portion potentially degraded by the caterpillar. Thus, although many cardenolides are sequestered intact by monarch caterpillars, it is clear that some are detoxified, as in the case of N,S-compounds, where the removal of the N,S-ring substantially reduces toxicity (Agrawal et al. 2021 [↗](#), Agrawal et al. 2022 [↗](#)). Likewise, several di-glycosides are stored as mono-glycosides by both monarchs and the seed-feeding milkweed bug *Oncopeltus fasciatus* (Agrawal et al. 2022 [↗](#), Rubiano-Buitrago et al. 2023 [↗](#)). Finally, Seiber et al. (1980) [↗](#) reported that genins (cardenolides without any sugar moiety) were not sequestered intact, but are rather stored as more polar metabolites, as was the glycoside uscharidin, neither of which were part of the current study.

In our current work, the addition of cardenolides to a diet nearly devoid of the compounds (leaf discs of *A. incarnata*) stimulated caterpillar feeding and growth. Although at first glance this result can seem counterintuitive, plant secondary metabolites, even when effective defenses, often function as stimulants for feeding and oviposition by specialist insects (Renwick 2002 [↗](#), Schoonhoven and Van Loon 2002 [↗](#), Bernays and Singer 2005 [↗](#), Wink 2018 [↗](#)). Although the extent of feeding stimulation was not predicted by whether the cardenolides were N,S-containing compounds, we found that the most stimulatory compounds not only increased caterpillar growth, but also increased the efficiency of conversion of ingested matter (ECD).

Sequestration efficiency

Sequestration efficiency is the proportion of ingested plant toxins that are stored by herbivores. In our study, N,S-cardenolides had the lowest sequestration efficiency (Figure 4 [4](#)), presumably due to the detoxification required. Additionally, we found two-fold variation among the three related non-N,S-compounds (Figures 4 [4](#), 1 [1](#)). Remarkably, sequestration efficiency on our positive control, *A. curassavica* leaves, was the same as the average of our individual compounds (Fig. S3 [3](#)). And finally, monarch caterpillars had 100-fold higher sequestration efficiency on our negative control (*A. incarnata*). This result is consistent with evidence demonstrating the upregulation of transporters and cardenolide concentration on this extremely low cardenolide species (Jones and Agrawal 2019 [4](#), Tan et al. 2019 [4](#)).

Sequestration efficiency may be determined by food intake rate, the extent of toxin processing (detoxification or transport) needed, and existing levels of the toxin already sequestered (Jones et al. 2019 [4](#), Beran and Petschenka 2022 [4](#)). For example, fertilization of tropical milkweed resulted in monarch caterpillars growing more slowly and having lower cardenolide sequestration efficiency than when feeding on controls (Tao and Hunter 2015 [4](#)). This effect seems to have been driven by an overall negative relationship between foliar cardenolide concentrations and sequestration efficiency. In this experimental approach, across other systems (e.g., Bowers and Collinge 1992 [4](#), Lampert 2020 [4](#)), and ours, caterpillar feeding and growth rates are often correlated with sequestration efficiency. These relationships may be due to general vigor, consumption rate and exposure to the plant toxins, or detoxification (and its associated costs). Thus far, in no plant-herbivore system have these relationships been disentangled.

Phytochemical diversity effects

Mechanistically, defensive compounds may act at different stages of attack by herbivores, and may reach different tissues in the consumer, ultimately providing greater defense than any one compound. For example, defensive phytochemicals can differentially impact herbivore oviposition, larval feeding, growth, sequestration, and mating (Landolt and Phillips 1997 [4](#), Macel and Vrieling 2003 [4](#), Macel et al. 2005 [4](#), Kim and Jander 2007 [4](#), Müller et al. 2010 [4](#), Agrawal et al. 2021 [4](#)). Additionally, the interaction diversity hypothesis posits that phytochemical diversity may not necessarily have non-additive effects on any one herbivore, but different compounds may target different species of attackers, thereby creating a benefit of phytochemical diversity against diverse insect pests (Whitehead et al. 2021 [4](#), Zaman et al. 2025 [4](#)). Early on, Berenbaum et al. (1991) [4](#) not only speculated that phytochemical diversity may be defensively beneficial via different pathways, but also that structural diversity within a compound class, in their case furanocoumarins, may be driven by environmental variation; such condition dependent effects may both be caused by available nutrients (e.g., Tao and Hunter 2015 [4](#), Agrawal et al. 2024a [4](#)) as well as distinct benefits of chemical diversity in different environments (Berenbaum et al. 1991 [4](#), Agrawal et al. 2012a [4](#)).

Other benefits of phytochemical diversity from the plant's perspective can be manifold. Although “synergism” between phytochemicals is often invoked, there is little to no evidence of synergistic effects of plant defense compounds in a pharmacological sense, especially when they are in the same chemical class (see Table 1 [4](#) in López-Goldar et al. 2024 [4](#)). Because some researchers interpret “synergism” with such a biochemical mechanism, we demure from using this term, as synergism per se has not been well-studied. Nonetheless, diversity effects, here defined as greater than additive impacts of multiple compounds compared to singletons, have been reported in several studies within a phytochemical class. In classic work, angular furanocoumarins showed relatively low toxicity in isolation, but had greater than additive effects with linear furanocoumarins, both in specialists and generalist herbivores (Berenbaum and Zangerl 1996 [4](#)). For amides and imides from *Piper* spp., evidence suggests diversity effects depend on the level of specialization of herbivores, and the type of assay (growth, parasitism, final size) (reviewed in Richards et al. 2016 [4](#)). Some studies with *Jacobaea* pyrrolizidine alkaloids provide evidence for greater effects in mixture compared to single compounds, although this effect was primarily found with sub-organismal (in vitro) assays (Nuringtyas et al. 2014 [4](#)).

Two studies have measured impacts of phytochemical diversity not only on caterpillar performance and sequestration, but also on immunity to parasites. Richards et al. (2012) [found](#) that specialist *Junonia* caterpillars grew faster, had lower mortality, sequestered higher concentrations, and had reduced immune responses on artificial diets containing mixtures of two of iridoid glycosides (acubin and catalpol) compared to those containing individual compounds. In other words, this study reported a lower than expected defensive function of mixtures, but this may come with reduced immunity of caterpillars to enemies. In our studies of *A. curassavica*, we find the reverse: not only do mixtures of cardenolides reduce caterpillar performance and sequestration, but monarchs have greater immunity against the protozoan parasite. More work will be needed to address the causes of these effects and trade-offs between performance, sequestration and immunity (Hoogshagen et al. 2023 [found](#)).

In previous research (López-Goldar et al. 2024 [found](#)), we demonstrated effects of phytochemical diversity using three abundant cardenolides from common milkweed. These effects were most evident in terms of a dominance effect of highly inhibitive compounds at the target site (Na/K-ATPase), as well as physiological effects of phytochemical diversity in assays with transgenic flies that were tolerant of cardenolides but lack other forms of specialization. When diverse cardenolides were briefly fed to monarch caterpillars, the main cost of chemical diversity was through an enhanced cost of sequestration compared to when monarchs were feeding on single compounds. In the current work, diversity of cardenolides imposed multiple limitations for caterpillar development, spanning growth and sequestration efficiency.

Empirical evidence from several studies (Berenbaum et al. 1991 [found](#), López-Goldar et al. 2024 [found](#), Zaman et al. 2025 [found](#)) suggests that relatively minor constituents of the natural blend of compounds in leaves may be contributing to the observed toxicity and greater effect of mixtures compared to single compounds. Even when the mixture is not more toxic than the most toxic individual compounds, mixtures are often more than additive in their effects. This distinction is not based on mechanism, and therefore we have not been using the phrase synergistic, but rather “non-additive diversity effect “ in a statistical sense. This distinction is parallel to the plant diversity literature in ecology which aims to separate multiple mechanisms of diversity effects (Petchey 2003 [found](#)). Our key point is that non-additivity in phytochemical defense may be adaptive because small amounts of highly toxic compounds may complement, synergize, or dominate the mixture, thereby providing an economic advantage in terms of potentially lower production costs with the benefits of diversity (López-Goldar et al. 2024 [found](#)).

Conclusion

In milkweeds, as in many other groups, diversification of secondary metabolites appears to occur via the chemical evolution of glycosylation, oxidation, cyclization (i.e., ring formation) and addition of complex functional groups onto core existing chemical structures (Fig. 1 [found](#)). For the group of coroglaucigenin based cardenolides, molecular mass, complexity, polarity, and toxicity progress through each biosynthetic step leading to N,S-cardenolides (Rubiano-Buitrago et al. 2025 [found](#)). Although we have only tested one herbivore species here, our past work has shown similar effectiveness of these compounds against seed-feeding specialists as well, especially in vitro (Agrawal et al. 2021 [found](#), Agrawal et al. 2022 [found](#)). Here, two N,S-cardenolides had unequal effects, with only uscharin having a stimulatory effect on feeding and growth; nonetheless, these compounds were both modified and showed poor sequestration efficiency of the resulting cardenolides. When in a realistic mixture, monarch caterpillars fed and grew the least and sequestered the least compared to caterpillars feeding on equimolar concentrations of any of the five single compounds (Fig S3 [found](#)). Thus, cardenolide mixtures presented in *A. curassavica* appear to be a compromise between reducing growth and sequestration of highly specialized monarchs and providing benefits to the monarch in fighting off enemies (Hoogshagen et al. 2023 [found](#)).

Materials and Methods

As our main goal was to compare the major cardenolides isolated from *A. curassavica* alone and in mixture, we took the approach of painting equimolar concentrations of total cardenolides in all treatments (Table 1 [↗](#)). These painting treatments were imposed on leaf discs of *A. incarnata*, a closely related species (both species within the Incarnatae clade), but with cardenolide abundance at trace levels (Agrawal et al. 2015 [↗](#)). Fully randomized within our experiment were a negative and positive control fed to caterpillars, *A. incarnata* painted with ethanol alone and *A. curassavica* leaves, respectively. Although these treatments were not included in most statistical analyses comparing the different individual cardenolides or comparing single compound and mixtures, we specifically compare these treatments to controls where appropriate (i.e., *A. incarnata* alone in the analysis of growth and *A. curassavica* in the analysis of sequestration).

Plant growth

Seeds of *Asclepias incarnata* (collected from Tompkins Co., NY) and *A. curassavica* (purchased from Everwilde Farms, Fallbrook, CA) were surface-sterilized with 10% bleach for 10 minutes, rinsed, nicked, and stratified at 4°C in petri dishes lined with moist paper towels for 7 days. Seed dishes were then moved into an incubator at 30°C for 3–4 days. Germinated seedlings were transplanted into 10 cm pots containing moistened Lambert all-purpose mix (LM-111; Riviere-Ouelle, Quebec, Canada) and placed in a growth chamber with a 14:10 day:night cycle, with a day temperature of 27°C and a night temperature of 24°C. Plants were grown for approximately 5 weeks, fertilized once at first dry down and again approximately 10 days later with 20:20:20 N:P:K (Jack's all-purpose fertilizer, JR Peters, Allentown, PA), and watered as needed.

Caterpillar sourcing and rearing

Monarch (*Danaus plexippus*) eggs were obtained from a laboratory colony in late January 2024, and were kept in the lab until hatching. Immediately upon hatching, each caterpillar was placed in a 1 oz deli cup containing one small piece of moist cotton (to prevent leaf tissue from drying) and a leaf disc (0.95 cm diameter) of *A. incarnata* or *A. curassavica*, prepared with the appropriate treatment. Twenty caterpillars were reared on each of nine treatments, consisting of eight treatments applied to *A. incarnata* leaf discs (negative control, five single compound treatments, and 2 compound mixture treatments) and one treatment applied to *A. curassavica* leaf discs (positive control), for a total of 180 caterpillars. Caterpillars were checked regularly and were given a fresh leaf disc corresponding to their assigned treatment each day or as needed to ensure they did not run out of food. Each time the caterpillar was given a new leaf disc, the previous disc was saved for leaf area determination. On the sixth day of the experiment, we increased the leaf punch diameter size to 1.27 cm, as the caterpillars' feeding increased. All caterpillars were reared in a randomized array on a lab bench, away from a window, at ambient lab temperature.

Cardenolide solution preparations

Cardenolides from *A. curassavica* foliage were available by previous isolation from aerial material (See methods in Agrawal et al. 2021 [↗](#)). All compounds were >90% absolute purity by ¹H NMR (except 15-hydroxy-calotropin which was 65% pure) and all compounds were >90% relative purity by UV data at 218 nm. Cardenolides were diluted to approximately 0.5 mM based on estimated amounts; frugoside, calactin and 15-hydroxy-calotropin in methanol, whereas uscharin and voruscharin were diluted in acetonitrile due to solubility constraints. Each compound was then quantified using an external HPLC-DAD digitoxin calibration curve (in the appropriate solvent) at 218 nm.

Both the calibration curve and compound quantification were performed on an Agilent 1100 HPLC (Santa Clara, CA, USA), using the column, specifications and gradient described below in the 'cardenolide extractions and HPLC-DAD analysis' section (and also in Petschenka et al. 2022 [↗](#)). Molar concentrations were converted to mass, based on the known molecular weight of each compound. For single compound treatments the volume of solution representing 1.2636 mg of

each compound was then pipetted into a separate tube and taken to dryness in a rotary evaporator (Labconco, Kansas City, MO). For the equimolar mixture, this same mass (1.2636 mg) was split, in equimolar fashion, between all 5 compounds, while the realistic mixture contained relative proportions of the five compounds similar to what is found in *A. curassavica* leaf tissue (see Table 1 [↗](#)). For our realistic measure, we combined knowledge from published studies [Agrawal et al. \(2021\) \[↗\]\(#\)](#) and unpublished work on the relative abundance of cardenolides in *A. curassavica*. In both cases, the total concentration of the mixtures was equivalent to that of individual compounds. We made a second set of these preparations halfway through the experiment.

For each treatment, dried cardenolide was brought up in 95% ethanol to a concentration of 0.675 mg/mL. Due to low solubility of some compounds in ethanol, these mixtures were treated as suspensions, vortexed and sonicated for homogeneity before each use, and stored at 4°C in between uses. Just before each feeding, experimental leaf discs (0.95 cm diameter) were prepared by taking leaf punches directly from fresh leaves in the growth chamber, and then using a pipet to apply 6 µL of the appropriate suspension evenly to the top surface each leaf punch. Punches were collected from each plant on one day only, to prevent any potential effects of induction. Punches were dried in a fume hood and then a second aliquot of 6 µL was applied, for 12 µL total, to bring the concentration of each experimental cardenolide application to 3 mg/g dry leaf tissue, which approximates the total cardenolide concentration of *A. curassavica* leaves based on our previous work and the literature. *A. incarnata* and *A. curassavica* control leaf discs received 12 µL of 95% ethanol alone, in the same manner as above. *A. incarnata* leaves were chosen as the substrate because they are low in cardenolides (mean of 0.0028 mg/g dry mass based on five samples used in this study), while *A. curassavica* leaf discs for this experiment averaged 3.8 mg/g dry leaf tissue. From Day 6 on, when larger punches were used, the amount of cardenolide mixture applied to each punch was adjusted to keep the overall cardenolide concentration at 3 mg/g dry leaf tissue.

Data collection

On each day that leaf discs were changed, notes were taken on caterpillar mortality or molting, and old leaf discs were taped to a piece of paper for estimation of remaining leaf area, using LeafByte ([Getman-Pickering et al. 2020 \[↗\]\(#\)](#)). Remaining leaf area was subtracted from known starting punch area to estimate total leaf area per day, and these amounts were summed for the 10 days of the experiment. On Day 10, the final day of the experiment, caterpillar instar (2 or 3) was recorded and each caterpillar was weighed, and stored at -80°C for subsequent cardenolide analysis. All loose frass was collected from each caterpillar (not counting early instar frass stuck to the cotton piece) and stored at -80°C as well. While we started with 180 caterpillars, 21 caterpillars died during the experiment (final N=141). Caterpillars and frass were freeze-dried prior to extraction.

Cardenolide extraction and HPLC-DAD analysis

Freeze dried caterpillar and frass samples were weighed and ground to a powder in a Mixer Mill (Retsch, Haan, Germany) in screw cap tubes (Sarstedt, Nümbrecht, Germany) using 1 stainless steel bead per sample. Samples were extracted as per [Petschenka et al. 2022 \[↗\]\(#\)](#), using 1mL methanol spiked with 20 ug hydrocortisone standard per sample, and extracted using zirconia-silica beads and a FastPrep-24 (MP Biomedicals, Santa Ana, CA) twice for 45sec at 6.5 m/s. Samples were centrifuged at 20,817g for 12 min, and each supernatant was transferred to a clean 2 mL tube (caterpillars) or a 1.4 mL racked tube (frass). Caterpillar samples were dried and then brought up in 250 µL methanol, and defatted using 750 µL hexanes, and then taken to dryness, along with the frass samples, in a rotary evaporator (Labconco, Kansas City, MO) at 35°C. Residues were reconstituted in 200 µL methanol, filtered through a hydrophobic filter plate (Millipore, Burlington, MA), and sealed for chromatography analysis.

Fifteen microliters of extract were injected into an Agilent 1100 series HPLC-DAD, and compounds were separated on a Gemini C18 reversed phase column (3µm, 150 x 4.6 mm, Phenomenex, Torrance, CA). Cardenolides were eluted on a constant flow of 0.7 ml/min with an acetonitrile-

water gradient as follows: 0-2 min 16% acetonitrile, 25 min 70% acetonitrile; 30-40 min 95% acetonitrile, followed by a 10min post-run in 16% ACN. Peaks were recorded by diode array at 210, 218, 280, 320 and 360 nm, with hydrocortisone as the standard. Using the 218 nm detection data, peaks with symmetrical absorption maxima between 217 and 222 nm were recorded as cardenolides (Petschenka et al. 2022 [↗](#)).

Concentrations were calculated using peak areas of hydrocortisone (and a conversion factor to expected peak area of digitoxin), and total cardenolide concentration was calculated as the sum of all individual cardenolide peaks.

Statistical approach

To understand impacts our individual cardenolide treatments on growth of the caterpillars, we took two approaches. First, on a dry mass basis, we analyzed caterpillar mass as a measure of growth rate, leaf mass ingested, and mass of frass excreted. The effect of cardenolide treatments on the efficiency of conversion of digested matter (EDC) (Waldbauer 1968 [↗](#)), was calculated as caterpillar growth/(ingested leaves – frass). One-way ANOVA was used to test the differences between the five cardenolide treatments. In addition, we built models with cardenolide treatment, ingested matter, and their interaction as predictors. These analyses address the extent that feeding impacts growth and digestion differentially among treatments. A parallel set of analyses was conducted on cardenolide sequestration and excretion. Specific contrasts assessed differences between cardenolide types.

To test for a phytochemical diversity effect, our single cardenolide treatments were contrasted to our mixture treatments. Here we employed fixed effect nested ANOVAs with the five individual cardenolide treatments nested within a “single” effect and our two mixtures nested within a “mixture” effect. In this way, response variable was tested as a function of treatment group (single vs. mixed), with the specific treatments nested within each of these two groups. This approach is functionally equivalent to a contrast of the five single vs. two mixture treatments, and but also assesses variation among the nested effects. All analyses were conducted using JMP Pro Ver. 16.

Estimation of ic-50 of 15-hydroxy calotropin

The compound identity was verified by NMR (Table S1 [↗](#)). Inhibition of both a cardenolide-adapted (monarch butterfly) and unadapted (porcine) Na/K-ATPase by 15-hydroxy calotropin was tested using an in vitro enzyme assay described in Petschenka et al. 2022 [↗](#). Monarch nervous tissues were dissected, homogenized in Millipore water and freeze-dried, while the porcine ATPase was obtained commercially (Millipore Sigma, Burlington, MA). A stock solution of our 1mM 15-hydroxy calotropin prep was prepared by HPLC quantification (as described above in the ‘cardenolide solution preparation’) in 20% DMSO in Millipore water. Serial dilutions were prepared with 20% DMSO - one at 0.5 mM, then 4 additional 1/10 dilutions, to prepare a 6-point inhibition curve, and each curve was run in triplicate for each enzyme (monarch and porcine), alongside ouabain. Assays were run, as described in Rubiano Buitrago et al. 2025 [↗](#), but using 0.75 monarch brain tissue per mL Millipore water instead of *Oncopeltus* nervous tissue. Absorbances were determined spectrophotometrically at 700 nm, corrected by background values, and dose-response curves were fit using a nonlinear mixed effects model with a four-parameter logistic function in R version 4.5.1, in order to determine the concentration of 15-hydroxy calotropin that inhibits the activity of each ATPase by 50% (Rubiano Buitrago et al. 2025 [↗](#), Petschenka et al. 2022 [↗](#)).

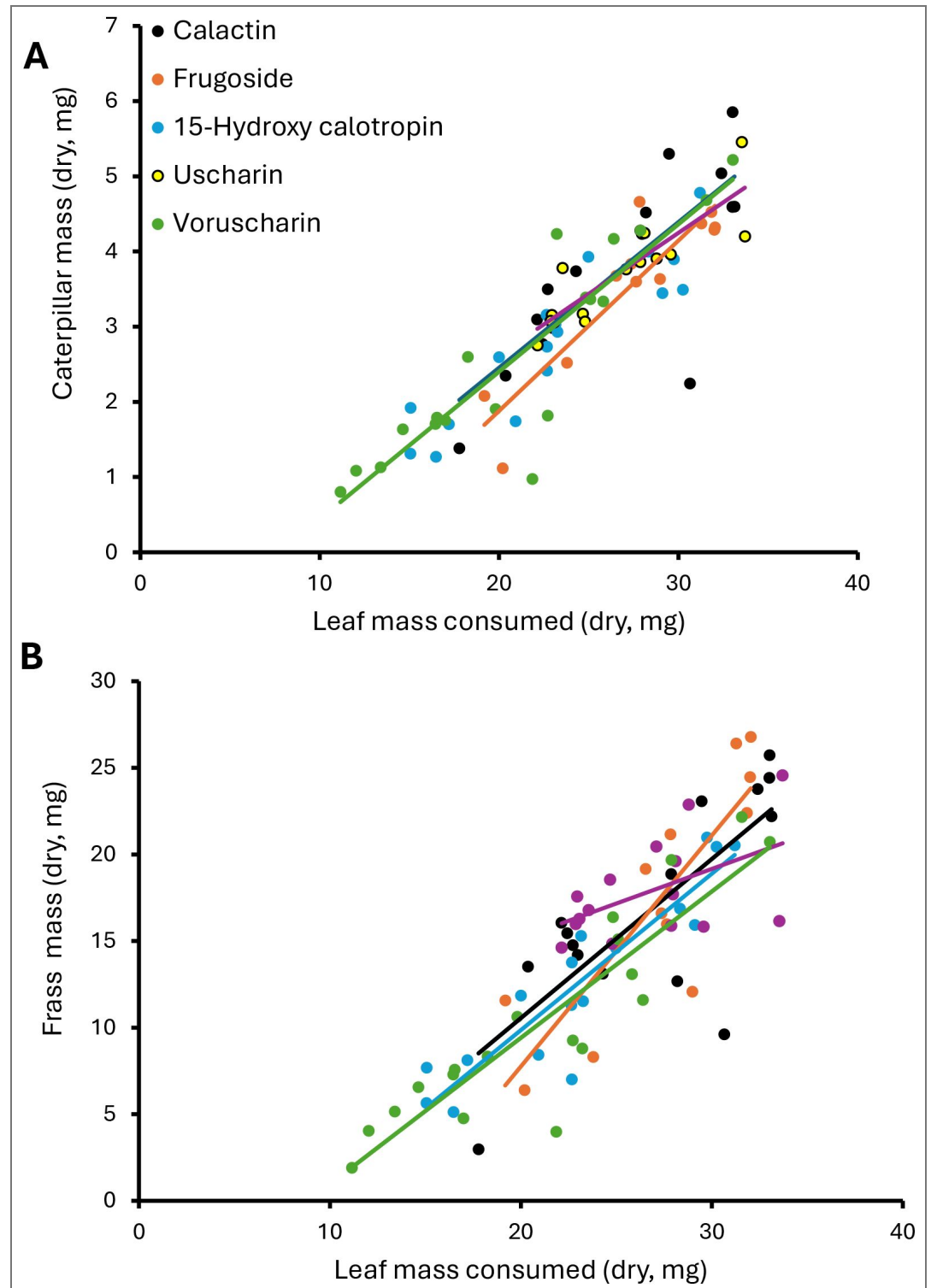


Figure S1. Monarch caterpillars are impacted by feeding on leaf discs painted with isolated cardenolides from *Asclepias curassavica*. A) growth is dependent on leaf mass consumed, with the same slope for all treatments, and B) excretion is dependent on leaf mass consumed, but with different slopes depending on treatments.

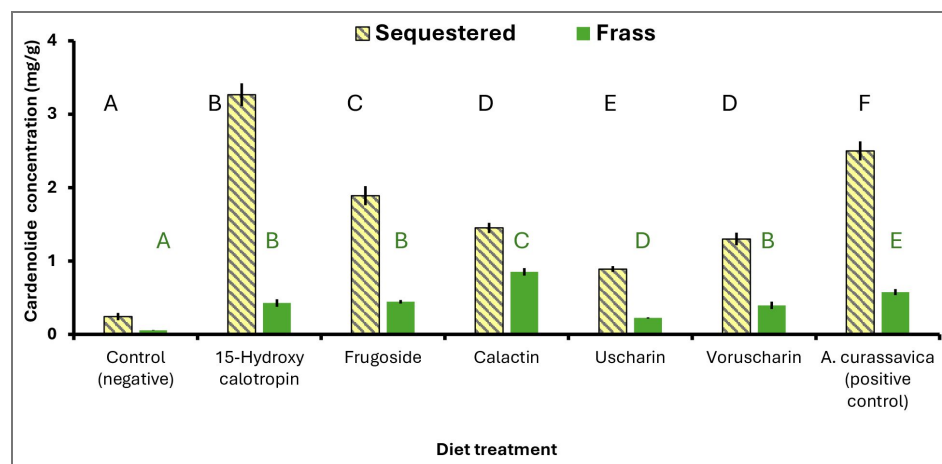


Figure S2. Monarch caterpillars differentially sequester ($F_{6,102}=102.22$, $p<0.001$) and excrete ($F_{6,100}=37.12$, $p<0.001$) cardenolides when feeding on leaf discs painted with isolated compounds from *Asclepias curassavica*.

Shown are means $\pm$ SEs of cardenolide concentration (on a per body mass basis) and different letters indicate a significant difference ($p<0.05$, Fisher's LSD, letters are only comparable within a tissue type). Orange shading indicates nitrogen-containing cardenolides. Data on absolute cardenolides sequestered are given in the main text. The negative control is *A. incarnata*, the very low cardenolide plant species that was used to paint on individual cardenolides. The positive control was *A. curassavica* leaves, the species from which cardenolides were isolated what we were simulating in our experimental treatments.

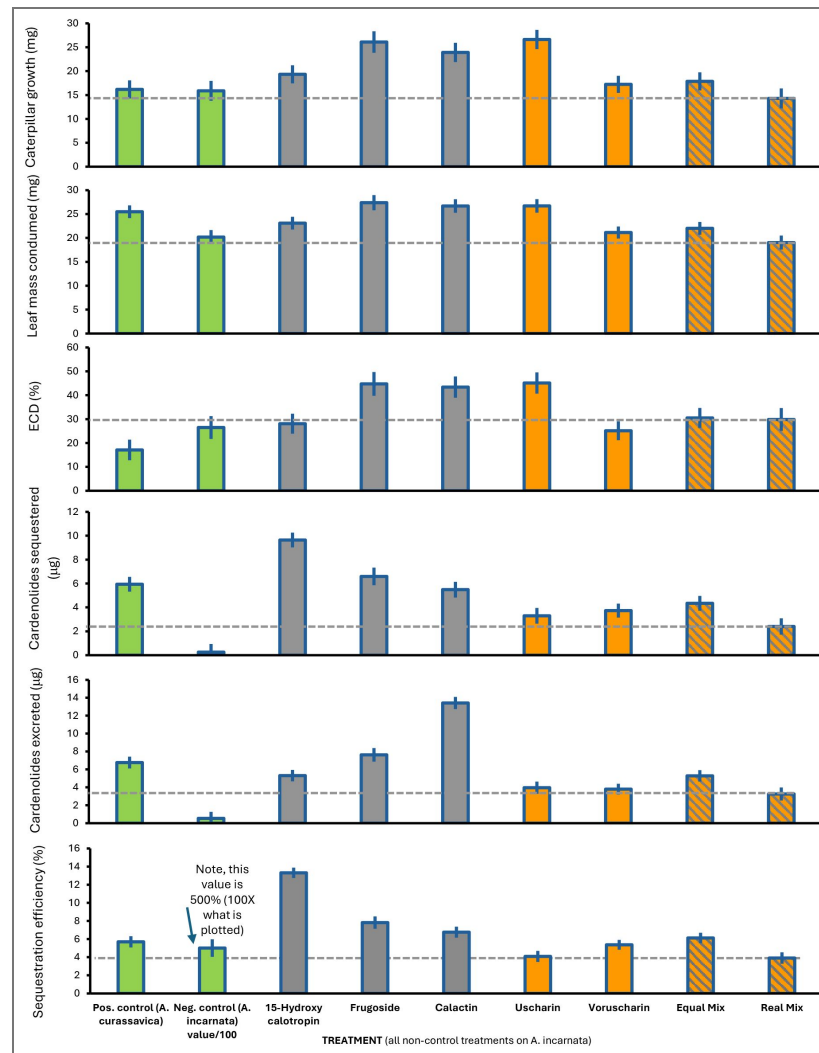


Figure S3. Mean $\pm$ SE of nine treatments growing monarch caterpillars on two controls (green: positive on *A. curassavica* and negative on *A. incarnata*), three non-N,S-cardenolides (gray), two N,S-cardenolides (orange), and 2 types of mixtures (striped).

Dashed line indicates the response of the realistic mixture, that which approximates the composition of cardenolides in *A. curassavica*. For all non-control treatments the total cardenolides applied to leaf discs was of equal concentration.

Figure S4. Inhibition curves estimated from six concentrations of 15-hydroxy calotropin on a sensitive (porcine) and adapted (monarch) sodium-potassium ATPase in vitro.

Shown are Means $\pm$ SEs.

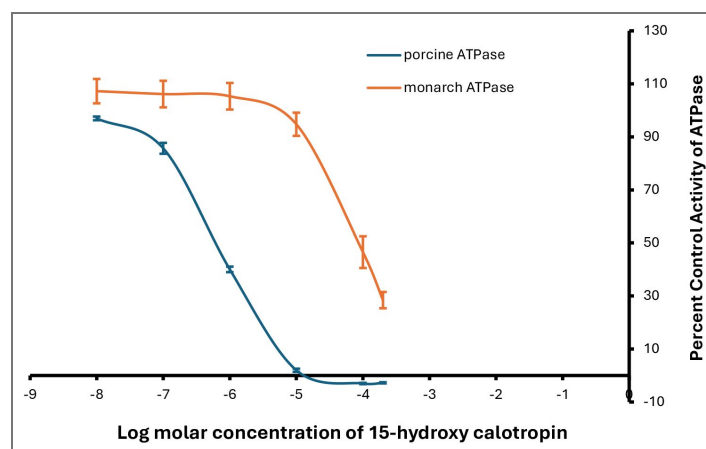


Table S1. 1D NMR assignment of key positions for the identification of 15 β -hydroxy-calotropin.

Chemical shifts with an asterisk correspond to values in CDCl₃, whereas non-designated ones correspond to values in CD₃OD. Data was compared with published records (El-Askary et al. 1993 [DOI](#), Rubiano-Buitrago et al. 2022 [DOI](#)).

Position	δ_H (J in Hz)	δ_C
1	1.12 (t,12)	36.5
	2.46 (dd,12,4)	
2	3.92 (m)	73.1
3	3.88 (m)	69.8
15	4.51 (dd,9,7)	72.9
	4.50 (t,7.6)*	
16	1.67(m)	37.6
	2.62 (m)	
17	2.7 (dd,10,4.7)	49.3
18	0.84 (s)	16.5
19	10.07 (s)	208
1'	4.44 (s)	97
2'		91.1
3'	3.58 (dd,12,4.9)	73.7
4'	1.54 (m)*	38.5*
	1.85 (m)*	
5'	3.64 (m)*	67.7*
6'	1.23 (d,6.4)	20.9

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Additional information

Author contributions

AAA conceived the study, AAA & APH designed the research, APH carried out the experiments, data collection, and sample analyses, PR-B assisted with caterpillar rearing, compound identification by NMR, and proposing the biosynthetic pathway in Fig. 1, AAA conducted the statistical analyses and wrote the first draft of the manuscript, AAA, APH, and PR-B contributed to revisions.

Author ORCID iDs

Anurag A Agrawal: <http://orcid.org/0000-0003-0095-1220>

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Peer reviews

Reviewer #1 (Public review):

Summary:

In the ecological interactions between wild plants and specialized herbivorous insects, structural innovation-based diversification of secondary metabolites often occurs. In this study, Agrawal et al. utilized two milkweed species (*Asclepias curassavica* and *Asclepias incarnata*) and the specialist Monarch butterfly (*Danaus plexippus*) as a model system to

investigate the effects of two N,S-cardenolides - formed through structural diversification and innovation in *A. curassavica*-on the growth, feeding, and chemical sequestration of *D. plexippus*, compared to other conventional cardenolides. Additionally, the study examined how cardenolide diversification resulting from the formation of N,S-cardenolides influences the growth and sequestration of *D. plexippus*. On this basis, the research elucidates the ecophysiological impact of toxin diversity in wild plants on the detoxification and transport mechanisms of highly adapted herbivores.

Strengths:

The study is characterized by the use of milkweed plants and the specialist Monarch butterfly, which represent a well-established model in chemical ecology research. On one hand, these two organisms have undergone extensive co-evolutionary interactions; on the other hand, the butterfly has developed a remarkable capacity for toxin sequestration. The authors, building upon their substantial prior research in this field and earlier observations of structural evolutionary innovation in cardenolides in *A. curassavica*, proposed two novel ecological hypotheses. While experimentally validating these hypotheses, they introduced the intriguing concept of a "non-additive diversity effect" of trace plant secondary metabolites when mixed, contrasting with traditional synergistic perspectives, in their impact on herbivores.

Weaknesses:

The manuscript has two main weaknesses. First, as a study reliant on the control of compound concentrations, the authors did not provide sufficient or persuasive justification for their selection of the natural proportions (and concentrations) of cardenolides. The ratios of these compounds likely vary significantly across different environmental conditions, developmental stages, pre- and post-herbivory, and different plant tissues. The ecological relevance of the "natural proportions" emphasized by the authors remains questionable. Furthermore, the same compound may even exert different effects on herbivorous insects at different concentrations. The authors should address this issue in detail within the Introduction, Methods, or Discussion sections.

Second, the study was conducted using leaf discs in an in vitro setting, which may not accurately reflect the responses of Monarch butterflies on living plants. This limitation undermines the foundation for the novel ecological theory proposed by the authors. If the observed phenomena could be validated using specifically engineered plant lines-such as those created through gene editing, knockdown, or overexpression of key enzymes involved in the synthesis of specific N,S-cardenolides - the findings would be substantially more compelling.

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Reviewer #2 (Public review):

This study examined the effects of several cardenolides, including N,S-ring containing variants, on sequestration and performance metrics in monarch larvae. The authors confirm that some cardenolides, which are toxic to non-adapted herbivores, are sequestered by monarchs and enhance performance. Interestingly, N,S-ring-containing cardenolides did not have the same effects and were poorly sequestered, with minimal recovery in frass, suggesting an alternate detoxification or metabolic strategy. These N,S-containing compounds are also known to be less potent defences against non-adapted herbivores. The authors further report that mixtures of cardenolides reduce herbivore performance and sequestration compared to single compounds, highlighting the important role of phytochemical diversity in shaping plant-herbivore interactions.

Overall, this study is clearly written, well-conducted and has the potential to make a valuable contribution to the field. However, I have one major concern regarding the interpretations of the mixture results. From what I understand of the methods, all tested mixtures contain all five compounds. As such, it is not possible to determine whether reduced performance and sequestration result from the complete mixture or from the presence of a single compound, such as voruscharin for performance and uscharin for sequestration. For instance, if all compounds except voruscharin (or uscharin) were combined, would the same pattern emerge? I suspect not, since the effects of the individual N,S-containing compounds alone are generally similar to those of the full mixture (Figure S3). By taking the average of all single compounds, the individual effects of the N,S-containing ones are being inflated by the non-N,S-containing ones (in the main text, Figure 4). In the mix, of course, they are not being 'diluted', as they are always present. This interpretation is further supported by the fact that in the equimolar mix, the relative proportion of voruscharin decreases (from 50% in the 'real mix'), and the target measurements of performance and sequestration tend to increase in the equimolar mix compared to the real mix.

Despite this issue, the discussion of mixtures in the context of plant defence against both adapted and non-adapted herbivores is fascinating and convincing. The rationale that mixtures may serve as a chemical tool-kit that targets different sets of herbivores is compelling. The non-N,S cardenolides are effective against non-adapted herbivores and the N,S-containing cardenolides are effective against adapted herbivores. However, the current experiments focus exclusively on an adapted species. It would be especially interesting to test whether such mixtures reduce overall herbivory when both adapted and non-adapted species are present.

It remains possible that mixtures, even in the absence of voruscharin or uscharin, genuinely reduce sequestration or performance; however, this would need to be tested directly to address the abovementioned concern.

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Author response:

Thanks for these insightful reviews and your summary assessment. We certainly agree that ours was a laboratory study with a single specialized insect, and both mixtures types had all five compounds (controlling for total toxin concentration). Thus, our conclusion that combined effects of naturally occurring toxins (within the cardenolide class) have non-additive effects for the specialized sequestering monarch are constrained by our experimental conditions. In our assay we used two mixture types, equimolar and "natural" proportions. We acknowledge that the natural proportions will vary with plant age, damage history, etc. of the host plant, *Asclepias curassavica*. Our proportions were based on growing the plants a few different times under variable conditions. Although we did not conduct these experiments on non-adapted insects, we discuss a related experiment that was conducted with wild-type and genetically engineered *Drosophila* (Lopez-Goldar et al. 2024, PNAS). In sum, we appreciate the reviewers' comments.

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