Adipogenesis and epicardial adipose tissue: A novel fate of the epicardium induced by mesenchymal transformation and PPARγ activation

Yukiko Yamaguchia, Susana Cavalleroa, Michaela Pattersona, Hua Shenb, Jian Xub, S. Ram Kumarc, and Henry M. Sucova1,1

1Broad Center for Regenerative Medicine and Stem Cell Research, 1Department of Surgery, Keck School of Medicine, and 2Center for Craniofacial Molecular Biology, Ostrow School of Dentistry, University of Southern California, Los Angeles, CA 90089

The hearts of many mammalian species are surrounded by an extensive layer of fat called epicardial adipose tissue (EAT). The lineage origins and determinative mechanisms of EAT development are unclear, in part because mice and other experimentally tractable model organisms are thought to not have this tissue. In this study, we show that mouse hearts have EAT, localized to a specific region in the atrial-ventricular groove. Lineage analysis indicates that this adipose tissue originates from the epicardium, a multipotent epithelium that until now is only established to normally generate adipose tissue originates from the epicardium, a multipotent epithelium (i.e., the epicardium), an anatomical relationship that distinguishes EAT from thoracic or paracardial fat (3). In all cases, the adipose tissue lies underneath an outer mesothelium (i.e., the epicardium), an anatomical relationship that distinguishes EAT from thoracic or paracardial fat (3). In

EAT is present in fetal and newborn stages in humans (4, 5) and other species (6), which implies that its derivation is under developmental control. In adults, there is a tendency for more EAT with increasing obesity, which led to early speculation that EAT is pathological, and “fatty heart” was a common diagnostic explanation in the 17th and 18th centuries (7). With recognition that all humans have at least some EAT, attention turned in more recent times to the possibility that EAT might serve beneficial or detrimental functions in heart metabolism, insulation, response to injury, coronary artery disease, or many other speculated possibilities (1–3, 8). A major limitation in understanding the biology of EAT is the seeming absence of this tissue in virtually all commonly used experimental animal models. Thus, rodents, avians, and amphibians are generally considered to not have EAT, although all have noncardiac thoracic (paracardial) fat (which has sometimes been mischaracterized as EAT).

The epicardium is the outer noncontractile mesothelium of the heart. It migrates onto and spreads over the heart surface during early embryonic development, which in mice occurs during the embryonic day E9.5–10.5 period (9). Once formed, the epicardium serves as a source of secreted factors that influence mitogenic expansion of the ventricular myocardium and assembly of coronary blood vessels (10). In addition, the epicardium is a multipotent progenitor cell population (11). Starting from almost the time of its formation, the epicardium undergoes transformation to generate mesenchymal cells that first reside in the subepicardial space between the epicardium and myocardium. One subset of these cells assembles around nascent coronary endothelial tubes and constitutes the smooth muscle layer of the mature coronary vessels. A separate subpopulation migrates as single cells into the myocardium and becomes the predominant source of cardiac fibroblasts that secrete extracellular matrix needed for mature heart structure (12). These two fates are well established; additional speculated fates for the epicardium, including serving as a source of cardiomyocytes and of coronary endothelium, have not been confirmed, at least under normal conditions.

In this study, we show that mice have EAT, which is normally limited to a very specific location in the heart. Using genetic fate mapping approaches, we provide evidence that this tissue is derived by mesenchymal transformation of the epicardium. We explain differences in the presence and amount of EAT between species based on deployment of the peroxisome proliferator activated receptor gamma (PPARγ) molecular pathway, and we show in mice that we can eliminate normal EAT and induce the formation of ectopic ventricular EAT, based on these insights.

Significance

A layer of fat surrounds the heart in most mammals, including humans. The biology of this tissue has been speculated for centuries, but never subjected to experimental analysis because common experimental model species are thought to not have this tissue. In this study, we show that rodents have cardiac fat, albeit in a very specific location in the heart. We implicate the origin of this tissue from the epicardium (the outer epithelium of the heart) and the underlying mechanisms that account for its derivation. By comparing human and mouse epicardial cells, we provide an explanation for the prominent species differences in the presence and amount of cardiac adipose tissue.

Author contributions: Y.Y., S.C., and H.M.S. designed research; Y.Y., S.C., M.P., and H.S. performed research; Y.Y., J.X., S.B.K., and H.M.S. analyzed data; and Y.Y., S.C., M.P., H.S., and H.M.S. wrote the paper.

The authors declare no conflict of interest.

*This Direct Submission article had a prearranged editor.

1To whom correspondence should be addressed. Email: sucov@usc.edu.

This article contains supporting information online at www.pnas.orglookup/suppl/doi:10.1073/pnas.1417232112/DCSupplemental.
contrast, adult hearts from mice and rats show no ventricular EAT, and have direct contact of the epicardium to the myocardium (Fig. S1 J–O), even in locations where coronary vessels lie (Fig. S1P). We observed that adult mouse (Fig. I4) and rat (Fig. S1Q) hearts have a depot of fat, limited to a very specific location between the atrial and ventricular chambers [the atrial–ventricular (AV) groove] on the dorsal to dorsolateral sides of the heart. Histologically, this tissue is subepicardial (Fig. 1 B and C). A second defining feature of EAT is its perfusion by coronary vasculature rather than from systemic vessels (3). By isolation of adult mouse hearts and retroaortic ink perfusion, we observed ink infiltration into small vessels of the AV groove tissue as well as into major and minor coronary vessels of the ventricle (Fig. S2 A–C). Human EAT is known to have brown adipose tissue character (1, 13), and several BAT-specific genes, as well as pan-adipose genes, were expressed in mouse AV groove fat (Fig. S2D). These three criteria imply that mouse AV groove fat tissue is EAT.

We examined the time course of appearance of AV groove adipocytes using the marker perilipin (PLIN), a protein present in lipid vacuoles of mature adipocytes (Fig. 1 D–F). Mesenchyme is particularly abundant in the AV groove at mid- and late-embryonic stages, but there were no PLIN+ cells in this region or elsewhere in the ventricle during embryonic stages or during early postnatal life. PLIN+ cells were first seen in the AV groove at 2 wk (Fig. 1E), intermixed with unlabeled mesenchymal cells. Positive cells at this time had multiple relatively small vacuoles rather than a single large vacule with minimal cytoplasm that typifies mature adipocytes, suggesting that these cells are immature adipocytes or had only recently initiated terminal maturation. PLIN+ adipocytes increased in size and maturation progressively thereafter.

We used fate mapping to address the tissue origin of mouse EAT. In the heart, a Tbx18Cre driver line labels the entire epicardium from the time of its initial formation (14). In hearts from adult mice with Tbx18Cre and the conditional reporter allele R26YFP, highly efficient labeling of the epicardium, coronary vessel smooth muscle, and interstitial cells (likely fibroblasts) was evident (Fig. S3 A–C). Inspection of the AV groove revealed complete colocalization of PLIN+ cells with the lineage marker (Fig. 1 G–I). A parallel analysis with Tie2Cre, which labels endocardium and endothelium and their derived cells (Fig. S3 D–F), did not label AV groove EAT (Fig. 1 J–L). Thus, EAT adipocytes originate from a Tbx18Cre-expressing progenitor, which is likely to be the epicardium (see Discussion).

Primary epicardial cells can be grown from embryonic heart tissue explants (15). We compared mouse and human primary embryonic ventricular epicardial cells for their propensity to undergo adipogenesis in culture, either spontaneously or in response to an adipogenesis-inducing mixture containing dexamethasone, insulin, isobutyl methylxanthine (IBMX), rosiglitazone, and palmitate (DIIR-Pal). Adipocytes were visualized by Oil Red O staining, which accumulates in lipid vacuoles. In the absence of induction, human and mouse primary cell cultures showed no sign of adipocyte differentiation. Human cells treated with DIIR-Pal for 2 wk showed prominent adipocyte differentiation, whereas mouse cells under the same conditions did not at all (Fig. 2 A–D).

Fig. 1. Tbx18Cre lineage-derived EAT is present in the AV groove of adult mice. (A) The dorsal surface of an adult (5 mo old) mouse heart. A small amount of fat (arrowheads) is visible in the AV groove. (B and C) Histological appearance of EAT. H&E-stained sections of adult mice show the presence of fat in the AV groove underneath the epicardium. The boxed regions are shown at higher magnification in B′ and C′. At, atrium; IVS, interventricular septum; Vent, ventricle. (D–F) Time course of appearance of AV groove fat. Staining with the adipocyte marker perilipin (PLIN) is absent until 2 wk after birth and increases with age. Autofluorescence (AutoFL) of the myocardium is included to delineate tissue borders. The boxed regions are shown at higher magnification in D′–F′. (G–L) Lineage origins of EAT adipocytes. The same sections are shown in the YFP (lineage marked) channel alone, the PLIN channel alone, or the merge of these and the DAPI channel. PLIN and YFP immunostaining colabel in Tbx18Cre lineage-mapped tissue; small interstitial cells of unknown fate in the AV groove are labeled by Tie2Cre but these do not overlap PLIN+ adipocytes. The sections shown in G–L are adjacent to those used in B′ and C′. The boxed regions of I and L are shown at higher magnification in I′ and L′.
This outcome corresponds with the presence and absence of EAT on the ventricular surface of human vs. mouse hearts and implies that primary epicardial cell culture could be a faithful means of studying epicardial cell differentiation in vivo. Adipocytes were also not evident when mouse epicardial cells from the embryonic AV groove were induced to differentiate (Fig. S4 A and B), showing that both subpopulations of mouse embryonic epicardial cells lack adipogenic potential. Therefore, the presence of EAT in the mouse AV groove starting around 2 wk after birth (Fig. 1E) represents a unique postnatal aspect of epicardial differentiation.

The ligand-dependent nuclear receptor PPARγ is a master regulator of adipocyte differentiation and lipid metabolism in several cell types (16). We compared cultured human and mouse primary embryonic ventricular epicardial cells and found that Pparg was expressed prominently in human cells, but at a 10-fold lower level in mouse cells (Fig. 2 E and F). Pparg was also minimally expressed in MEC1 immortalized mouse embryonic ventricular epicardial cells (Fig. 2 E and F), which also do not undergo adipogenic differentiation in response to the inducing mixture (Fig. S4 C and D). We virally expressed Pparg in primary mouse epicardial cells and in MEC1 cells; this resulted in a low degree of spontaneous adipocyte differentiation in the absence of treatment and a prominent level of differentiation in the presence of the PPARγ ligand rosiglitazone (Fig. 2 G–J and Fig. S4 E–H). PPARγ has a moderate level of ligand-independent transcriptional activity (17), which may explain the sporadic induction of differentiation when overexpressed in the absence of added ligand (Fig. 2H). Thus, the differential ability of human vs. mouse epicardial cells to initiate adipogenic differentiation can be explained by the level of endogenous PPARγ expression, and mouse epicardial cells are able to undergo adipogenic differentiation when they express PPARγ. Consistent with these observations, PPARγ was not present in the embryonic mouse heart and only became detectable in AV groove mesenchyme starting around 2 wk after birth (Fig. 2 K–M), coincident with the appearance of EAT (Fig. 1E).

To determine if PPARγ activity is required for mouse EAT formation in vivo, we crossed a Pparg conditional loss-of-function allele with Tbx18Cre. Mutant mice survived into adulthood and were seemingly normal and contained typical amounts of fat in noncardiac locations, including the kidney fat pad, intestinal omentum, and female reproductive tract (Fig. S5). In the AV groove of Tbx18Cre/Pparg conditional mutant mice, mesenchyme was still present (Fig. 3B), but mature PLIN+ adipocytes were eliminated (Fig. 3D). A small number of interstitial PLIN+ cells of uncertain identity were observed, but the absence of adipocytes was confirmed by immunostaining for fatty acid binding protein 4 (FABP4) as a second adipocyte marker (Fig. 3F). Thus, the PPARγ pathway is required for formation of mature PLIN+ adipocytes. We cannot yet say if mesenchymal cells in the AV groove in these conditional mutants are of an alternative lineage (e.g., fibroblast) that is also present in normal hearts, or if these are immature preadipocytes that have failed to mature to the point of expressing PLIN or FABP4. In either case, these observations imply that PPARγ activity does not influence the process of epicardial mesenchymal transformation, but rather the ability of mesenchyme to adopt an adipogenic fate.

Conversely, we also used Tbx18Cre to force PPARγ expression in a gain-of-function manipulation. Because an allele that conditionally expresses wild-type PPARγ is not available, for this purpose we used a conditional allele that expresses a Pax8–PPARγ fusion protein (PPFP) after recombination; the fusion protein was previously shown to behave like normal PPARγ, including the induction of adipogenic differentiation, in the presence of an appropriate PPARγ ligand (18). We first tested this genetic manipulation in primary cell culture. Primary ventricular epicardial cells derived from Tbx18Cre/PPFP embryos displayed typical epicardial morphology and showed no basal adipogenic differentiation, but underwent active adipogenic differentiation in the presence of rosiglitazone (Fig. 4 A–D). This mirrors the behavior of genetically normal primary mouse epicardial cells in which PPARγ expression is forced by viral infection (Fig. 2 G–J).
We raised control and Tbx18Cre/PPFP mice to adulthood. The amount of AV groove EAT was comparable in both (Fig. S6 A and B), and ventricular chamber EAT was absent in both even when mice were provided a high fat diet supplemented with rosiglitazone starting in adulthood (Fig. S6 C and D). Thus, forced expression and ligand activation of PPARγ (specifically PPFP) in the Tbx18Cre lineage in the adult mouse does not override the signals that support or prevent adipogenic differentiation in vivo.

The observation that Tbx18Cre/PPFP epicardial cells in culture efficiently initiate adipogenesis in response to rosiglitazone (Fig. 4 A–D) suggested that epicardial–mesenchymal transformation (EMT) might be an important step in adipogenic differentiation, as primary epicardial cell cultures are at least partially mesenchymal during early growth (e.g., as manifested by expression of vimentin and smooth muscle actin). To test this model, we cryoinjured the heart surface of adult mice, because injury is known to activate EMT in the adult heart (19). Transdiaphragmatic cryoinjury may be a preferable injury model for these studies, as primary epicardial cell cultures are at least partially mesenchymal (20).

In control mice provided with a high fat diet supplemented with rosiglitazone, the injured surface of the heart was covered with an extensive fibrotic matrix. This scar tissue lacked adipocytes, although a small localized cluster of adipocytes was typically observed in the injury-adjacent area (Fig. 4 E and G and Fig. S6 E and F). The observation that this occurs at all implies that injury is accompanied by some degree of relaxation in the negative control of adipogenesis, and presumably also in expression of PPARγ. In contrast, in injured Tbx18Cre/PPFP, mouse, prominent accumulations of adipocytes were present in the injury-adjacent region (Fig. 4 F), although fibrotic scar tissue was still extensive. Quantitation indicated a more than 50-fold increase in the amount of postinjury EAT in Tbx18Cre/PPFP hearts compared with littermate controls (Fig. 4 G). A layer of epicardium identified by the marker podoplanin overlaid the induced ventricular EAT (Fig. S6 G and H), which was immunolabeled with both PLIN and FABP4 (Fig. S6 H and I). Mesenchyme induced by adult heart cryoinjury therefore almost completely adopts a fibrotic fate in control mice, whereas the expression of PPARγ (PPFP) promotes or allows significantly more adipogenic differentiation.

Mesenchymal transformation of the epicardium occurs only sporadically if at all during normal postnatal life, but occurs actively during normal embryonic development. We therefore treated mice with rosiglitazone in utero during the period when this process is actively underway. When examined in adulthood, rosiglitazone exposure of control embryos had no effect (Fig. 4 H and I), whereas in Tbx18Cre/PPFP embryos, this treatment resulted in broad domains of ventricular fat in the subepicardial space (Fig. 4 J–L and Fig. S6 J and K). The morphology of the adult heart (Fig. 4 J) and the viability of these mice into adulthood was not compromised, which implies an adequate level of differentiation by the embryonic epicardium to coronary vascular smooth muscle and fibroblast fates, even though a subpopulation of the epicardium-derived mesenchyme underwent adipogenic differentiation.

**Discussion**

Our results demonstrate that rodents normally have EAT in the atrial–ventricular groove and support the conclusion that this tissue is derived from the epicardium. Very recently, two reports reached related conclusions. Using Wt1Cre as a lineage marker, Chau et al. (20) demonstrated that all visceral fat, including epicardial fat, is derived from the mesothelium; as noted above, the epicardium is the mesothelium of the heart. Liu et al. (21) described mouse AV groove EAT as we observed, and using MshnCre and Wt1Cre, concluded derivation from the epicardium. Our analysis with Tbx18Cre, bolstered with our demonstration that epicardial cells can become adipocytes in vitro, complements these recent reports. Although any single Cre driver can have experimental caveats related to efficiency and specificity, the aggregate data show that adipogenic differentiation is a third and new fate of the multipotent epicardium. In both of the recent reports, the experimental approaches resulted in only moderately efficient epicardial recombination, so the possibility of a mixed derivation could not be addressed. Tbx18Cre is highly efficient in the epicardium, and we observed virtually complete labeling of AV groove EAT with this lineage marker (Fig. 1 G–I), indicating that the epicardium is likely to be the sole source of EAT adipocytes. However, it should be cautioned that additional nonepicardial sites of Tbx18Cre expression (22, 23) could in principle contribute cells to this tissue.

Collectively, our cell culture and in vivo results indicate that mouse epicardium-derived cells can adopt an adipocyte fate after mesenchymal transformation and if they express PPARγ. The requirement for PPARγ is not unexpected, as adipogenic differentiation requires PPARγ activity (24). Our conclusions related to the requirement for EMT rest on several observations: adult heart injury induces EMT and induces fat when PPARγ is expressed (Fig. 4 F); an adipocyte fate is induced in embryonic...
heart during the period when normal EMT is underway (Fig. 4 J–L); epicardial cells in culture are partially mesenchymal and this unleashed adipogenic potential when they express PPARγ (Figs. 2 G–J and 4 A–D); and finally, adipocytes are a mesenchymal cell type and if derived from the epicardium, it is almost inescapable that EMT must occur before adipocyte differentiation can occur. Formally, however, definitive confirmation of this model would require experimental manipulation to prevent EMT and then determine the consequence to adipogenesis. Liu et al. (21) also noted induction of ventricular fat after adult heart injury, although interestingly, their observations were conducted in mice and four control littermates (bearing only Tbx18Cre, or only PPFP, or neither allele).

**Fig. 4.** PPARγ activity and mesenchymal transformation induce adipogenesis in vivo. (A–D) In vitro induction of adipogenesis by PPFP. Primary embryonic ventricular epicardial cells were derived from Tbx18Cre/PPFP embryos and control littermates, treated with rosiglitazone, and lipid accumulation visualized by Oil Red O staining. (E and F) In vivo adult adipogenesis. Cryoinjury was performed on Tbx18Cre/PPFP adult mice and controls; a high fat diet and rosiglitazone were provided for 3 mo after surgery. In transverse sections near the apex, ventricular adipocytes were detected by PLIN immunostaining and DAPI counter-staining. Images of uninjured hearts are in Fig. S6 C and D. The boxed areas at the transition between normal myocardium (myo) and scar tissue are shown in E, E′, F, and F′; E and E′, and F and F′, are adjacent or nearby sections. (G) Quantitation of ventricular fat. The volume of ventricular fat per heart was derived from PLIN- and Oil Red O-stained serial sections. There was no fat in uninjured control (n = 4) or uninjured Tbx18Cre/PPFP (n = 5) hearts, a small amount in injured controls (n = 4), and substantially more (*P = 0.0004) in injured Tbx18Cre/PPFP (n = 5) hearts. Fat was only present in injury-adjacent areas. (H–L) Induction of embryonic adipogenesis. Pregnant females were fed a high fat diet and provided with rosiglitazone starting at E10.5 and continuing during lactation, and weaned pups were kept on the same regime until 3 mo of age. The H&E images (H and I) show normal morphology of the treated adult hearts; immunostaining for PLIN (L, K, and L) shows the presence of adipocytes in two different locations of the ventricle in Tbx18Cre/PPFP mice but not in controls. These observations were repeated in six Tbx18Cre/PPFP mice and four control littermates (bearing only Tbx18Cre, or only PPFP, or neither allele).
such that most of the right dorsal ventricular wall and a part of the dorsal left ventricular wall were in direct contact with the diaphragm. A 0.5-mm metal blunt probe precooled in liquid nitrogen for 1 min was inserted through the sleeve and pushed against the heart and diaphragm for 10 s. Sham-operated mice received the same procedure but with a room temperature probe.

High Fat Diet and Rosiglitazone Administration. A single daily dose of rosiglitazone (10 mg/kg body weight) was administered intraperitoneally immediately after surgery and for the next 3 d. 1 μM rosiglitazone was also coadministered in the drinking water. The high fat diet was purchased from Harlan Laboratories (TD06414). The water and diet were provided for 3 mo from the day of surgery until tissue harvest. To administer rosiglitazone in utero, a single daily dose of 10 mg/kg body weight was administered intraperitoneally to pregnant females from E10.5 to E13.5. The females were provided ME with a high-fat diet and with drinking water containing 1 μM rosiglitazone from E10.5 until pups were weaned. Pups continued to receive the water and high fat diet until tissue harvest at 3 mo of age. Control mice were littersmates of the experimental mice and therefore were exposed to the same conditions.

Quantification of Fat Volume. Serial sections of samples were generated and stained with Oil Red O. Sections were then photographed, and Oil Red O stained tissue was quantified using ImageJ. Volume was calculated, and Student’s t test was performed for statistical significance.

ACKNOWLEDGMENTS. The authors acknowledge the technical assistance of Mr. Danny Lee in these studies. This work was supported by NIH Grant HL070123 and by a pilot award from the University of Southern California Diabetes & Obesity Research Institute (to H.M.S.). Y.Y. was supported by a predoctoral fellowship from the American Heart Association and H.S. was supported by a postdoctoral fellowship from the California Institute for Regenerative Medicine.