Scanning electron microscopy of human islet cilia

Alexander J Polino
Sanja Sviben
Isabella Melena
David W Piston
Jing W Hughes

Follow this and additional works at: https://digitalcommons.wustl.edu/oa_4

Part of the Medicine and Health Sciences Commons

Please let us know how this document benefits you.
Human islet primary cilia are vital glucose-regulating organelles whose structure remains uncharacterized. Scanning electron microscopy (SEM) is a useful technique for studying the surface morphology of membrane projections like cilia, but conventional sample preparation does not reveal the submembrane axonemal structure, which holds key implications for ciliary function. To overcome this challenge, we combined SEM with membrane-extraction techniques to examine primary cilia in native human islets. Our data show well-preserved cilia subdomains which demonstrate both expected and unexpected ultrastructural motifs. Morphometric features were quantified when possible, including axonemal length and diameter, microtubule conformation, and chirality. We further describe a ciliary ring, a structure that may be a specialization in human islets. Key findings are correlated with fluorescence microscopy and interpreted in the context of cilia function as a cellular sensor and communications locus in pancreatic islets.

Primary cilia are antenna-like sensory organelles that extend from the centriole and project into the extracellular space, serving important functions in sensory detection and signaling. Since their initial description by electron microscopy (EM) in the rodent pancreas (1, 2), primary cilia have been increasingly examined for their link to human metabolic disease, where altered cilia expression or function disrupt pancreatic development and glucose homeostasis (3–7). The pancreatic islets of Langerhans are spherical clusters composed of multiple endocrine cell types. Most cells in the islet, as in the rest of the body, project a single primary cilium, its structure most definitively studied by EM (8–12) in addition to targeted protein staining by immunohistochemistry. In human islets, primary cilia were first described in beta cells, in 1964 in a case of insulinoma (13), whereas validation in nondiseased islet tissues took decades to follow (14–16). More recently, islet primary cilia were found to have glucose- and ATP-regulated motility (16, 17), suggesting that primary cilia are not simply static sensory organelles. These findings have fueled growing interest in islet primary cilia in diabetes research, paralleled by an increasing need to understand the basic structure and composition of these cellular projections. What are their shape and size? What is the mechanical basis for their movement? How might their signaling be compartmentalized? Toward answering these questions, we first aimed to determine the physical parameters of islet primary cilia.

Structurally, primary cilia are divided into several subdomains that are conserved among species and cell types. Lengthwise, cilia are composed of a basal body and an axoneme. The basal body is the mother centriole of the cell, a microtubule-based cylinder that emanates from near the Golgi and remains entirely intracellular. It anchors the primary cilium via a connecting transition zone, a morphologically distinct segment that projects out of the ciliary pocket and controls cargo trafficking to the axoneme. The primary cilium axoneme has a highly conserved conformation of nine outer microtubule doublets with a lack of central microtubules, hence named “9+0,” a pattern consistently observed at the axonemal base and thought to continue through the length of the cilium. This model has required revision, however, due to reports of non-9+0 cilia cross-sections captured by transmission electron microscopy (TEM) as well as 3D tomography of primary cilia in cells of both pancreatic and nonpancreatic origin (9, 18–22). The updated view is that the canonical ciliary “9 + 0” configuration changes soon after leaving the cell body, and that microtubule number and arrangement may be modified along the length of the cilium. We recently reported on the non-9+0 arrangement of human islet cilia, but only in sporadic thin sections of resin-embedded tissue prepared for TEM (16), while how the axoneme structure might evolve from base to tip in islets is unknown. Many other morphometric features of human islet primary cilia also remain undefined, including axonemal length and diameter, geometry of microtubule filaments, whether they assemble higher order conformations, and the presence of any accessory components contributing to the maintenance of axonemal integrity, dynamics, and sensory signaling function. Thus, elucidating islet cell cilia structure and especially visualizing the entire axonemal cytoskeleton is a priority in islet research.

Significance
Primary cilia are vital cell-surface sensory organelles, but their physical dimensions have eluded characterization due to difficulty in isolating and studying these enigmatic structures, especially in human tissues. In the present study, we use a multi-scale surface scanning approach to enable a 3D architectural study of human primary cilia, the first in pancreatic islets and in any human tissue. Results provide a morphometric basis for understanding ciliary function including signaling capacity, axonemal motility, and cell-cell communication.


The authors declare no competing interest.

This article is a PNAS Direct Submission.

Copyright © 2023 the Author(s). Published by PNAS.

Author affiliations: “Department of Cell Biology and Physiology, Washington University School of Medicine, Saint Louis, MO 63110; “Washington University Center for Cellular Imaging, Washington University School of Medicine, Saint Louis, MO 63110; and “Department of Medicine, Washington University School of Medicine, Saint Louis, MO 63110

To whom correspondence may be addressed. Email: jing.hughes@wustl.edu.

Published May 19, 2023.
Advances in superresolution and electron microscopy combined with increasing availability of human donor specimens makes it now possible to perform ultrastructural studies on primary human islets. Large open-access databases such as the nPOD nanotomy atlas have curated scanning transmission EM images of human islets (23). While these images are useful for demonstrating disease-related morphological alterations across cellular compartments, cilia and centrosomes were not examined in these studies owing to their rarity and low likelihood of identification. To enable ultrastructural characterization of islet primary cilia, we used surface topography imaging by scanning electron microscopy (SEM) to directly visualize the ciliary axoneme in human pancreatic islets. This approach allows cilia identification based on their extracellular localization and distinct morphology and preserves full-length cilia structure in situ. We examined cilia with or without demembranation to compare surface versus cytoskeletal details of primary cilia, both contextually within intact human islet tissues (Fig. 1). Ciliary components post-demembranation were preserved with clarity, allowing quantification of microtubule size, number, chirality, and observation of structural motifs not previously described in other mammalian cilia.

Results

We examined individual cilia on 15 islets from three human donors (Table 1). The islets in our study were isolated from human donor pancreata and ranged in size from 100 to 300 μm in diameter, consistent with dimensions reported in literature. Intact islets exhibited round shape and well-preserved surface contour, with the contrasting textures among adjacent cells exhibiting a soccer ball-like appearance (Fig. 2A). Primary cilia are prominent structures on the islet surface, typically measuring 3 to 6 μm in length, are flexible and display varying curvature. Other membrane protruberances such as microvilli and filopodia are also observed on the islet surface. The vast majority of human islet cells bear a single cilium each, a feature that we and others have previously demonstrated by immunofluorescence microscopy (14, 16, 24, 25).

In the present study, primary cilia were imaged with or without demembranation, the latter revealing a clear view of the axonemal structure from base to tip. Attrition in cilia number occurred at each step of sample preparation and image acquisition. In total 60 full-length cilia including 50 membrane-extracted and 10 non-extracted samples were selected for morphometric analysis. Multi-scale SEM was used to visualize structural details of each cilium and its subdomains with progressively higher resolution.

Ciliary Length, Diameter, and Volume Estimations. Most cilia on the islet periphery project away from the cell, while some project horizontally along the membrane (Fig. 3). We sought to determine the linear dimensions of islet primary cilia, keeping in mind the caveat that measurements of 3D object length in 2D images are by nature underestimations. By virtue of the object tilt, planar-extrapolated lengths can only be equal to or shorter than actual object length. To achieve the best estimation of ciliary length, we selected mostly cilia that were captured in profile perpendicular to the imaging detector. Average axonemal length measured 5.74 ± 2.3 μm among 50 extracted cilia and 4.21 ± 1.5 μm among 10 unextracted cilia, with comparable distributions (Fig. 3A). As human islet cilia are wider at the base than tip, we measured ciliary diameter both proximally and distally, obtaining an average of 237 ± 52 nm at the base and 148 ± 37 nm at the tip for extracted cilia (Fig. 3D). These dimensions were similar in unextracted cilia, 240 ± 80 nm at the base and 121 ± 51 nm at the tip. Overall length and diameter measurements were comparable to those determined by cryoelectron tomography (cryo-ET) in cultured kidney cell cilia and those reported for other mammalian cilia (19–21).

Using averaged length and diameter measurements from all cilia, we estimated the human islet primary cilium volume and surface area based on the shape of either a truncated circular cone or a right cylinder (Fig. 3F). As a truncated cone with a base and tip diameter of 0.24 and 0.14 μm (radius 0.12 and 0.07 μm) and height of 5 μm, the cilium volume is estimated to be 0.15 μm³, with a surface area of 3.05 μm². As a cylinder, with an average axonemal diameter of 0.2 μm (radius 0.1 μm), the cilium has a volume of 0.16 μm³ and surface area of 3.2 μm². Thus the two model methods are in good agreement with each other.

We next measured the average diameter of single islet cells in our dataset, which was 12 to 14 μm (radius 6 to 7 μm), and determined the cytoplasmic volume to be ~1,000 μm³. This allowed us to calculate the ratio of cytoplasmic volume to ciliary volume for human islet cells as approximately 6,000 to 7,000. The ratio of plasma membrane surface area to ciliary membrane area is about 150 to 200. These ratios were previously estimated to be 5,000 and 500, respectively, for a typical 0.3 × 5-μm primary cilium (26). The minor discrepancy between our and previous
estimates is due to a smaller diameter of human islet cilia than that generally assumed (0.24 μm versus 0.3 μm at the base). Thus, human islet cilia appear slightly more slender than what is typical for other mammalian primary cilia. Nonetheless, the values we obtained for human islet cilia are comparable to prior estimates from nonislet cells (26, 27), indicating that the physical dimensions of primary cilia are largely conserved across tissues and organisms.

The actual shape of the human islet primary cilia, of course, is not a uniform cone or cylinder. Our SEM imaging revealed remarkable cilia geometries and structural motifs, which we now describe from base to tip. Special attention is given to axonemal domains with known functional importance, as well as previously uncharacterized structural elements.

**The Ciliary Pocket and Transition Zone Are Well-Visualized by SEM.** The ciliary pocket is an invagination of the plasma membrane from which the primary cilium emanates. It is a specialized domain for ciliary protein trafficking, endocytosis, and actin cable docking, among other functions (28, 29). Ciliary pockets had been observed in mouse islets more than a half century ago by TEM, in beta cells exhibiting deeply rooted cilia where the ciliary pocket was interpreted as a site of basal body docking to the plasma membrane, with a presumed role in primary ciliogenesis (1). In the present study, the ciliary pocket of human islets is seen to form a circular pit surrounding the axoneme in some cells, while in others the cilium erupts from the surface without an appreciable ciliary pocket, and there are other cells still that have such deeply rooted cilia within recesses of the cell that a ciliary pocket is not visible (Fig. 4). From these observations we conclude that the ciliary pocket adopts specialized configurations among different human islet cells, a theme that is consistent with the differential membrane organization at the cilia base that has been described in other systems (28–31).

Higher up, the transition zone is the proximal-most region of the axoneme and is captured in striking clarity in our study. There is a section of 5 to 6 parallel rows of particles that encircle the cilia base orthogonal to the direction of axonemal microtubules (Fig. 4 B–D), corresponding to the “ciliary necklace” that has been described at the neck of the cilium in many species, where it supports the microtubule transition from triplets to doublets and acts as a membrane diffusion barrier (32–36). The necklace in human islet cilia measures approximately 230 nm in diameter and 140 nm in height in the visible portion above the cell. Individual bead-like particles comprising the necklace measure approximately 12.5 nm in diameter each, similar to the dimensions of the intramembrane particles previously described for kinocilia in the respiratory epithelium (37). The distance between the necklace rows, or strands, is about 18 to 22 nm, and the number of extracellular strands ranges between 3 and 6 per cilium. Whereas the ciliary necklace had best been previously demonstrated by freeze-etching EM (32, 37), a technique that relies on a carbon cast replica of the structures in study, our imaging approach visualizes the structure itself, allowing direct morphometric measurements of its components.

Distal to the ciliary necklace, we see cylindrical strands of microtubule doublets forming the ciliary axoneme. Small knob-like densities with diameters of ~20 nm are observed.
decorating the microtubules and are similar in size and shape as the beads that constitute the ciliary necklace (Fig. 4). These particles are also present on microvilli and cortical cytoskeleton, in a dense albeit irregular way, thus appear nonspecific to cilia and may represent membrane remnants or debris postextraction and post-fixation. Occasional larger appendages are seen to associate with the axoneme surface (Fig. 5 A), but we did not observe regular repeats of polymer complexes resembling intraflagellar transport trains (19, 38), which may not have survived demembranation.

Cilia Are Tethered by Microfilaments at the Base. We observe physical interactions between the ciliary axoneme and ciliary pocket as well as with the cell surface. The base of the primary cilium is connected by thin filaments to the surrounding microvilli and cytoskeleton surface (Figs. 4 A–C and 5 A and B). These microfilaments have a diameter of 7 to 10 nm and a branching form, consistent with actin fibers which are known to associate with primary cilia—particularly at the base (28, 39–41). Location and morphology-wise, they are distinct from the Y-links in the transition zone of cilia and flagella (21, 32, 38, 42, 43), which are symmetrically present on all microtubules and serve to tether the microtubules to the ciliary membrane. In contrast, these external filamentous strands observed in our study are located above the transition zone, attach to distant targets including neighboring microvilli and cell surface, and are asymmetrically arranged around the axoneme, predominating on one side of the cilium (Fig. 4 A and B). Similar filaments are also present on microvilli, where they are more symmetrically distributed and create an aster-like network. Neither cilia- nor microvilli-associated filaments are visible in nondemembranated samples (Fig. 5 C and D), suggesting that these structures are normally sheathed by the ciliary and plasma membranes. Fluorescence imaging of mouse islets using structured illumination microscopy provides supporting evidence that the base of islet primary cilia is embedded in an F-actin network (Fig. 5 E and F).

Microtubule Doublet-to-Singlet Transition. In all observed cilia, individual microtubule filaments start as doublets at the cilium base and transition to singlets at the one-third to half-way mark along the axoneme, between 1 and 2 μm from the base, at which point the cilium takes on a noticeable change in diameter and width (Fig. 6). There is a clear demarcation where this change takes place, beyond which point the microtubules become thinner and more loosely associated, reflecting a change in filament organization and intraflagellar cross-linking. Such microtubule doublet-to-singlet transitions have been reported in cultured kidney cell cilia (19, 21) and can be observed in open-access volumetric imaging of rodent and human pancreatic islets (20, 22, 23). Direct measurements from our dataset showed that human islet cilia microtubule singlets possess an average diameter of 37 ± 4 nm, approximately half that measured of microtubule doublets (68 ± 10 nm). These numbers are consistent with those previously reported in kidney primary cilia (19, 21). Small globular densities are seen coating the microtubule doublets in the proximal axoneme, whereas they are generally sparse or absent on distal portions of microtubules (Figs. 6 and 7).

Cilia Handedness and Secondary Structure. As microtubule filaments evolve from doublets to singlets, they undergo conformational changes with respect to each other. Whereas they begin as a tight straight bundle in the ciliary basoproximal region, the microtubules begin to separate distally and twist about a central axis before coalescing at the tip (Fig. 7). There is a periodicity to the microtubule rotations, measuring 900 to 1,000 nm in arc length with a pitch of 500 to 1,000 nm between turns. Within the loose microtubule bundle, actin-like thin filaments with a
diameter of ~8 nm are occasionally observed stretching along the long microtubule axis (Fig. 7C). Occasional microtubules break or splay apart (Fig. 7E and F), likely as a consequence of physical disruptions from extraction and fixation. Given its helical nature, the axoneme is intrinsically chiral. Human islet cilia are predominantly left-handed (>80%), while a minority have a mild right-handedness (~8%), and a similar minority have no discernable rotation (Fig. 8A). This is in contrast to the predominant right-handed spins reported for kidney cell cilia (19, 21). To be sure of our observations, we confirmed using fiducial markers that our SEM images are not mirror images. The presence or absence of taxol also did not alter the rotational nature and direction (Fig. 8B), suggesting that these higher order microtubule conformations are fairly stable.

A Ciliary Ring Is Occasionally Observed in the Proximal Axoneme. We observe a rare but potentially important feature, which is that a ring is present in the proximal axoneme, at about 1 to 1.5 μm from the base. It wraps around the axonemal bundle as a continuous strand with similar texture and diameter as a microtubule singlet or doublet (30 or 60 nm) (Fig. 9). We name this structure the “ciliary ring” to differentiate it from the ciliary necklace and ciliary bracelet, which are separate structures (32, 44, 45). Rings are observed on 4 out of 50 demembranated cilia (8%) and were present on islets from multiple donors, thus not an isolated phenomenon. All rings occurred on cilia from cells that possessed short microvilli and therefore were likely beta cells. One cilium was observed to bear two rings that subdivide the axoneme into equidistant thirds (Fig. 9C–E). Distal to each ring, the axoneme exhibits greater separation and twist, appearing as if the ring is acting as a band to keep the microtubule bundle together. Rings are not observed on unextracted cilia, indicating that the structure occurs submembrane.

Ciliary Tip as a Special Domain. The distal ciliary domain, or tip segment, is captured intact in demembranated islet cells, showing structural organization that differs from the rest of the axoneme. The microtubules decrease in number toward the distal cilium before terminating into a dense cap structure (Fig. 10A–C). There is diversity in the number of microtubules that make it to the tip, typically between 3 and 5, but up to 7 have been counted. A greater diversity in cilia tip morphology is seen in nonextracted samples in which the cilium retains its membrane covering, where the enclosed tip shape ranges from pointed to rounded to bulbous (Fig. 10D–F). The highly dilated tip seen in Fig. 10F resembles those previously described in normal kidney collecting duct cilia (46) as well as cilia bearing mutations affecting retrograde transport that cause a buildup of cargo and vesicles in the tip region (47–49). Our specimens came from healthy donors with no known metabolic disease or ciliopathies, suggesting that these morphological differences may represent a normal spectrum of pancreatic islet cilia in various stages of growth and remodeling. Overall, the unique morphology of the cilia tip observed in our study is consistent with its role as a dedicated sensory and signaling domain, where the diversity of tip structure may be related to specific ciliary roles in different islet cells.

Multiciliated Islet Cells. Occasional compound cilia are observed by SEM where a single cell may project two or even three cilia from the same base. These are rare, estimated <5% frequency among human islet cells. We observed three cases of double cilia and one case of triple cilia. One clear example each is shown in.
In both cases, the double and triple cilia projected axonemes that were similar in length and diameter and exhibited varying degrees of physical interactions among the axonemes. These project as independent axonemes, analogous to the human dermatologic condition of *pili multigemini* or “tufting” where...
multiple hair fibers emerge from the same follicle (50). As with differences observed in cilia tip morphology, these multicilia may represent either normal heterogeneity or aberrant variations among islet primary cilia, as the mechanisms governing centrosome duplication and ciliogenesis are yet unclear in human islets.

Fig. 7. Microtubule configurations in the distal axoneme. In demembranated cilia, microtubules rotate and separate in the distal portion of the axoneme, creating a helical cage (A–D). Occasional actin-like thin filaments are seen coursing along the microtubules (C). Splaying of individual microtubule filaments is likely due to specimen damage (E and F). Scale, A and B, 500 nm; C–F, 300 nm.

Fig. 8. Cilia handedness in human islets. Primary cilia in human islets exhibit chirality in the microtubule bundle, examples shown in (A) from mid-axoneme where rotations begin. Quantitation among 50 membrane-extracted cilia shows that the majority have left-handed turns, while a small minority have right-handed turns or no appreciable rotation. (Scale, 300 nm.) (B) The rotational nature of islet cilia is not altered by the microtubule-stabilizer taxol; shown are two representative images each of ± 1 μM taxol, distal axoneme where rotation is most pronounced. (Scale, 200 nm.)

<table>
<thead>
<tr>
<th>Cilia helix</th>
<th># out of 50</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Left</td>
<td>41</td>
<td>82</td>
</tr>
<tr>
<td>Right</td>
<td>4</td>
<td>8</td>
</tr>
<tr>
<td>Non-rotational</td>
<td>5</td>
<td>10</td>
</tr>
</tbody>
</table>
Discussion

Our data provide the highest-resolution view to-date of human primary cilia in any tissue and organ system. Recent advances in volumetric electron microscopy including focused ion beam scanning electron microscopy (FIB-SEM) and electron tomography (ET) has allowed 3D examinations of mammalian primary cilia including in rodent islets (15, 20, 22); however, no human islet data are available from these approaches. Our study fills this important gap, combining 3D approaches to visualize cilia in situ and multiscale imaging to resolve the full ciliary axoneme. The SEM protocol works reliably for fresh human donor islets and generates images suitable for both qualitative and quantitative analyses. We did observe some attrition in cilia number during the workflow, including islet loss from coverslips, cilia breaking off during specimen preparation, and sample charging during image acquisition, which are practical matters to keep in mind when adopting this imaging method. Nonetheless, among islets from three healthy male donors, we identified a robust number of cilia including 60 with full-length axoneme, which will serve as a reference dataset for future studies of islet cilia from patients with clinical disorders. Some key limitations and implications of our work are discussed below.

An important question that we could not formally address is the heterogeneity of cilia morphology among islet alpha/beta/delta cells. The identification of islet cell types, while a simple task by light microscopy and TEM, are trickier on SEM as there are not many identifying features unequivocally established for endocrine cell groups. We attempted to classify cells by surface morphology, relying on ultrastructural features reported by previous studies of human islets (23, 51, 52). For example, beta cells are known to express abundant microvilli (53, 54), which corresponded to a characteristic velvety appearance on their surface (Figs. 2 and 9). Alpha cells, in contrast, have smoother surfaces that are largely devoid of microvilli (Figs. 2 C and 4 D). Delta cells in human islets tend to be elongated and form long filopodial extensions (55), but whether they possess microvilli and whether these correspond to the much longer villi (Figs. 4 F and 6 B) than those observed on beta cells remains a question. Cellular abundance and distribution offer helpful clues, as the surface of human islets contains intermingling beta, alpha, and delta cells in roughly 5:3:1 ratio (51). Among the 60 cilia that were sampled in entirety in our dataset, we putatively identified 37 beta, 8 alpha, and 3 delta cells based on the postulation that beta cells express uniform short microvilli, alpha cells have few or no microvilli, and delta cells are rare cells with long villi. In other cases, the surface morphology was not clearly visualized to allow cell type identification. This was a limitation in our dataset that can be addressed by future studies where true cell identity might be ascertained by immuno-SEM or correlative light and electron microscopy using hormonal or cell

Fig. 9. A ciliary ring encircles the axoneme. Ring-like structures found on human beta cell cilia, shown here from different vantage points and marked by arrowheads (A–E). Note the contrasting cilia morphology proximal versus distal to the ring, where the ring location marks where the axonemal bundle begins to unwind and twist. (B, D, and E) are high-magnification views of (A, C). Scale, (A, C) 1 μm; (B, D, and E) 200 nm.
surface protein markers. Elemental analysis using energy dispersive X-ray spectroscopy (EDS/EDX) is another possibility, which could be used to identify islet cell type by relative abundance of phosphorus, sulfur, and nitrogen, corresponding to alpha, beta, and delta cell secretory granules (23, 52).

The arrangement of actin-like connections at the ciliary base offers clues to the role of actin in cilia dynamics. The ciliary pocket has been described to be a docking site for actin cables (28, 56, 57), where interactions between the actin cortex and the cilium may serve to either stabilize cilia or generate forces to influence cilia movement (40, 58, 59). The asymmetry and sometimes one-sidedness of these fibers observed in our SEM data (Fig. 4 A–C) suggest that they may act as cables to pull the cilium via extraciliary forces, akin to a model proposed by Lippincott–Schwartz and colleagues (40). Such forces may contribute to primary cilia motility in addition to intraciliary microtubule-dynein activities (16, 40). Other potential functions of actin and associated myosin at the cilia base include regulation of ciliogenesis, establishment of cilia orientation, and regulation of endocytic uptake, as demonstrated in several experimental systems (58, 60–64). The observation of actin-like filaments along the central axonemal axis (Fig. 7 C) suggests that ciliary

Fig. 10. Diversity of cilia tip morphology in human islet cells. (A–C) Demembranation reveals a conserved central microtubule organization where the microtubule filaments coalesce and terminate at a dense tip complex. The number of microtubules present at the tip is usually between 3 and 5. (D and F) Unextracted samples with intact ciliary membrane show greater heterogeneity including ciliary bulbs and bulges. Scale, A–C, 300 nm; D–F, 400 nm.

Fig. 11. Multicilia in human islet cells. (A) Double cilia emanating from a single enlarged pocket with two parallel axonemes similar in length and diameter, both possessing defined tips. (B) Three axonemes sharing a connected base, physically uncoupled but interactive in their tip regions. (Scales, 500 nm.) Minor background distortion is seen in both images from charge build-up during scanning.
actin could be a constitutive component that provides mechanical stability to the axoneme or partitions the cilia into functional microdomains, as has been proposed for non-islet cilia (19, 39). It would be informative to test these models experimentally in islet cells.

The helical nature and chirality of axonemal microtubules was an unexpected feature that may have functional significance. Our finding that islet cilia are mainly L-handed helices contrasts with earlier reports in kidney cell cilia, which produce mild R-handed spirals (19, 21). Centriolar triplets typically exhibit an anti-clockwise spin in most tissues (65–67). If such a rotation were continued along microtubule doublets and singlets, this would indeed translate to a L-handed axoneme. While the functional difference of L- versus R-handed turns is yet unclear, we could at least conclude that cilia handedness can differ among cells, and speculatively, we may propose that it could differ among the same cell type under different growing conditions, a phenomenon well-described in the Arabidopsis plant (68). The more fundamental question is, what is the function of coiling? In contrast to tubulin protofilaments in which curved tips are thought to facilitate individual strand growth (69), collective coiling by the microtubule bundle likely contributes to overall axonemal stability and dynamics, acting as a compression shock absorber and a stretch and length modulator. With the microtubules fixed at both base and tip, the ciliary axoneme can be considered a topologically closed system, where changes in its helical conformation would allow elastic deformation without breaking strands of the microtubule bundle. In other words, the helical unwinding would be a quick way to increase ciliary length without polymerizing new subunits, and theoretically permit coil direction switching. Additionally, having loose microtubule helices in the distal axoneme would produce greater cilia surface area and luminal volume, which may alter the capacity for ciliary signaling, cargo trafficking, and exocytic vesicle release. However, as the degree of microtubule separation may be related to demembranation, it would be prudent to confirm these observations using more precise methods such as cryo-ET, which keeps tissues in a more accurate native state. Among the most intriguing observations from our dataset is the “ciliary ring,” previously not described in any mammalian primary cilia. The low frequency of the ring suggests that it may not always assemble, or survive demembranation as in our procedure. Ring complexes around individual microtubules have been reported in budding yeast (70), namely the Dam1 complex that aids in microtubule stability, but key differences are that 1) Dam1 ring thickness is about the size of tubulin protofilaments (~5 nm), thinner than that of a microtubule (25 nm), 2) Dam1 rings wrap around a single microtubule, whereas the ring observed on islet cilia wraps the entire axoneme bundle, and 3) Dam1 rings are not seen in other fungal species besides budding yeast nor in higher organisms, thus unlikely to represent the ring feature observed in this study. The ciliary ring that we describe is also distinct from the ciliary necklace and bracelet, which have different morphological and localization, for example the ciliary bracelet has only been observed in the flagella of the unicellular green alga Chlamydomonas (32). Nor does it resemble the gamma-tubulin rings reported in Xenopus, whose function is to cap the minus end and block MT growth (71). The only known ring-like structure that wraps around the entire axoneme is the membrane-bound septin ring at the clitum base (72). Septins are a family of guanosine-5′-triphosphate-binding proteins that act as diffusion barriers, and in primary cilia the septin ring is found below the transition zone. Thus location-wise, the septin ring is too low to be the structure that we observe, which is microns above the cell surface. While the composition of the ring remains unknown, we suggest based on its morphological resemblance to microtubules that this could be a tubulin-based complex that contributes to the structural integrity of the axoneme. Having a mid-axonemal ring could support microtubule bundling, promote cilia assembly, and stabilize against disassembly. Future studies might clarify the prevalence of the ring as well as regulatory factors in its formation. Whether the ring is fixed in place or could slide to dynamically control axonemal architecture or helical cage diameter is another interesting biomechanical question, but beyond the scope of this study.

Finally, the observation of compound primary cilium raises questions about their purpose and the mechanism of numerical control. The majority of cells in human pancreatic islets contain a single primary cilium, consistent with the idea that there may exist evolutionary mechanisms to ensure single primary cilium number for proper function (73). While our images clearly demonstrate the existence of double and triple cilia, we do not know their functional significance, nor can we glean from surface scans their anatomical origin, for example whether they are joined together at the basal body or accompanied by extranumerary centrioles. The Polo-like kinase (PLK) family proteins are well-studied regulators for centriole biogenesis in human cancer cell lines, with PLK4 being an established driver of centrosome duplication (74–76). While these pathways have not been experimentally studied in pancreatic islets, our survey of published human single-cell RNAseq datasets (77, 78) shows that all Plk1–5 genes including Plk4 have detectable and cell type-dependent expression in healthy human islets, suggesting that there indeed exist mechanisms for regulated cilia/centrosome duplication. The multicilia observed in our study were healthy-appearing, with normal lengths and diameters and individual ciliary membranes, surrounded by a well-formed ciliary pocket. Thus we posit that, rather than being a diseased odyssey, these multicilia may be a normal feature among islet cells, perhaps retained over evolution to maintain cilia density per unit surface area or volume. Given the heterogeneity of Plk gene expression, multiciliation may also be differentially regulated among islet cell subsets or be under hormonal control. Gain- and loss-of-function studies in the future might demonstrate a role for PLK or related factors in centrosome amplification, cilia biogenesis, and islet function.

**Methods**

**Iset Preparation for SEM.** Human islets were obtained from the Integrated Islet Distribution Program (IIDP). After retrieving islets from the shipping bottle, we washed and rested islets overnight in islet medium (RPMI 1640 with 11 mM glucose, 10% FBS, and 1% Penicillin–Streptomycin (Gibco, #15140122)). Islets were then plated on laminin-coated coverslips (treated with 0.5 μg laminin/cm² for 2 h), and allowed to adhere for 3 d. For fixation/extraction, islets were washed in successive incubations, first for 5 min in cytoskeleton buffer (50 mM imidazole, 50 mM KCl, 0.5 mM MgCl₂, 0.1 mM EDTA, 1 mM EGTA, pH 6.8) with 0.5% Triton X-100 and 0.25% glutaraldehyde, then extracted for 10 min in cytoskeleton buffer with 2% Triton X-100 and 1% CHAPS, with or without 1 μM taxol (Cell Signaling #9807). Islets were washed three times in phosphate-buffered saline (PBS), then membrane-extracted with successive incubations, first for 5 min in cytoskeleton buffer (50 mM imidazole, 50 mM KCl, 0.5 mM MgCl₂, 0.1 mM EDTA, 1 mM EGTA, pH 6.8) with 0.5% Triton X-100 and 0.25% glutaraldehyde, then extracted for 10 min in cytoskeleton buffer with 2% Triton X-100 and 1% CHAPS, with or without 1 μM taxol (Cell Signaling #9807). Islets were washed three times in cytoskeleton buffer before fixation overnight in a solution containing 2% glutaraldehyde in 0.15 M cacodylate buffer, pH 7.4 at 4 °C.

**SEM.** Coverslips containing adhered and fixed islets were transferred into 0.1% tannic acid in ultrapure water for 20 min at room temperature. Following this, samples were rinsed four times for 5 min each in ultrapure water and incubated in 0.2% aqueous uranyl acetate for 20 min. Samples were rinsed three times for 10 min each in ultrapure water and dehydrated in a graded ethanol series (10%, 20%, 30%, 50%, 70%, 90%, and 100% x 3) for 5 min in each step. Once dehydrated, samples were loaded into a critical point drier (Leica EM CPD 300), which was set to perform 12 CO₂ exchanges at the slowest speed. Samples were...
then mounted on aluminum stubs with carbon adhesive tabs and coated with 5 nm carbon and 5 nm iridium (Leica ACE 600). SEM images were acquired on a FIB SEM platform (Helios S U DualBeam Fisher Scientific, Brno, Czech Republic) using the SEM imaging mode at 1.8 kV and 0.1 nA using a TLD detector. False coloring was rendered in Adobe Photoshop. To promote data accessibility and sharing, original high-resolution TIFF images from this study have been deposited at the public database BioImage Archive [https://www.ebi.ac.uk/biostudies/Bioimages/studies, accession #S-BIAD664 (79)].

Immunohistochemistry, Confocal and Superresolution Microscopy. Isolated human islets were washed with PBS and fixed with 4% paraformaldehyde (PFA) for 15 min and permeabilized with 0.3% Triton X-100 in PBS (phosphate-buffered saline with Triton X, PBST) for 10 min at room temperature. After incubation with blocking buffer (PBS with 10% normal goat serum) for 1 h at room temperature, islets were incubated overnight at 4°C with primary antibodies diluted in PBST. Primary antibodies included mouse-anti-acetylated tubulin (Proteintech #66200-1-Ig, 1:400 dilution), rabbit-anti-ARL13b (Proteintech, 1:400 dilution), and mouse-anti-DNAC1 (NeuroMab #75-372, 1:50 dilution). The next day, islets were washed, incubated with secondary antibodies for 1 h at room temperature, and washed again in PBST. DAPI provided nuclear counterstain.

Islets were mounted on glass slides with Prolong Gold Anti-fade (Thermo Fisher P36930) and imaged using an inverted Zeiss LSM880 and Ti2 Eclipse SIM microscope on Airyscan mode (Nikon, Ti-E).

Morphometric Analysis. Linear measurements of human islet cilia length and diameter were performed in ImageJ using the straight line or freehand line measurement tool. Intact full-length cilia images were selected for analysis, excluding those that were fragmented or disrupted during sample processing. Results were compiled in GraphPad Prism and calculated for average and SD. Volume (V) and surface area (SA) calculations are derived by approximating the shape of the primary cilium to a truncated cone with base radius r1 and tip radius r2, or a right cylinder with radius r and height h, using the following equations:

\[
V = \frac{1}{3} \pi (r_1^2 + r_1 r_2 + r_2^2) h,
\]

Lateral area: \(F = \pi (r_1 + r_2) \sqrt{(r_1^2 + r_2) + h^2}\).

Surface area: \(S = 2\pi rh + r_1^2 + r_2^2\).

For a circular truncated cone:

\[
V = \pi r^2 h,
\]

\[
SA = 2\pi rh + r^2.
\]

Data, Materials, and Software Availability. Imaging data have been deposited in BioImage Archive [https://www.ebi.ac.uk/biostudies/Bioimages/studies, accession #S-BIAD664 (79)]. All study data are included in the main text.

ACKNOWLEDGMENTS. We are indebted to Dr. Ursula Goodenough along with Dr. Bob Schmidt, Robyn Roth, and the WashU Cilia Group for help with data interpretation. We acknowledge the assistance of Drs. James Fitzpatrick and Praveen Krishnamoorthy at the Washington University Center for Cellular Imaging in electron and light microscopy studies, supported by Washington University School of Medicine, The Children’s Discovery Institute of Washington University and St. Louis Children’s Hospital (CDI CORE 2015-505 and CDI CORE 2019-813), the Foundation for Barnes-Jewish Hospital (3770) and the Washington University Diabetes Research Center (DK020579). Human pancreatic islets were provided by the NIDDK-funded Integrated Islet Distribution Program (IDIP) (RRID:SCR_014387) at City of Hope, NIH grant #2U44DK090805 and the JDRF-funded IDIP Islet Award Initiative. A.J.P. is supported by the Diabetes and Related Metabolic Diseases training grant T32DK007120. Other funding included NIH grants DK115795 and DK127748 to J.W.H., Doris Duke Charitable Foundation grant DFFRC to J.W.H., and DK115972 and DK123301 to D.W. Graphs were created with BioRender.com.


