Both human and nonhuman organisms are sensitive to statistical regularities in sensory inputs that support functions including communication, visual processing, and sequence learning. One of the issues faced by comparative research in this field is the lack of a comprehensive theory to explain the relevance of statistical learning across distinct ecological niches. In the current review we interpret cross-species research on statistical learning based on the perceptual and cognitive mechanisms that characterize the human and nonhuman models under investigation. Considering statistical learning as an essential part of the cognitive architecture of an animal will help to uncover the potential ecological functions of this powerful learning process.

**Finding Structures in the Sensory World**

A central problem in the study of cognition and development concerns the ways in which organisms compute sensory inputs and discover patterns in the environment. Statistical learning (see Glossary) has been proposed as a key mechanism for extracting regularities distributed across sensory modalities and cognitive domains, and as a process that constitutes the foundation of further abilities ([1–3] for recent reviews). Over the past 20 years a substantial body of research has explored statistical learning across myriad domains in both human and nonhuman learners.

In the present review we identify constraints on statistical learning: differences in the amount of statistical information acquired or types of computations performed over a structured input by a given organism. We examine similarities and differences between species, and interpret statistical learning findings based on what is known about domain-general mechanisms possessed by the species in question.

Researchers in the field of human statistical learning consider these mechanisms to be an essential component of our perceptual and cognitive systems [3]. In humans, statistical learning is constrained in the way in which it operates across modalities and domains [1,4,5]. In addition, data from human infants suggest that statistical learning abilities may emerge at distinct points in development of different sensory modalities. This trajectory is likely explained, at least in part, by the ways in which the perceptual and cognitive skills of infants develop as a function of age and experience.

In nonhuman organisms, it is still unknown how statistical learning relates to other perceptual and cognitive capacities and to ecological niches. Studies have largely focused on the types of statistical computations performed by each species, and the extent to which such computations mirror those performed by humans. Less attention has been directed towards the ways in which statistical learning is constrained in nonhuman animals. As in prior work with humans, our goal is to examine statistical learning abilities through the lens of domain-general abilities that differentiate species.
Who Can Do What?
Statistical learning facilitates the detection of sequential patterns in visual, auditory, and tactile information, as well as of spatial patterns in the visual domain [6–27]. In humans, and some songbird species, statistical learning also supports communicative functions. This process has been implicated in numerous aspects of early language development, including discovering word boundaries, prosodic and phonotactic patterns, syntactic structures, and label–object mappings [28–36]. In a similar vein, vocal learning in songbirds may involve statistical learning processes. For example, juvenile Bengalese and zebra finches learn their own songs by tracking the probability distribution of syllables within songs sung by adult tutors [37–39].

We have chosen to focus this review on three specific statistical learning abilities: tracking sequential statistics, generalizing sequential patterns, and acquiring simple syntactic structures. These three abilities are particularly relevant because they have been studied using comparable behavioral methods across species. Note that there is also a large literature on testing human adults. For the purposes of the current comparative review, however, we focus primarily on human infant studies because the methods are more comparable to those used with nonhuman animals (Box 1).

Tracking Sequential Patterns

Human neonates detect frequencies of co-occurrence of streams of syllables [40] and shapes [41]. The youngest ages for which there is published evidence showing detection of transitional probabilities is 4.5 months [42] for visual materials and 7 months for auditory materials [43]. Detecting transitional probabilities is more complex than detecting co-occurrence frequencies because transitional probabilities entail predictive relations among items, whereas frequencies involve simple co-occurrence of elements. Given the studies published to date, it is unclear whether the detection of sequential statistics emerges earlier than is reported in the literature.

Sensitivity to sequential statistics has also been demonstrated in nonhuman species. Zebra finches (Taeniopygia guttata) track transitional probabilities to encode sequences of song syllables [44]. When learning songs from adult tutors, juvenile Bengalese finches (Lonchura striata domestica) select chunks of notes with greater internal transitional probabilities over groups of notes spanning chunk boundaries [37]. Following mere exposure without reward, newborn domestic chicks (Gallus gallus) detect patterns in streams of visual elements [13]. Although it remains unclear exactly which statistics chicks are computing (e.g., transitional probabilities or co-occurrence frequencies), this species does appear to be tuned to distributional information in its postnatal environment.

Among mammals, statistical learning from linguistic materials has been demonstrated in rats (Rattus norvegicus), who utilize frequencies of co-occurrence rather than transitional probabilities to track syllables in speech streams [8]. Cotton-top tamarins (Sanguinus oedipus) show similar capacities; however, it is unknown whether tamarins detect co-occurrence frequencies or transitional probabilities between syllables [45].

Generalizing Sequential Patterns

The sequential learning tasks described in the previous section require learners to track specific sequences of elements. Other experimental paradigms assess whether learners can abstract beyond specific sequences, thus requiring generalization. Human neonates generalize the structure of triplets of syllables arranged in an ABB pattern (e.g., ga-ti-ti, we-fo-fo, la-gu-gu), discriminating novel ABB sequences from random sequences of the same syllables [46]. A few
subgroups such as noun-phrases and verb-phrases. Subphrases can be nested into other subphrases to confer hierarchical organization to a sentence. Which word pertains to which category, and the position of subphrases within a sentence, is determined by statistical dependencies (an example of rudimentary phrase structures in the literature: A^B^C^D). Statistical learning: learning mechanism enabling detection of regularity (e.g., co-occurrence frequencies, transitional probabilities, nonadjacent dependencies, etc.) in sensory inputs. Transitional probability: a form of conditional probability defined by the formula: probability of B|A = frequency of AB/frequency of A. This conditional statistic is sensitive to the order with which one item predicts the following one within a pair (AB).

Box 1. Behavioral Procedures

(i) Habituation

Infants are exposed to structured visual or auditory sequences until they reach a habituation criterion (diminished looking). Each test trial consists of presentation of a single sequence that is either consistent or inconsistent with the habituation materials. Discrimination is measured as a function of looking time to the consistent versus inconsistent test items.

(ii) Head-Turn Preference

Infants are exposed to sound sequences for a few minutes in the presence of a neutral visual stimulus. The test items consist of sequences that are either consistent or inconsistent with the exposure materials (e.g., words vs non-words, the same syllables in novel orders). Infants control the duration of each test trial via a head turn in the direction of the auditory stimulus. Discrimination is measured as in habituation. In cotton-top tamarins the orienting response towards the speaker playing the stimuli serves as measure of discrimination.

(iii) Go/No-Go

Animals are first trained to discriminate a sound associated with the Go response (e.g., pecking on a sensor) from a sound associated with the No-Go response (no pecking) via reinforcement and/or punishment. Test items alternate with training stimuli which are reinforced to avoid response extinction. Subjects categorize test stimuli as consistent with Go or No-Go stimuli experienced during training; discrimination is measured as the proportion of correct responses.

(iv) Spontaneous Discrimination

Animals are familiarized with artificial languages for hours, and are then tested on strings of sounds consistent versus inconsistent with the familiarization language; discrimination is measured as a function of changes in calling behavior in response to the stimuli. In the newborn chick task, animals are exposed for hours to a structured visual stream of objects. Discrimination between consistent versus inconsistent stimuli is measured by the proportion of time spent near the screen playing the consistent stimulus (Figure I).

(v) Operant Conditioning

Animals are trained to select one type of stimuli associated with food (S+); responses toward S− are associated with bland punishment or no-reward. Discrimination is measured as a function of choice of consistent versus inconsistent stimuli with respect to S+. Alternatively, the training phase consists of presentation of stimuli associated with food reinforcement. Test items are words consistent with the language versus non-words, and the behavioral response (e.g., lever-pressing) associated with test stimuli is measured.

Figure I. Apparatus and Sample Stimuli Used to Investigate the Detection of Statistical Patterns in Chicks [13]. In the familiar sequence, the shapes are structured into pairs such that the first shape in a pair is always followed by the same second shape. In the unfamiliar sequence, the same shapes are presented in random order.
months later, infants’ generalization abilities become more robust. After familiarization with a pattern such as ABB, 7-month-old infants discriminate between novel syllables arrayed in this familiar structure from the same syllables in a new structure (e.g., ga-ti-ti vs ga-ti-ga [47]).

Infants perform similar computations in visual and auditory nonlinguistic domains, although with some constraints. It has been suggested that infants perform better if previously exposed to the same regularities implemented by speech ([48]; for counterexamples, see [49]), or, more broadly, communicative signals [50]. An alternative hypothesis suggests that successful learning is impacted by familiarity and/or ease of processing. For example, infants are more apt to generalize sequences of animals or upright faces than geometric forms or inverted faces [9,21,51]. Other perceptual and cognitive factors appear to constrain this form of learning in infants, including the presence of immediate repetitions of individual elements [9,21,52].

Nonhuman animals also generalize in sequential learning tasks, but with notable limitations. Zebra finches generalize based on the perceptual similarity between training and test materials rather than syllable order. In fact, in the absence of acoustic properties shared between familiar and novel streams, finches fail to generalize at all [53–55]. Similarly to human infants, finches seem to privilege patterns containing adjacent repetitions such as ABB and AAB [54]. Newly hatched chicks demonstrate robust generalization given training on triplets of visual objects arranged according to ABA, AAB, ABB, and BAA patterns, regardless of the presence of immediate repetitions [26]. Among mammals, rats trained on ABA sequences instantiated by strings of tones generalize to novel stimuli [56]. Similar results have been obtained with consonant–vowel alternations implementing ABB patterns compared to random sequences [57,58]. Rhesus macaques (Macaca mulatta) discriminate novel AAB versus ABB strings implemented by their own calls [59]; a recent extension of this study involved another primate species, the cotton-top tamarin, which can generalize structures presented in both speech and musical tones [60].

**Acquiring Simple Syntactic Structures**

By 12 months of age, human infants can learn rudimentary syntactic structures in laboratory tasks. Using artificial grammar learning paradigms, infants are first familiarized with simple miniature grammars that generate sets of sentences. Infants subsequently discriminate grammatical versus ungrammatical strings containing either violations of internal syllable pairs or violations at the edges of the grammar [32]. In a similar paradigm, 12-month-olds learned phrase structures grammars that mimicked the basic statistical patterns of natural languages (e.g., a determiner, such as ‘the’ or ‘a’, requires a noun somewhere within a sentence [34]). In this latter study, nonsense words from an artificial language were clustered in categories (e.g., determiners) whose presence depended on the presence of other categories (e.g., nouns). After exposure to the language, infants distinguished grammatical versus ungrammatical test strings which violated predictive dependencies between word categories. This task also required generalization of grammatical knowledge beyond the trained sentences.

The ability of songbirds to learn and generalize syntactic structures is somewhat more restricted. European starlings (Sturnus vulgaris) learn structures as complex as finite-state grammar (e.g., AB^n) and hierarchical grammar (e.g., A^nB^n) formed by their own song syllables [61]. Finite-state grammars generate strings of items repeated linearly (e.g., ABABAB) whereas hierarchical grammars comprise categories of items grouped into substrings that are nested into other substrings (e.g., phrase structure, AABBBB). However, at least in starlings, generalization to novel instances of the grammars (e.g., CDCDCD) is driven by acoustic similarity between training and test syllables, rather than by detection of the underlying patterns [62].
When the computations require transfer of structured information to novel inputs, these species exhibit limited capacities. Bengalese finches (*Lonchura striata domestica*) demonstrate more advanced generalization skills, taking advantage of statistical (predictive) dependencies between categories of song syllables, and generalize to strings composed of novel syllables [10]. This evidence is consistent with the previously described results from 12-month-old human infants, pointing to common processing of basic syntactic patterns between the two species.

Among nonhuman primates, tamarins learn finite-state and phrase-structure grammars [34,63]. However, in the latter case, monkeys failed to extract the structure when it required generalization to novel sentences, unlike human infants, suggesting that the operation is limited to the specific stimuli with which they were familiarized. In a series of artificial grammar learning tasks involving phrase-structure grammars (similar to [34]), marmosets (*Callithrix jacchus*) primarily encoded regularities involving the beginning of sentences, whereas macaques could also track violations throughout the strings [11]. Macaques have also been compared to human adults in tasks investigating learning of nonadjacent dependencies. In line with some findings on nonhuman primates [63,64], but not others [65], macaques exhibited no sensitivity to regularities involving nonadjacent syllables, only responding to violations of contiguous syllables. Humans performed better than macaques overall, detecting nonadjacent patterns [66].

**Species Comparisons and Constraints**

Despite many similarities, human infants and nonhuman animals diverge in the facility with which they perform various statistical learning tasks. Our interpretation of these cross-species differences considers perception and cognition, focusing on the extent to which these general abilities constrain statistical learning. A similar approach has been taken to explain early learning of grammar with respect to similarities between human and nonhuman performance. According to this perspective, grammar acquisition is supported by specialized perceptual and memory systems that are possibly shared with other species [67]. In the present review article we examine and interpret cross-species differences in comparable tasks based on domain-general abilities possessed by the species in question. Considering the creature under investigation *tout court* has the potential to provide a window into the ecological functions of statistical learning. To this end, we will examine statistical learning cross-species comparisons through the lens of three aspects of perception and cognition: vocal learning, perceptual processing, and memory.

**Relationship between Vocal Learning and Statistical Learning**

Vocal learning requires animals to modify the acoustic structure of the vocalizations of their own species and produce novel patterns of sounds. Through this mechanism, the young of some species acquire fundamental communicative signals that facilitate social interactions with conspecifics. Vocal learners include passerine songbirds, seals, cetaceans, and some bats. However, there are substantial differences in the way this process unfolds across species. Some species acquire a single novel vocalization as juveniles, whereas others learn new sounds even in adulthood [68,69]. Differences also include the structure of the learned sound patterns. The songs of some species provide more structural variability (e.g., European starlings), whereas others consist of fixed sequences of notes (e.g., zebra finches) [70,71]. Differences in vocal learning across species may predict the complexity of the patterns animals can learn and generalize in laboratory tasks [55]. In particular, species whose songs are syntactically structured and composed of a wide range of notes should be better at performing complex computations requiring, for instance, generalization of a given pattern to novel exemplars.
Consistent with this hypothesis, recent findings comparing two avian species in an artificial grammar learning task reveal differences that are consistent with general aspects (syntactic-like organization, phonological variability) of the vocal behavior of birds [55]. Zebra finches learn regularities such as ABA and AAB presented as short sequences of song syllables, but fail to generalize to structured strings formed by novel song syllables. However, the budgerigar (common parakeet, *Melopsittacus undulatus*) goes beyond item-specific information and generalizes these patterns, recognizing them even when they are implemented in novel sounds. This difference in performance can be explained in light of what is known about vocal learning in these species. The vocal repertoire of zebra finches includes highly stereotyped songs formed by rigid syllable sequences, resulting in repetitive, linear patterns [70–73]. Budgerigars are open-ended vocal learners, with high vocal plasticity, and their songs show greater phonological variability and syntactic-like organization [74,75]. We hypothesize that the perceptual and learning skills possessed by the finches may not be suited to performing computations more complex than rote memorization of specific syllable order. By contrast, the vocal learning abilities of budgerigars suggest the presence of learning mechanisms that can detect the underlying structure of sound sequences, later recognizing that structure when implemented by novel syllables (e.g., generalization). This hypothesis is supported by findings showing that budgerigars possess generally superior memory for acoustic complex stimuli [76]. The directionality of this relationship remains unknown. In line with theorizing about human language, we hypothesize that the learning abilities themselves have shaped the song structure of these species [4,77].

Visual Statistical Learning in Neonates

The development of perceptual processing also provides insights into the learning outcomes observed across species. Precocial and altricial species differ in the ontogeny of a range of physiological and behavioral functions. Precocial animals are biologically mature from hatching or birth, and are generally independent from parental care. For instance, superprecocial animals such as megapode birds (e.g., Australian brushturkey, *Alectura lathami*) leave the nest shortly after hatching, and display fully developed brain structures and motor behavior, being able to fly few hours after hatching [78–80]. As a consequence, the early perceptual processing performed by precocial organisms exceeds that of altricial species (see also [81]). These ontogenetic differences may be linked to the statistical learning skills present in a given species at birth. It is likely the case that altricial animals would have to deal with biological limitations (e.g., neural development, sensory and perceptual processing) that constrain what can be learned at the outset of postnatal life.

From this perspective, consider the comparison between human neonates – an altricial species with an extended developmental timeline – and newly hatched domestic chicks, a precocial bird species. In a visual statistical learning study, newly hatched chicks discriminated structured from random streams of four and six shapes [13]. Human neonates, however, succeeded only with streams of four shapes, failing with six-shape streams [41]. Limited perceptual abilities are likely to constrain visual statistical learning in humans at birth, whose visual system is immature (e.g., severely reduced acuity [82]), generally limiting visual learning [83]. Such restrictions are typical of the altricial human primate, whose offspring stay immature longer than other mammals [84,85]. Unlike humans, chicks hatch in an advanced stage of development, with completely developed visual pathways from the very first days of postnatal life [86–88]. In chicks, vision is the predominant sensory modality, allowing full processing of complex visual stimuli immediately after hatching. Compared to chicks, reduced visual skills in human infants appear to constrain statistical learning, limiting the extent to which infants can process statistics over particular inputs. Indeed, a classic theory of perceptual development hypothesizes that
limitations on the amount of information computed – owing to the protracted maturation of some sensory systems relative to others – actually facilitate human perceptual and cognitive development [89].

Statistical learning is likely to guide different functions in humans and chicks. In humans, early learning of regularities plays a key role in domains other than vision, especially language processing. In chicks, however, extracting regularities might be linked with early social learning, a fundamental capacity that allows newly hatched chicks to recognize the mother hen and siblings. This process, filial imprinting, occurs immediately after birth via learning of invariant visual features of their social companions, such as plumage color patterns as well as beak and head shape [86,90]. We hypothesize that statistical learning works in tandem with filial imprinting in chicks, and leads to an integrated representation of relevant social objects that will allow further recognition and identification. Consistent with our hypothesis, a recent study shows that imprinting in chicks promotes the learning of multimodal regularities such as XX versus XY implemented by sound–shape pairs as well as generalization to novel instances [91], see also [92]. Being precocial requires chicks to process salient visual stimuli at birth, whereas altricial humans do not have to rely on early visual learning to identify relevant social inputs. In this view, statistical learning abilities are influenced by the ontogeny of the species, which determines the functions supported by the learning process (i.e., language acquisition, recognition of social objects, etc.).

Memory and Statistical Learning
Memory is a fundamental component of statistical learning. Learners must keep track of sequences of elements that rarely persist over time, and temporarily hold information necessary for future processing (i.e., working memory [93]). According to the memory-based framework presented in [94], regularities extracted from structured input lead to representations in long-term memory even after short exposure (see also [95]). Memory traces then become the foundation for subsequent learning operations, driving detection of further patterns, and integrating stored information to find common regularities across exemplars [96,97]. Computations such as extraction, storing, and integration seem to be fundamental for both acquiring sequential regularities and generalizing to novel exemplars. This framework also points to the way in which learners retain and retrieve information as a constraint on statistical learning, suggesting that sensitivity to patterns in the sensory input is shaped by the memory skills of learners (e.g., storing, access). Following this path, we hypothesize that species with reduced memory skills would be sensitive to a restricted set of structured information from sequentially presented inputs compared to species with enhanced storing and retention abilities.

Among nonhuman primates, marmosets and macaques have been directly compared in artificial grammar learning experiments to test learning of sequential syntactic patterns. When presented with strings violating the familiar grammar at multiple locations, marmosets, unlike macaques, detect only violations at the very beginning of the grammar [11]. One explanation for this pattern of performance points to cross-species differences in memory skills. These species belong to separate groups of the primate order, with distinct physical features and cognitive abilities. Macaques are equipped with superior general learning and memory capacities, outperforming marmosets in basic learning tasks such as the discrimination of learned stimuli [98,99]. In particular, some species of the Callitrichidae family, including the common marmoset, seem to use spatial memory while foraging, concurrently tracking several object locations containing food [100–102]. It is likely that such differences affect other computations that closely involve learning, such as statistical processing of grammatical sequences. In this view,
marmosets developed better memory skills in the spatial modality than in the temporal modality, thus performing worse on tasks requiring retention of sequential stimuli.

The general learning systems of nonhuman primates may not be tailored to acquire linguistic structures as complex as hierarchical grammars (see also [103]), but do support learning of sequential patterns in other domains such as tool-use [104,105], motor actions [106,107], and social communication [108] which require the processing of events occurring over time and play a prominent role in the behavioral repertoires of primates. It is possible that the use of linguistic materials in studies investigating sequential statistical learning has hindered the discovery of more advanced abilities in nonhuman primates (see Concluding Remarks). Experiments using ‘grammars’ created not from words but from sequences of tools or actions might reveal statistical learning abilities that are not observed with linguistic materials. More generally, interactions between statistical learning abilities and the elements over which learning occurs (e.g., speech syllables vs motor actions) point to important constraints on statistical learning, and may help to explain divergent learning outcomes across species.

Concluding Remarks
In this review we have framed cross-species differences in statistical learning in the context of perceptual and cognitive mechanisms that vary across organisms. It is clearly the case that reduced perceptual abilities in a given modality lead to limited learning of structured inputs, as in the case of human neonates. In a similar vein, reduced memory capacity affects regularities detected from sequential inputs, confining learning to a restricted portion of the information available in the input.

According to this perspective, we might expect similar computations in species that share cognitive functions supported by statistical learning. For instance, statistical learning is likely to drive vocal learning in organisms that must learn to produce structured vocalizations [71,109,110]. Similarly to human infants acquiring languages, juveniles of some songbird species track the statistical properties of songs, and recombine statistically coherent patterns into new songs [37,38]. These observations suggest that both acoustic features and the structural organization of the songs are acquired during vocal learning, making this process similar to the beginning of human language acquisition. It is thus not surprising that some bird species and humans exhibit the most advanced statistical learning of sound sequences.

Vocal learning is not limited to birds and humans. Bats, seals, cetaceans, and elephants meet the requirements for vocal learners: they can learn novel vocalizations and imitate the sounds of other species. Among cetaceans, bottlenose dolphins (Tursiops truncatus) possess impressive vocal learning and mimicry abilities: they learn to produce new vocalizations, have sophisticated communication systems composed of extended call repertoires, and generally possess remarkable cognitive skills [111–113]. For these reasons we would expect dolphins to show similar statistical learning capacities to human neonates and songbirds [114]. In a similar vein, we would expect non-vocal learners to show restricted abilities in artificial grammar learning tasks. For example, domestic chickens are limited to short repetitive calls [115,116] whose usage is acquired following social experience with conspecifics [110]. Given that there is no vocal learning in this species, we predict that young chicks would fail to track probabilistic structures from auditory stimuli, performing worse than vocal learners.

Statistical learning may support communicative functions even in the absence of vocal learning. Fieldwork suggests that free-ranging putty-nosed monkeys (Cercopithecus nictitans) can track co-occurrences between specific combinations of natural calls and the presence of an eagle,
eventually emitting vocalizations to signal the predator presence to the rest of the group [117]. Black-fronted titi monkeys (*Callitecia nigirfrons*) produce call sequences whose combination and type vary based on the context (e.g., aerial vs terrestrial predators [118,119]). Female baboons (*Papio cynocephalus ursinus*) appear to possess a combinatorial system of communication reflecting the complex hierarchy of their social group [108]. For example, female baboons notice inconsistent call sequences with respect to the rank of the caller, recognizing when a dominant female emits unusual vocalizations when interacting with subordinates [120].

Statistical learning abilities also impact on non-communicative domains across species. Guinea baboons (*Papio papio*) demonstrated learning of hierarchical structures from visual stimuli [121] and orthographic patterns [23]. Baboons learned statistical relations between printed letters and their positions within a word, distinguishing English words and nonwords with remarkable accuracy. The main theoretical implication of this work is that orthographic processing, an essential computation in reading, might be rooted into more general statistical learning mechanisms shared across baboons and humans, and appears to be constrained by domain-general abilities necessary to discriminate visual objects (e.g., detecting feature combinations).

Further consideration of the role of the perceptual and cognitive mechanisms supporting statistical learning will help to clarify how this process unfolds in different ecological niches (see Outstanding Questions). In human infants, the evidence suggests that statistical learning abilities have impacted on the types of structures that are observed in human languages [4,33,34]. It is also the case that what human infants have already learned impacts on the types of structures that they subsequently learn most readily [96,122]. Both of these directions of effects may influence statistical learning in non-human animals as well. Interpreting cross-species findings in this light links our approach to perspectives on human learning that considers statistical learning as a core component of cognitive systems, rather than as an independent computational process [3]. This view is also aligned with modern neurocomputational theories which treat brains as sophisticated prediction machines [123], internalizing probabilistic models of the environment to anticipate the sensory stream and generate inferences [124,125]. Shifting the focus of comparative research to the systems within which learning is embedded will us allow to develop a much deeper understanding of the ecological relevance of statistical learning.

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