INTRODUCTION

Isolated crinoid ossicles, reflecting complete post-mortem disarticulation of individuals during extended residence within the taphonomically active zone (Lewis 1980), represent one of the most abundant bioclasts in the Paleozoic rock record (Lowenstam 1957; Ausich 1997). Although used to some degree in regional biostratigraphy (e.g., Moore 1938; Fearnhead et al. 2013; Thomka and Brett 2017) and erection of descriptive morphotaxa (e.g., Moore and Jeffords 1968; Donovan 1986), fragmentary crinoid remains are largely under-utilized data sources, particularly in studies of taphonomy, paleoecology, and facies analysis (see Jeffords 1978; Holterhoff 1997; Thomka et al. 2012 and references therein). In addition, isolated crinoid ossicles contribute to greater understanding of the internal morphology and intraspecific variability of taxa that are known from a limited number of specimens. Consequently, careful attention to disarticulated crinoids shows promise for enhancing paleoenvironmental reconstructions, improving understanding of biotic interactions, and increasing records of echinoderm biodiversity and paleobiology.

Nevertheless, articulated or partially articulated material is generally needed to formally describe new crinoid taxa. This can leave associated skeletal material in the form of isolated ossicles collected alongside more complete specimens insufficiently studied or completely ignored in favor of the articulated specimens. This study is an exploration of crinoid material consisting exclusively of loose ossicles representing supplementary material to the articulated cups that served as the sole basis for description of a crinoid species. These specimens provide additional documentation of a taxon with a relatively sparse fossil record and, more importantly, provide paleoecologic and taphonomic information not presented in the initial description.

METHODS AND MATERIALS

Specimens described here are reposited in the Cleveland Museum of Natural History invertebrate paleontology collection, all under lot number CMNH 16597. A diverse collection of isolated basal plates, radial plates, brachial and anal sac spines, and columnals, collectively representing multiple crinoid taxa, is present. In total, 153 plates are included in this specimen lot, with the distribution of ossicle types given in Fig. 1. Although many plates cannot be reliably identified, 35—including 15 basal plates and 20 radial plates—are herein attributed to the pinnulate cladid crinoid Diphuicrinus ohioensis Burke, 1976 (Fig. 1).
All plates were recovered from the Middle Pennsylvanian Putnam Hill Limestone of the Allegheny Group from an abandoned strip mine north and east of a tributary of Elk Fork, about 3.4 km northwest of McArthur, Vinton County, southeastern Ohio (lat 39°16´35˝N, long 82°29´40˝W). This locality is the same as “Locality 3” of Burke (1976), a site from which several of the articulated cups that serve as type material for *D. ohioensis* were collected, namely paratypes USNM (United States National Museum of Natural History) 166575 and USNM 166576. However, the specimen label of CMNH 16597 states that the assemblage of isolated ossicles was collected by J. J. Burke in 1965, over a decade before publication of the description of *D. ohioensis*. Consequently, these ossicles have long been listed as unidentified and were not included in the formal description of this species (Burke 1976).

**RESULTS**

A sample of isolated cup plates in CMNH 16597 closely matches the description for corresponding portions of *D. ohioensis* given by Burke (1976). The most distinctive features are: (1) the relatively large forefacet (aboral ligament fossa) on radial plates, an inherent feature of diphuicrinids (Strimple and Moore 1971), with a diagnostically shallower forefacet relative to internal facetal areas (Burke 1976); and (2) the prominent sculpture on exteriors of basal (Fig. 2) and radial plates (Fig. 3), which is coarser and characterized by more discrete nodes than in most other crinoids with otherwise similar calyx plating patterns and cup morphologies, such as *Graffhamicrinus* Strimple, 1961 (Strimple 1977). As indicated in Fig. 1, isolated radial and basal plates are described in this study; although some non-spinose brachials may represent *D. ohioensis*, the arms of this species are currently unknown, so brachials cannot be reliably identified.

Approximately 74.3% of *D. ohioensis* ossicles (12 of 15 basal plates, 14 of 20 radial plates) are encrusted by minute, threadlike, calcareous tubes (Fig. 2B-F, Fig. 3B-D). These are superficially similar to those produced by serpulid (polychaete annelid) worms in modern marine environments. However, true serpulid structures are restricted to younger deposits (Vinn and Mutvei 2009; Ippolitov et al. 2014). Crinoid ossicles vary from being relatively densely encrusted—close to 10% of surface area covered; Fig. 2F, Fig. 3D—to only having 1 or 2 short fragments preserved (Fig. 2B, Fig. 3B). Plates are most commonly encrusted on the exterior surfaces, which could have occurred either before or after the death of the crinoid; however, a number of specimens also have encrusting tubes on the interior and/or articular surface of plates, which could only have occurred after the death and total disarticulation of the crinoid. Regardless, 2 patterns are worth documentation: first, where the exterior of plates are encrusted, tubes are present primarily in the low areas in between elevated nodes; second, that long, continuous lengths of calcareous tubes are completely absent and, instead, encrusting calcareous tubes are present in the form of short, incomplete portions (Fig. 2, Fig. 3).
DISCUSSION

As no isolated ossicles were discussed in the original description of *D. ohioensis* (Burke 1976), the discovery of a sample of cup plates represents an additional record of a crinoid taxon known only from a limited number of specimens. Appropriate material within CMNH 16597 can be properly identified, and loose basal and radial plates can now be used to supplement articulated cups in future studies concerned with the morphology of diphuicrinid crinoids. Continued attention to isolated ossicles and other forms of fragmentary material is strongly encouraged. Nearly all museums have an abundance of echinoderm material catalogued as indeterminate loose plates (or the like), and a wealth of information on biodiversity, distribution, and preservation can potentially be gained through scrutiny of specimen lots that are largely overlooked in favor of articulated material.

The high frequency of encrustation of *D. ohioensis* ossicles by small, calcareous tubes was not previously documented in the initial description and interpretation of this taxon. Scanning electron microscopy of these tubes revealed at least 2 recognizable forms (Fig. 4). Most can be identified as microconchids (Fig. 4A-E), which are enigmatic.
tentaculitoids that are coiled, at least initially (Zatoń and Vinn 2011; Zatoń et al. 2016; Fig. 4A); a smaller proportion of tubes can be attributed to agglutinated foraminifera (Fig. 4E-F). The microconchids are essentially hollow calcareous tubes showing evidence of accretionary growth (Fig. 4C). The foraminifera are characterized by poorly sorted tests composed of angular sediment grains (Fig. 4F). Although the presence of encrusting organisms offers little in terms of taxonomic information related to crinoids, encrustation represents an important paleoecological and taphonomic phenomenon, and one of the most useful indicators of paleoenvironmental parameters (Brett and Baird 1986; Parsons and Brett 1991). Elevated relative frequencies of encrustation of isolated crinoid ossicles, particularly when encrustation occurs on both the exterior and interior/articular surfaces of cup plates, as in the collection studied here, is a strong indicator of stratigraphic condensation associated with a very low sedimentation rate (Thomka et al. 2012). Although this interval of the Putnam Hill Limestone has yielded articulated cladid crinoid cups, as described by Burke (1976), these reflect relatively rare obrution (rapid burial) events that episodically punctuated an environment with an otherwise minimal sediment influx (e.g., Brett and Baird 1986; Thomka et al. 2012).

The preferential encrustation of smooth, depressed areas in between the elevated nodes on basal and radial plate exteriors is an unusual taphonomic feature. There are 3 potential explanations for this pattern. The first interpretation is that the lower areas offered some advantage in terms of strength and/or persistence of currents. This seems least likely given the tendency of suspension-feeding organisms to more commonly encrust elevated areas over depressed ones (Taylor and Wilson 2003); however, the orientation of isolated plates on the seafloor or the overall microtopographic complexity of the coarsely sculptured plates may have somehow contributed to unusual water flow in the areas between nodes. The second interpretation is that the spaces in between nodes represented broader, flatter areas that were more suitable as sites for encrustation. Although a few portions of calcareous tubes extend from lower areas up onto the bases of nodes, these segments are no more than a few millimeters in length and never wrap around lower portions of nodes, suggesting that such structures might have represented substrata that were not preferred. The third interpretation is that

**FIGURE 3.** Examples of radial plates belonging to *Diphuicrinus ohioensis* from the Middle Pennsylvanian Putnam Hill Limestone of Vinton County, Ohio (all reposited as CMNH 16597). (A) Well-preserved, clean plate lacking encrusting organisms. (B) Sparsely to moderately encrusted plates with segments of tubes occurring primarily in the right side of the specimen. (C) Moderately encrusted plate with numerous, highly arcuate partial tubes at the base of nodes primarily in the center of the specimen. (D) Poorly preserved, heavily encrusted plate with most of the surface area in between prominent nodes covered in thin, highly incomplete tubes. All scale bars = 5 mm.
FIGURE 4. Scanning electron microscope images of encrusting vermiform tubes on isolated plates of *Diphuicrinus ohioensis*. (A) Example of a contorted microconchid tube on a basal plate, with a highly weathered specimen visible on the bottom of the node in the background. (B) Relatively straight microconchid tube in between 2 elevated nodes of a basal plate. (C) Close-up view of the aperture of a microconchid tube on a radial plate. Note that the morphology is a simple, undivided, hollow tube. (D) Larval settlement area for a microconchid showing the diagnostically coiled orientation. (E) Agglutinated foraminiferan encrusting the node in the left foreground, with several heavily weathered microconchid tubes in the center and upper portion of the image. (F) Close-up view of elongate agglutinated foraminiferan test showing the poorly sorted texture.
encrustation was initially more ubiquitous across the exterior of cup plates, but post-mortem abrasion by currents removed portions of the calcareous tubes that were exposed on elevated nodes. See similar phenomenon described in Donovan (1999) for encrustation of pumice fragments. Given the evidence for slow sedimentation and consequent extended residence time of bioclasts in seafloor sediment prior to burial (see above), prolonged exposure to natural sandblasting by currents may have effectively stripped away encrusters from the nodes, thereby restricting their preservation to the protected inter-node areas. This may be supported by the incomplete and seemingly fragmented state of encrusting tubes (Fig. 2, Fig. 3, Fig. 4). Similar patterns are likely to be documented if careful attention is paid to the distribution of encrusting organisms on sculptured echinoderm ossicles in future studies. Information derived from such studies has the potential to clarify the precise cause(s) of the patterns described here.

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