Exosomal microRNA communication between tissues during organogenesis

Toru Hayashia and Matthew P. Hoffmanb

aDepartment of Anatomical Science, Kitasato University School of Allied Health Sciences, Sagamihara, Kanagawa, Japan; bMatrix and Morphogenesis Section, National Institute of Dental and Craniofacial Research, National Institutes of Health, DHHS, Bethesda, Maryland, USA.

ABSTRACT

Epithelial-mesenchymal interactions are required to coordinate cell proliferation, patterning, and functional differentiation of multiple cell types in a developing organ. This exquisite coordination is dependent on various secreted molecules that provide developmental signals to mediate these tissue interactions. Recently, it was reported that mature mesenchymal-derived microRNAs (miRNAs) in the fetal mouse salivary gland are loaded into exosomes, and transported to the epithelium where they influence progenitor cell proliferation. The exosomal miRNAs regulated epithelial expression of genes involved in DNA methylation in progenitor cells to influence morphogenesis. Thus, exosomal miRNAs are mobile genetic signals that cross tissue boundaries within an organ. These findings raise many questions about how miRNA signals are initiated to coordinate organogenesis and whether they are master regulators of epithelial-mesenchymal interactions. The development of therapeutic applications using exosomal miRNAs for the regeneration of damaged adult organs is a promising area of research.

CONTACT Toru Hayashi htoru88@kitasato-u.ac.jp

Intercellular communication with RNAs

Intercellular communication is an essential feature required for multicellular organisms.1 The communication often involves various proteins, such as soluble factors, their receptors, and secreted extracellular matrix.1 RNA was initially not considered a mediator of intercellular communication because of its instability and rapid degradation by endogenous RNases.2 However, since the late 1990s RNA was discovered to have functional roles as a mobile signal across cell and/or tissue boundaries in multicellular organisms,3-5 including plant, nematode, and insect cells.1,2,6-8 In plants, RNA transport was demonstrated by grafting tissue from a genetically modified plant into a control plant, allowing the transgene signals to be tracked over long distances through phloem.4,9 The report of RNA-transport and RNA-interference (RNAi) in Caenorhabditis elegans (C. elegans) were awarded the 2006 Nobel Prize in Physiology or Medicine.5 C. elegans expresses SID-1, an RNA channel, to import double-stranded RNA (dsRNA) into cells.8,10,11 and the dsRNA is also transported from cell to cell throughout the whole body and surprisingly from one generation to the next.8,12

An important question is whether RNA transfer has been implicated in vertebrate organs. In 1974, experiments were performed to examine whether RNA transfer is transferred between lung tissues in embryonic chick and mouse.13 Isolated lung mesenchyme was incubated with ribonucleosides labeled by stable heavy isotopes. Then the mesenchyme was cultured with unlabeled-epithelium separated with a filter. Subsequent equilibrium centrifugation of RNA from the epithelium showed no detectable transfer of macromolecular RNA such as messenger miRNAs (miRNAs).13 However the extensive transfer of labeled ribonucleosides was detected from mesenchyme to epithelium with subsequent incorporation into epithelial RNA, in spite of thoroughly washing the mesenchyme to remove the remaining labeled ribonucleosides.13 The authors speculated that this radioactivity was noise, comprised of heterogeneous RNA including some small RNAs.13 Therefore the “noise” detected in the unlabeled-epithelium may have been functional small RNAs transported from the mesenchyme in the lung.

Similarly, a report in 2012 may have implied that micro-RNAs (miRNAs) were transported between tissues in the mouse mammary gland.14 miRNAs are small noncoding regulatory RNAs that involved in various biologic processes and function by silencing mRNA expression at post-transcriptional levels.15-17 The authors eliminated the miR-17/92 cluster from genomic DNA specifically in mammary epithelial cells and assessed the functional roles of this miRNA cluster during mammary gland development. Unfortunately, the mammary epithelium showed no defects during development from puberty to lactation. Therefore, the authors concluded that the miR-17/92 cluster was dispensable for mammary gland development, and suggested that other miRNA paralogues may compensate for the miR-17/92 cluster in the mammary epithelium.14 In this study, it is interesting to note that miRNAs in the miR-17/92 cluster were still detected in the mammary epithelial cells, contrary to an expectation that they were specifically deleted in the mammary epithelium.14
argued that the unexpected miRNA detection was due to other cells contaminating the epithelial cell purification. However, it may be possible that the miRNAs were transported to the mammary epithelium from surrounding tissues such as mesenchyme and fat pad, although the authors did not suggest the possibility. Taken together, these studies using lung and mammary gland, in hindsight, suggest small RNA transfer may occur between different tissues within vertebrate organs.

**RNAs in extracellular vesicles**

Last decade, a major advance in understanding intercellular communication in mammalian cells was the discovery that miRNAs are involved. This showed that cells *in vitro* secreted extracellular vesicles (EV) containing functional miRNAs in a stable form as a cargo and that the EV were delivered to other cells. Later studies reported that miRNAs packaged in EV were also found in body fluids *in vivo*, which was promising for clinical applications, such as liquid biopsy. EV are categorized into three groups: apoptotic bodies of 1–4 μm in diameter, which are generated from plasma membrane when apoptosis occurs in damaged cells; microvesicles of 200 nm-1 μm in diameter, which are formed by outward blebbing of plasma membrane; and exosomes (30 to 100 nm in diameter), which are formed by invagination and budding of late endosome membranes and released by fusion of multivesicular bodies with the plasma membrane. To date, open source resource for EVs catalogs RNAs and other biomolecules that detected in these EVs (Vesiclepedia; http://www.microrna.org). As of July 2017, there were 4,934 entries for miRNAs, 27,642 entries for miRNAs, 584 entries for lipids, and 92,897 entries for proteins including development signals such as WNTs and Hedgehog proteins. Among these EVs, exosomes have been of particular interest since the first discovery that RNAs are “signaling molecules” in *cell-to-cell* communication via exosomal secretion and uptake.

**Epithelial-mesenchymal interactions in embryonic salivary gland**

It has been suggested that the miRNAs in exosomes are used as mobile signals for *cell-to-cell*, and even for fetal-maternal communication. However, most studies have examined exosomal miRNA transport systemically by injection of exogenous exosomes *in vivo* or by using cultured cell lines *in vitro*, which may be different from physiologic conditions. Recently, we demonstrated that endogenous miRNAs are transported via exosomes between tissues during mammalian organogenesis. We used *ex vivo* culture of fetal ICR mouse submandibular glands (SMG) at embryonic day 13 (E13), which has been a well-studied model organ to study epithelial-mesenchymal interactions. The E13 SMG is easily cultured in minimal media and serum-free conditions *ex vivo* and undergoes branching morphogenesis, which is robust and consistent (Fig. 1A). The E13 SMG is composed of a condensed neural crest-derived mesenchyme containing an epithelial rudiment, which can be physically separated and recombined in *ex vivo* culture, to enable us to treat each tissue independently with reagents and recombine them after the treatment. Additionally, the isolated epithelium can be cultured in the absence of mesenchyme and undergoes branching morphogenesis as long extracellular matrix and growth factors are provided.

Classic recombination experiments with epithelium and mesenchyme from different organs suggest that the embryonic SMG mesenchyme plays an instructive role for epithelial branching and differentiation. Thus, it has been accepted that the SMG mesenchyme provides mobile signals to the epithelium leading to a change in epithelial progenitor cell fate, although all the signals of the tissue-tissue interactions have not been defined (Fig. 1B). We previously profiled the miRNAs expressed in both SMG epithelium and mesenchyme. Since recent studies suggested that exosomal miRNAs mediate intercellular communication, we therefore investigated whether the E13 SMG in culture releases exosomes containing miRNAs and discovered that they act as mobile genetic signals to mediate epithelial-mesenchymal interactions.

**Exosomal miRNAs are transported from mesenchyme to epithelium in fetal SMG**

We cultured E13 SMG without serum on top of floating filters with a 100 nm pore size for 48 hrs and collected the conditioned medium. The filter pore size may help to prevent transfer of large extracellular vesicles secreted to the conditioned medium. Sequential centrifugations were performed to remove cell debris and larger EV such as apoptotic bodies, and microvesicles. Finally, an exosome-containing pellet was acquired by ultracentrifugation and analyzed by transmission electron microscopy to observe the size of the exosomes, western blot to confirm the presence of exosome markers, and electrophoresis for RNA detection. Together, our results confirmed that E13 SMG produce exosomes carrying small RNAs.

To examine if exosomes are transported between epithelium and mesenchyme, we recombined tissues labeled with fluorescent BODIPY-ceramide, which labels cell membranes and exosomes. We observed ceramide-signals in the non-labeled epithelium but the primary transcript of the miRNA was not, although all the signals of the tissue-tissue interactions have been defined (Fig. 1B). We previously profiled the miRNAs expressed in both SMG epithelium and mesenchyme. Since recent studies suggested that exosomal miRNAs mediate intercellular communication, we therefore investigated whether the E13 SMG in culture releases exosomes containing miRNAs and discovered that they act as mobile genetic signals to mediate epithelial-mesenchymal interactions.

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but was in the mesenchyme, suggesting that the mature miR-133b-3p is processed and transported from the mesenchyme to the epithelium. Furthermore, tissue recombination experiments with brefeldin A treatment to suppress exosome release, reduced mature miR-133b-3p levels in the epithelium, further supporting that transmission of mature miR-133b-3p occurs from mesenchyme via exosomes across the tissue boundary. The other 80 exosomal miRNAs could be also transported to the epithelium, and analysis of all their primary and mature forms remains to be investigated. Importantly, miRNA transport was observed and measured without using any exogenous or transgenic overexpressed miRNAs, suggesting that the miRNA transport occurs under physiologic condition during SMG organogenesis (Fig. 1C).

We showed that miRNA mobile signals from the mesenchyme interact with the acceptor tissue, e.g., the SMG epithelium, although we cannot exclude the possibility that miRNA transport occurs in the other direction, from epithelium to mesenchyme. Recent reports showed that EVs contain other non-coding small RNAs such as piRNA, snRNA, snoRNA, and tRNAs as well as miRNAs. Furthermore it has been shown that there are various ways to transport RNAs between cells such as nanotube tunnels, gap junctions, and intercellular bridges formed by incomplete cytokinesis, suggesting that cells within complex tissues also communicate with each other with potentially multiple ways to transport RNAs.

**Regulation and functional roles of exosomal miRNAs during SMG development**

Given that SMG epithelium is composed of heterogeneous cells and multiple cell layers, some open and important questions remain. Which epithelial cells can accept the mesenchymal exosomes? Our data suggested uptake of exosomal miR-133b-3p from mesenchyme by epithelial cells expressing KIT that is a progenitor marker of end buds. This was demonstrated by knockdown of miR-133b-3p in epithelium, resulted in both reduction of end bud morphogenesis and an increase of DIP2B, a target of miR-133b-3p, in KIT \(^{C} \) cells. However, KIT \(^{C} \) cells in an endbud are heterogeneous, i.e., KIT \(^{C} \) K5 \(^{C} \) cells, KIT \(^{C} \) K14 \(^{C} \) cells, and KIT \(^{C} \) K19 \(^{C} \) cells. Additionally, KIT \(^{C} \) cells are distributed throughout an end bud, indicating that some KIT \(^{C} \) cells are located in more peripheral regions while others are in the center of the end bud. Therefore, which KIT \(^{C} \) cell populations accept the exosomal miR-133b-3p in vivo remains unclear.

Another remaining question: how can the exosomes target KIT \(^{C} \) cells or potentially subset of KIT \(^{C} \) cells? There may be two possibilities for uptake of the exosomes in KIT \(^{C} \) cells. First, it was shown that basement membrane (BM) in E13 SMG was perforated by numerous microscopic holes at tips of the expanding end buds (KIT \(^{C} \) cells). The holes (average 1.6 \( \mu \)m \(^{2} \) in diameter) at the end buds are potentially large for exosomes to pass through. Importantly, the microperforations were absent in duct areas. Therefore, if the mesenchymal exosomes are mainly transported through the holes to epithelium in E13 SMG, holes at tips in endbuds may allow KIT \(^{C} \) cells to accept mesenchymal exosomes. Interestingly, BM in E11 lung and kidney was also perforated by holes, suggesting that epithelial-mesenchymal interaction with exosomes could also occur in these organs.

Second, adhesion molecules in exosomes might mediate exosome uptake in KIT \(^{C} \) epithelial cells. It was shown that human tumor cells secrete exosomes with integrins that are
selectively packaged and target to extracellular matrix-enriched cellular areas in distant organs, suggesting that unique exosomal integrins may selectively adhere and establish a niche for sites of future metastasis. In fetal mouse SMG, various integrins are expressed in epithelium and mesenchyme. Although it is not examined whether the mesenchymal exosomes in E13 SMG contain integrins as a cargo, these adhesion molecules in the exosomes may mediate exosome uptake in KIT+ epithelial cells that express integrins. For example, Itga4 expression levels are quite different between KIT+K5+ cells and KIT+ cells at E15 SMG (sgmap.nidcr.nih.gov), suggesting that subset of KIT+ cells express distinct integrins. Taken together, both microperforation at end buds in BM/or and differential integrin expressions in KIT+ cells could affect where mesenchymal exosomes are delivered.

What is the role of exosomal miRNAs on differentiation during organ development? A major function of the fetal mesenchyme that has been proposed is to instruct the epithelium. Exosomal miRNAs from the mesenchyme may regulate epithelial differentiation as well as morphology, and thus play a major instructive role in early epithelial development. Functional differentiation with the occurrence of proacinar cells and secretory products begins in vivo at E15 and continues to birth. Interestingly, microperforations in the BM were associated with early developmental stages and were not detected at E15. Thus, exosome transport may be reduced at E15, which could potentially be one of triggers for epithelium to begin differentiation, which raises the possibility that miRNAs are an effective way to repress differentiation of the epithelium during branching morphogenesis. Given that gene expression during development is regulated by multiple growth factors, the transcriptome of the immature epithelium must respond and integrate multiple growth, maintenance and differentiation cues simultaneously, using cassettes of genes in gene regulatory networks (GRNs). As reported, exosomal miR-133b-3p targets Disco interacting protein 2 (DIP2B), which is associated with methylated genomic DNA in hereditary chromatin (Fig. 1C). In vivo, may influence GRNs at the epigenetic level, rather than by downregulating the expression of an individual gene within a GRN. Thus, exosomal miRNAs downregulate expression of the epithelial epigenetic machinery to allow expansion of undifferentiated epithelial progenitor cells at early stages of development during growth and branching until the epithelium is ready for further cell differentiation.

The exosomal miRNAs from the mesenchyme may also play a role in specifying epithelial cell fate. This idea is suggested by tissue recombination experiments of either embryonic mammary or pituitary epithelium with embryonic SMG mesenchyme (C3H/HeM mouse), showing that the mesenchyme was instructive. The mammary epithelium at E16 was recombined with SMG mesenchyme at E14, and the mammary epithelium resembled SMG in morphology, although functional differentiation of the epithelium was similar to mammary gland. The exact mesenchymal signals that induce a change of mammary epithelial morphology are still unknown. Likewise, in the pituitary epithelium at E9-E10 (C3H/HeM mouse), the E14 SMG mesenchyme induced α-amylase and adrenocorticotrophic hormone expression, indicating that the SMG mesenchyme induced both salivary- and pituitary-specific differentiation. Distinct tissue-specific transcriptomes and/or GRNs could be responsive to common epigenetic regulators that specify or direct epithelial cell fate. Thus, embryonic SMG mesenchyme-derived exosomal miRNAs could direct a tissue-specific transcriptome toward SMG-like transcriptome, which might result in SMG morphogenesis or/differentiation.

Communication with exosomal miRNAs is also associated with salivary gland dysfunction. Sjögren’s syndrome (SS) is a systemic autoimmune disease of the exocrine glands, such as salivary and lacrimal glands. It has been reported that miRNA expression profiles in minor salivary glands were different between SS patients and healthy donors, and saliva from SS patients contained distinct exosomal-miRNAs. Interestingly, ebv-miR-BART13-3p, a miRNA expressed by the Epstein Barr Virus (EBV), was detected at high level in salivary glands of SS patient despite the EBV usually infects B cells. A following study revealed that the ebv-miR-BART13-3p was transported from the infected B cells to salivary epithelial cells via exosomes, and downregulated expressions of a protein involved in Ca2+ signaling, stromal interacting molecule 1 (STIM1), suggesting a functional correlation between the ebv-miR-BART13-3p and impairment of saliva production in the SS.

Exosomal miRNA-based diagnostics and therapies

Since EVs, including exosomes, are circulating in body fluids and reflect physiologic states of donor cells, they have been a focus of active research for use in diagnostics, understanding pathogenesis and for therapeutics (Fig. 2A). Cancer cells secrete exosomes, which act as messengers to communicate with themselves as well as other normal cells. In fact, melanoma cells secrete exosomes, microvesicles, and apoptotic bodies during culture. The EV subsets in the conditioned medium of melanoma cells were isolated by differential centrifugation and then small RNA sequencing detected 252 miRNAs in all samples of which 23 mRNAs were unique to exosomes. Another 125 miRNAs in exosomes were also detected in both/either microvesicles or apoptotic bodies, suggesting that many identical miRNAs were loaded in all EVs secreted by cancer cells. These data highlight that in the discovery and characterization phases of defining miRNA signatures of a particular cancer or cell type, standardization of exosome isolation methods will be required to avoid contamination of other EVs that may lead to erroneous interpretation.

It will be important to determine whether exosomes could enhance regenerative therapy. Tissue regeneration using either gene or cell-based therapy has been proposed as therapy to restore damaged-organ function including SMG (Fig. 2A). Exosomes could protect miRNA cargo from endogenous RNases, and allow the miRNAs to reach their target cells. In addition, exosomes usually contain CD47, an integrin-associated transmembrane protein, which plays a functional role to protect cells from phagocytosis as in a “don’t eat me” signal. It was reported that the half-life of exosomes overexpressing CD47 in circulation was longer than exosomes without CD47, which facilitated therapeutic effects in pancreatic cancer. Therefore, either exogenous exosomes engineered with CD47 and artificial miRNAs or
endogenous exosomes carrying miRNAs from an organ tissue could be useful for therapeutic applications during organ regeneration.

The advantage of using miRNA is that multiple genes or critical genes within a GRN may be targeted, and thus have a broader effect on a specific biologic process rather than on a single gene/signaling pathway (Fig. 2B).62 Also, miRNAs could be used to redirect the transcriptome in acceptor cells,48 tissues in vitro and organs in vivo, which could result in functional regeneration. However, a single dose of exosomes may have only transient therapeutic effect.63 Therefore, the identification of exosome-secreting cells may be useful, to either isolate exosomes from culture media, or to use the cells directly, in combination with a stem/progenitor cell for cell-based therapy (Fig. 2A). Indeed, mesenchymal stem cell (MSC)-based therapies are at the forefront of regenerative medicine, and accumulating evidence suggests that the regenerative effects of MSCs are at least in part due to extracellular vesicles that may contain miRNA.63,64

Conclusions

We have mainly discussed how the exosomal miRNAs as developmental signals participate in epithelial-mesenchymal interactions during SMG organogenesis. Identifying miRNAs that regulate the organogenesis provides potential therapeutic targets for tissue repair or regeneration of damaged adult organs. Future challenges include determining how many copies of particular miRNAs are in exosomes and how this is controlled,65 how the exosomal receptors target miRNA delivery to recipient epithelial cells, what biologic significance individual miRNA signals play in vivo organogenesis, whether functional redundancy occurs, whether other developing organs also communicate with exosomal miRNAs between tissues, and whether there is tissue-specificity of miRNAs transported via exosomes in other developing organs that orchestrate GRNs required for epithelial-mesenchymal interactions.
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