Review

Skin cell-derived extracellular vesicles: a promising therapeutic strategy for cutaneous injury

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Abstract

Wound healing refers to the healing process that occurs after the skin and other tissues are separated or damaged by internal or external forces. It is a complex combination of tissue regeneration, granulation tissue hyperplasia, and scar formation, and shows the synergistic effects of these processes. After skin damage, the environment around the wound and the cells at site of the damage respond immediately, and a range of cytokines and growth factors are released. In cutaneous injury, extracellular vesicle (EV) signaling plays a vital role in the healing process via paracrine and endocrine mechanisms. EVs are natural intercellular and inter-organ communication tools that carry various bioactive substances for message exchange. Stem cells and stem cell EVs facilitate tissue repair, showing promising potential in regenerative medicine. Nevertheless, EVs derived from specific skin tissue cells, such as epidermal cells, fibroblasts, vascular endothelial cells and inflammatory cells, also play important roles in cutaneous tissue repair. Here, we describe the characteristics of wound healing, concentrating on the production and functions of EVs derived from specific skin cells, and provide new ideas for wound therapy using EVs.

Key words: Extracellular vesicles, Specific skin tissue cell, Cutaneous injury, Wound healing

Highlights

• This article reviews EVs secreted by cells residing in skin tissue and their roles in wound healing.
• EVs derived from skin tissue cells reflect cell–cell cross-talk status in the microenvironment during the healing process.
• EVs provide new insights into the treatment of cutaneous injury and the current challenges, as well as pending questions in EVs application are also involved.
Background

The skin is a multilayered primary defense organ, comprising various cell types that intercommunicate to make it an effective barrier. Intimate intercellular communication in the skin is necessary for effective surveillance. The skin is the most vulnerable when exposed to the outside environment. Cutaneous wounds can be caused by trauma, burns and chronic diseases, and pose a great therapeutic challenge worldwide [1,2]. After cutaneous injury, the cells and microenvironment at the wound site respond rapidly to promote healing. Dynamic cellular events following cutaneous injury rely on bidirectional cell–cell communication for efficient wound healing. As intercellular mediators, skin cells transmit signals via immediate contact or release of soluble factors, including proteins, genetic material, such as microRNAs (miRNAs), and extracellular vesicles (EVs), transporting bioactive molecules to adjacent and distant cells for communication and organ regulation [3]. To some extent, EV-borne molecular signals drive the cross-talk between different cellular compartments in a way that directly determines the wound healing outcome. EVs originating from specific cells are being explored for their involvement in intercellular skin communication. In this review, we focus on the role of EVs secreted from different cells residing in skin tissues during the wound healing process. Furthermore, we summarize the recent research progress on EVs and emphasize their clinical relevance to facilitate the development of novel therapeutic strategies.

Review

Wound healing

Wound healing is a highly coordinated process in which numerous cell types are extensively mobilized to restore the injured tissue. It occurs via distinct overlapping phases: hemostasis, inflammation, proliferation and remodeling. In the early stages of the wound, there are varying degrees of tissue necrosis, blood vessel rupture and bleeding in the wound area. Within a few hours, an inflammatory response occurs, which is followed by local redness and swelling. Neutrophils predominate during early leukocyte infiltration, releasing toxic granules, producing an oxidative burst, initiating phagocytosis, and generating neutrophil extracellular traps (NETs) to eliminate infections [4]. Macrophages predominate approximately three days later. Fibrinogen promptly solidifies to form clots that stop the bleeding and protect the wound. Subsequently, the skin and subcutaneous tissue around the edges move toward the center to shrink the wound. Granulation tissue then grows from the bottom and edges of the wound to fill it, collagen fibers proliferate to form a scar. Eventually, the epidermis is re-epithelized [5].

During the entire repair process, various cell groups cooperate, contributing to the coordination of individual events and facilitating temporary and spatial control [6]. Monocytes, platelets, inflammatory cells, epidermal cells and dermal cells interact with each other to promote wound healing. Macrophages are conducive to cell growth, differentiation, extracellular matrix (ECM) formation, and remodeling by serving as reservoirs for the release of growth factors [7]. During transition from the inflammation phase to the granulation tissue phase, cellular interactions gradually become dominated by the interplay between keratinocytes and fibroblasts. Epidermal keratinocytes can interact with dermal fibroblasts either via soluble mediators or secreted vesicles that carry signaling molecules to transfer information between the two cell types [8]. Hence, the whole repair process is not a single-player process but requires cell-cell cross-talk and interaction between distinct cell types, in which EVs are most likely to be responsible for pathophysiologic message exchange.

Notably, the epithelial–mesenchymal transition (EMT) process during which epithelial cells acquire mesenchymal fibroblast-like characteristics, is an indispensable event involved in both wound healing and scar formation. Generally, EMT includes three diverse subtypes: type-1, type-2 and type-3 EMT. Type-1 EMT is associated with embryogenesis and organ development. Type-3 EMT occurs in neoplastic cells and promotes the growth of localized tumors via genetic and epigenetic modifications. The EMT linked to wound healing, tissue regeneration and organ fibrosis is classified as type-2 EMT [9]. In the early proliferative phase of wound healing and tissue repair, fibroblasts migrate into the wound and secrete collagen-rich ECM to stimulate granulation tissue formation. In the late period of the proliferative phase, fibroblasts are activated and differentiated into contractile myofibroblasts to promote wound contraction. Type-2 EMT, to some extent, can be understood that non-fibroblasts have the capability of trans-differentiation into fibroblasts and myofibroblasts with the profibrotic and proinflammatory activity to participate in wound healing and tissue repair. Type-2 EMT, to some extent, can be interpreted as that non-fibroblasts have the capability of trans-differentiation into fibroblasts and myofibroblasts with the profibrotic and proinflammatory activity to participate in wound healing and tissue repair. However, myofibroblasts act as double-edged swords, excessive ECM secretion can cause adverse scarring [10]. After the acute and moderate injury, the healing event is considered as remedial action. In the continuing chronic injury and inflammation, anomalous generation of myofibroblasts stimulates a progressive fibrosis, leading to the structure and function destruction of organ and tissue [11]. It is suggested that anti-EMT strategies could be used for antifibrotic/scar therapy during the specific period of repair.

Extracellular vesicles

EVs are lipid bilayer-encapsulated nanoparticles with a size of 50–1000 nm [7] that can be released by all cell types and found in various body fluids [12], including blood, urine, saliva, breast milk, amniotic fluid, ascites, cerebrospinal fluid, bile and semen [13]. Increasing evidence indicates that they play an essential role not only in the regulation of normal physiological processes, such as stem cell maintenance, tissue repair, immune surveillance and blood coagulation, but also in the pathology of several diseases [14–17].
Subpopulations and biogenesis EVs cover various subtypes of cell-released membranous structures. The classification standards for EVs subtypes have not yet been unified. According to MISEV2018 [18], EVs can be divided into medium/large EVs (>200 nm) and small EVs (sEVs < 200 nm) based on their physical properties. EVs can also be classified based on their origin (e.g. apoptotic bodies with 50–1000 nm diameter, microvesicles with 100–1000 nm diameter, exosomes with 40–160 nm diameter, and average 100 nm diameter) or biological composition (such as surface protein CD63+). Additionally, prevailing conditions are used to distinguish EVs as large oncosomes, hypoxic EVs and podocyte EVs. There are many EVs whose functions and composition have not yet been characterized [19,20].

Exosomes are widely studied, classical exosomes formed by the endosomal system that are formed via three stages: the plasma membrane of the cell invaginates for the first time to form endocytic vesicles, multiple endosomes fuse to form early endosomes, and intraluminal multivesicular bodies (MVBs) are formed in the early endosomes that enclose the intracellular molecules. Lastly, MVBs fuse with the plasma membrane, releasing intraluminal vesicles (ILVs), namely exosomes, into the extracellular space [21]. Endosomal sorting complex required for transport (ESCRT) drives the formation of MVBs and ILVs. ESCRT comprises approximately 30 proteins that assemble into four complexes (ESCRT-0, -I, -II and -III), which are linked to cargo recruitment and inward invagination of the late endosomal membrane [22]. ESCRT-0 complex recognizes and sequesters ubiquitinated transmembrane proteins in the endosomal membrane, whereas the ESCRT-I and II complexes are responsible for membrane deformation into buds with sorted cargo. ESCRT-III can be activated by ESCRT-II and subsequently drives vesicle scission [23]. In addition, ESCRT-independent pathways also regulate the inward budding of the MVB. Sphingomyelinase hydrolysis and ceramide formation are major processes, which can stimulate the spontaneous negative curvature of MVB membrane to form ILVs. Then the MVB fuses with the cellular membrane and releases the ILVs, named exosomes or ‘exosome-like vesicles’ [24,25].

The formation process of other EV types is as follows: apoptotic bodies are usually released in the process of cell apoptosis, formed by the wrinkling and invagination of the cell membrane and the subsequent division and wrapping of the cytoplasm [26]. Microvesicles (MVs) are related to the asymmetric distribution of phosphatidylcholine and sphingomyelin in the outer membrane, and phosphatidylserine and phosphatidylethanolamine in the inner membrane. This asymmetric distribution is mainly maintained by three types of proteins: flippase, floppases, and phospholipid scramblase. However, cytoplasmic calcium influx blocks phospholipid asymmetry by activating phospholipid scramblase. This activation leads to the redistribution of phospholipids in the bilayer of the cell membrane, encouraging the budding of the cell membrane [27] (Figure 1).

Components
EVs are enriched in various bioactive substances, mainly proteins, nucleotides (DNA, RNA, mRNA, miRNA, long non-coding RNA [lncRNA] and circular RNA), and lipids [28]. In recent years, with the rapid development of new detection technologies, bioactive components of EVs have been identified. Proteins are notable components of the EV cargo. EV proteins can be divided into two types: ubiquitous proteins involved in the formation of their structures, such as cytoskeleton components and members of the quaternary transmembrane protein superfamily, and those associated with their cell origin [29]. EVs contain several types of nucleic acids, including guideDNA [30], mitochondrial DNA [31] and RNA [23]. Moreover, EVs are abundant in cholesterol, phospholipids, phosphatidylethanolamine and other lipids, which are involved in the biosynthesis and uptake of EVs [32]. Various sorting mechanisms of EV cargo have been described, including the ESCRT machinery [33], syntenin-Alix pathway [34], tetraspansins [35], cytoskeleton [36], lipids [37], and arrestin domain-containing protein 1 [38]. These sorted pieces of machinery enrich specific sets of cargos into EVs, and their consumption can eliminate the production of a defined subpopulation of EVs [39].

EVs in regenerative medicine
Regenerative medicine is aimed to functionally regain or restore tissue and organs from damage, or to form them when they are lost [40]. Cell-based therapies are developed as effective means, in which viable cells, such as stem cells, progenitor cells, inflammatory cells, cartilage cells, islet cells and liver cells, are used for tissue repair [41–43]. Among these, stem cells are the most widely mentioned as they can differentiate into various cell types to mediate tissue repair and regenerative process [44]. In recent years, EVs are emerging as cell-free alternatives to cell therapies, which are exceptionally attractive due to their intrinsic biocompatibility, biodegradability, low toxicity, low immunogenicity and engineered modification [45]. EVs have been explored for in vitro and in vivo studies in various diseases treatment, such as the cardiovascular regeneration, nerve, kidney, liver, lung and skin tissue regenerative therapy [46]. Many clinical trials have also been conducted (https://clinicaltrials.gov/). The key molecules in EV-mediated tissue repair and regeneration processes include miRNAs, mRNAs and proteins. Several signaling pathways involved in regenerative processes, such as mitogen-activated protein kinase, Wnt/b-catenin, PI3K/Akt, Notch, TGF-β/Smad, STAT and Hedgehog signaling, CaMKII, and EfnA3 signaling, have been identified [47–50]. Given their advantages of nano-size and biocompatibility, EVs can cross natural biological barriers, such as the blood–brain barrier, which is a potential strategy for brain and central nervous system injury. In addition, EVs-integrated biomaterials can encourage bone regeneration [51]. EVs can also be loaded with biological macromolecules, short peptides, miRNAs, siRNAs and small molecule drugs.
to promote tissue repair capabilities [52]. For example, the anti-inflammatory drug, curcumin can be encapsulated into EVs to treat various tissue injuries, such as brain, lung and cutaneous injuries, and cerebral ischemia [53–56]. Catalase-loaded exosomes protect the neurons in a mouse model [57]. Some growth factors, such as TGF-β, fibroblast growth factor, vascular endothelial growth factor, and glial cell-derived neurotrophic factor are abundant in mesenchymal stem cell-derived EVs (MSC-EVs), and their delivery contributes to multiple tissue injuries [58,43]. Although this potential has not yet been well characterized, with the growing knowledge of functional roles of EVs, their applications in regenerative medicine will continue to expand.

EVs in cutaneous injury
Cutaneous injury is a common clinical issue, including acute or chronic cutaneous injury due to trauma, burns and diabetes. For chronic severe injury with continuous inflammation, there is a lack of better therapies. Increasing evidence has revealed that EVs are promising therapeutics and drug delivery platforms [59]. It is widely accepted that MSC-EVs are a promising candidate that aids and accelerates wound healing. Due to their extensive effects, MSC-EVs affect a variety of cells associated with wound healing, including keratinocytes, dermal fibroblasts, inflammatory cells and vascular endothelial cells. MSC-EVs participate in the entire healing process by regulating the immune response and inflammation, accelerating proliferation and re-epithelialization of skin cells, modulating ECM remodeling, and promoting angiogenesis [60,61] (Figure 2). EVs derived from other cells, such as plasma endothelial cells [62], induced pluripotent stem cells (iPSCs) [63], perivascular cells [64] and platelet [65]. In addition, in the context of the skin, studies have discovered in ex vivo sections of the human papillary dermis [66], sites of age-related cutaneous disorders [67], injury [68], and stroma of human skin carcinomas [69]. Moreover, in vitro vesicular cross-talk has been observed between several types of skin cells, including keratinocytes, melanocytes, human dermal fibroblasts (HDFs), dermal papilla cells (DPCs), outer root sheath (ORS) cells of the hair follicle, and microvascular endothelial cells [70–72]. How are EVs from specific skin tissue cells that participate in cutaneous repair linked to wound healing? To date, only a few studies have investigated the therapeutic potential of a population of specific skin cells that reside in or participate in wound healing (Table 1).

EVs derived from epidermal keratinocytes
Keratinocytes are highly specialized epithelial cells that play a major role in epidermal restoration after injury through proliferation and re-epithelialization [73]. Within cutaneous tissues, keratinocytes interact with small populations of other cell types, including melanocytes, Langerhans cells, intraepithelial lymphocytes and fibroblasts. Chavez-Muñoz et al. primarily reported that keratinocytes can produce exosomes through which some intracellular proteins such as stratifin, with matrix metalloproteinase (MMP)-1 stimulating activity for fibroblasts, are externalized into the keratinocyte microenvironment [74]. Keratinocyte-derived EVs (KC-EVs) contain a variety of bioactive molecules, such as nucleic acids, metabolic enzymes, cytoskeletal proteins, signaling proteins,
Table 1. Roles of extracellular vesicles derived from special skin tissue cells in wound healing

| EVs sources            | Interacting cells | Cargo(s) | Mechanism(s) and effect(s)                                                                 | Reference |
|------------------------|-------------------|----------|-------------------------------------------------------------------------------------------|-----------|
| Epidermal keratinocyte | Melanocyte        | miRNA-203| Raised the production of Tyr and Rab27a proteins and regulates pigmentation               | [56]      |
|                        |                   | miRNA-3196| Augmented the production of MITF-M and Rab27a and activates the MITF pathway to upregulate melanin production | [56]      |
|                        |                   | miR-330-5p| Downregulated Tyr to suppress melanocyte pigmentation                                      | [57]      |
| Fibroblasts            | Melanocyte        | miRNA-21 | Induced fibroblast function through MAPK/ERK signaling, amplified immune response, and reduced PTEN and RECK expression | [60]      |
| Vascular endothelial   | keratinocyte      |          | Regulated the expression of wound healing-related genes in DFs and induced fibroblast-mediated endothelial tube formation | [62]      |
|                        | keratinocyte      |          | Enhanced DFs and KCs migration via the MAPKinase pathway                                   | [63]      |
| Neutrophil             | Special proteins  |          | Promoted proinflammatory factor production via NF-κB and p38 MAPK signaling pathways      | [64]      |
| Macrophage             | Specific surface  |          | Facilitated the uptake of EVs by macrophage at the wound-edge                             | [65]      |
| T cells                | miR-381-3p        |          | Activated T-bet and RORγt transcription, inducing Th1 and Th17 polarization                | [67]      |
| HFSCs                  |                   |          | Triggered β-catenin and expression of AXIN2, induced HFSCs proliferation and hair growth   | [68]      |
| Epidermal stem cell    | Epidermal cells   | Proteins | Promote epidermal cell growth via various proteins involving epidermis development        | [72]      |
| Myofibroblast          | miR-425-5p        |          | Inhibited myofibroblast differentiation via reducing the TGF-β1 expression                 | [73]      |
|                        | miR-142-3p        |          |                                                                                          |           |
| Fibroblasts, macrophages| miRNAs           |          | Augmented wound cell proliferation and M2 Mϕ polarization through PI3K/AKT and TGFβ signaling pathways possibly | [74]      |
| Dermal fibroblast      | HUVECs            |          | Accelerated diabetic cutaneous wound healing through the Akt/β-catenin pathway            | [79]      |
|                        | DFs               |          | Increased DFs proliferation, inhibited production of ROS induced by UVB radiation due to upregulation of GPX-1 expression possibly | [80]      |
| HDFs                   | miRNA cargos      |          | Restored the function of aged HDFs and ameliorated skin photaging                        | [82]      |
| Keratinocyte           | miR-23a-3p        |          | Regulated epidermal homeostasis by reflecting cellular status                             | [84]      |
| DPSCs                  | Wnt3a             |          | Promoted cells function and elongation of the hair shaft via Wnt/β-catenin pathway        | [85]      |
| DPs, ORCs              | Frizzled4         |          | Stimulated HF growth through DP-secreted Norrin                                            | [86]      |
| Endothelial cells      | Proteins          |          | Increased endothelial cells growth, migration and capillary formation                     | [89]      |
| DFs                    | PLGF-1            |          | Stimulated collagen production and wound healing through PLGF-1 signaling potentially      | [90]      |
| Dermal papilla cell    | Hair matrix cells |          | Facilitated proliferation of hair matrix cells and hair growth via upregulating Wnt3a and β-catenin and downregulating of inhibitory molecule BMP2 | [93]      |
| DPCs                   |                   |          | Increased the expression of growth factors (IGF-1, KGF and HGF) and promotes hair growth  | [94]      |
| Outer root sheath cells|                   |          | Enhanced ORSCs proliferate and migrate to regulate HF development, and stimulated the expression of β-catenin and Shh | [95]      |
| MxCs, ORSCs            | miR-140-5p        |          | Accelerated HF elongation and cell proliferation through BMP/TGF-β signaling pathways     | [96]      |
| HFSCs                  | miR-22-5p         |          | Regulate HFSCs proliferation and differentiation via miR-22-5p-LEF1 axis                   | [97]      |
### Table 1. Continued

| EVs sources | Interacting cells | Cargoes | Mechanism(s) and effect(s) | Reference |
|-------------|-------------------|---------|----------------------------|-----------|
| Macrophage  | Endothelial cells | /       | Attenuated secretion of proinflammatory cytokine, and promoted proliferation and migration of endothelial cells to improve angiogenesis and re-epithelialization | [106]     |
|             |                   | miR-130a, VEGF, Wnt3a | Increased endothelial cellular proliferation, migration and new vasculature formation | [107]     |
| Epithelial cells |                   | miR-5-3p | Reduced inflammatory signals and promoted epithelial regeneration via LATS1/YAP/β-catenin signaling axis | [108]     |
| DFs         | lncRNA-ASLNCS5088 |         | Activated fibroblast and may cause hypertrophic scar formation via adsorbing microRNA-200c-3p, causing increased GLS and α-SMA expression | [109]     |
| DFs         | lncRNA-ASLNCS5088 |         | Promoted proliferation, migration and invasion of DFs via miR-493-3p/AKT signaling pathway | [110]     |
| DPCs        | Vascular endothelial cells | /       | Induced DPCs proliferation and hair growth | [71]      |
| Endothelial progenitor cell | | / | Accelerated re-endothelialization by enhancing endothelial function | [117]     |
|             |                   | miRNA-221-3p | Advanced wound healing by promoting angiogenesis through Erk1/2 signaling | [118]     |
|             |                   |         | Increased expression of angiogenesis-related factors via cell cycle, AGE-RAGE and the p53 signaling pathway probably | [120]     |

**Tyr** tyrosine, **MITF-M** microphthalmia-associated transcription factor melanocyte isoform, **HFSCs** hair follicle stem cells, **DPCs** dermal papilla cells, **ORSCs** outer root sheath cells, **HF** hair follicle, **MxCs** hair matrix cells, **HUVECs** human umbilical vein endothelial cells, **HDFs** human dermal fibroblasts, **EV** extracellular vesicle, **α-SMA** alpha-smooth muscle actin, **DF** dermal fibroblasts

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**Figure 2.** The diagram of EVs repair cutaneous injury and mechanism. (a) Cutaneous injury. The entire structure of the skin is destroyed, accompanied by an inflammatory response. In chronic wounds such as diabetic ulcers, the skin tissue is severely damaged, the epidermis is incomplete, dermal fibers are broken, blood vessel formation disorders with continuous inflammatory infiltration. (b) EVs treatment. EVs promote epidermal and dermal architecture and function recovery and angiogenesis and play an immunomodulatory role through RNA, such as microRNA and circRNA, as well as proteins and growth factors. Engineered EVs repair cutaneous injury better. **EV** extracellular vesicle, **ECM** extracellular matrix, **MSC** mesenchymal stem cell

 trafficking proteins and adhesion proteins, and have significant physiological consequences on target cells [75, 76]. Melanocyte–keratinocyte interactions are a well-studied system in which melanocyte activity, such as pigment production and pigment transfer, is assisted by EVs [77]. The study demonstrated that miRNA-203 and miRNA-3196 are two major functional molecules. KC-EVs conveyed miRNA-203 increases the production of tyrosine (Tyr) and Rab27a proteins, which subsequently induce Tyr activation and pigmentation. KC-EVs shuttled miRNA-3196 augments the production of microphthalmia-associated transcription factor (MITF) melanocyte isoform (MITF-M) and Rab27a, which then activates the MITF pathway to upregulate melanin production [78]. Liu et al. demonstrated that
keratinocytes secrete exosomes carrying miR-330-5p and decrease pigmentation in melanocytes [79]. It has also been reported that keratinocyte-derived MVs can mediate ultraviolet B radiation-induced systemic immunosuppression [80]. The interaction between keratinocytes and melanocytes reveals a potential mechanism of pigment production.

Additionally, the ECM of the epidermis is enriched with KC-EVs, and interactions with integrins expressed by intact cells regulate multiple cellular activities, such as adhesion, migration, differentiation, apoptosis and expression of specific genes [81,82]. It is hypothesized that EVs secreted by keratinocytes and incorporated into the ECM may deliver their cargo, including integrins, to dermal fibroblasts during the healing process. Li et al. found that human keratinocyte-derived MVs delivered miRNA-21 and promoted skin wound healing in diabetic rats by facilitating fibroblast function and angiogenesis [83]. EVs can also affect the gene expression in fibroblasts. For example, EVs derived from epidermal cells significantly regulate genes encoding MMP-1 and -3, IL-6 and IL-8, and genes associated with TGF-β signaling in fibroblasts. In addition, the presence of MV-like vesicles during active keratinocyte migration and early stages of granulation tissue organization in human wounded skin [8,84]. In addition, it has been proved that MV-like vesicles are present in the early stages of active keratinocyte migration and granulation tissue organization in injured human skin. Another study identified that HaCaT cell-derived EVs expedite the migration and proliferation of human keratinocytes and fibroblasts and may advance wound healing via activation of the mitogen-activated protein kinase pathway [85].

Moreover, EVs are emerging as mediators of intercellular communication between keratinocytes and immune cells. Cytokine-treated keratinocytes secrete exosomes that promote neutrophil pro-inflammatory factor production, containing IL-6, IL-8 and TNF-α [86]. Wound macrophages selectively absorb exosomes derived from wound-edge keratinocytes because of the specific surface N-glycan and miRNA packaging, inducing a wound inflammation response. This indicates close cross-talk between keratinocytes and macrophages through exosome-delivered miRNAs [87,88]. Jiang et al. demonstrated that EVs generated by cytokine-treated KCs promoted the polarization of Th1 and Th17 cells in psoriasis [89]. This highlights the importance of EVs as a communication strategy between KCs and T cells under psoriasis-like conditions. Cutting off these lines of communication by either inhibiting EVs secretion or blocking functional cargo could be a useful therapeutic strategy in patients with psoriasis and other inflammatory diseases. KC-EVs also participate in hair growth, and a study found that EVs derived from fisetin-treated KCs mediate hair growth promotion [90]. It has been suggested that exosomes can be considered as natural mediators that could be involved in hair cycle regulation and serve as promising delivery vehicles for improving skin and hair regeneration because of their potential to target various molecular processes and cells.

### EVs derived from epidermal stem cells

Keratinocytes proliferate from the basal cells of the innermost layer of the skin (stratum basalis) [91]. Epidermal stem cells (ESCs) are attached by hemidesmosomes to the stratum basalis, maintain skin homeostasis and hair regeneration, and provide the original power for re-epithelialization after injuries [92]. ESCs are morphologically similar to KCs, but have specific identification markers. These are of particular interest because they are numerous and accessible. Additionally, ESCs are easy to obtain without potential ethical and political issues compared with embryonic stem cells [93]. ESCs also possess a strong paracrine function; therefore, their effects on epidermal cells, dermal cells and angiogenesis in wound healing should not be ignored.

Leng et al. demonstrated that ESC-EVs were beneficial in epidermal cell growth and classical stemness regulators, and that Wnt signaling was always involved [94]. Duan et al. determined that ESC-derived exosomes enhance the wound healing rate and reduce scar formation in rats by delivering miR-425-5p and miR-142-3p [95]. Recent research has shown that exosomes originating from ESCs could improve diabetic wound healing and both ESCs and ESC-Exos have an equal effect. This indicates that ESC-Exo treatment may be a promising and technically advantageous stem cell-based therapeutic strategy [96].

### EVs derived from dermal fibroblasts

Cutaneous injury repair depends on the interaction between cells and the ECM. Fibroblasts are the main force in damage repair and ECM formation. Under the stimulation of several chemokines, fibroblasts migrate from the wound perimeter to the wound surface and secrete a large number of ECM components, such as collagen, fibronectin, laminin, vitronectin and proteoglycan. The capacity of dermal fibroblasts (DFs) in wound healing has been well clarified; however, their function seems to be underestimated. Fibroblasts are mesenchymal cells [97], which are gradually considered to be convenient alternatives to MSCs because of their common characteristics, such as expression of the same mesenchymal markers and multi-differentiative potential toward mesodermal lineage, and anti-inflammatory, immunomodulatory and regenerative effects [98,99]. Another explanation is that cultured DFs are mixed-cell populations, including stem cells at different developmental stages [100]. Their therapeutic potential is similar, it should be noted that DFs are more accessible than MSC and are easier to expand in vitro. Thus, the utilization of DFs may be more practical in clinical applications.

As active mediators of both the inflammatory and proliferative phases of wound healing, DFs communicate with all cell types in the wound bed through direct contact or in autocrine and paracrine manners [101]. EVs are key components of paracrine signaling and are responsible for communication between cells during wound healing. DFs-EVs have not been well investigated as they are generally accepted as terminally differentiated cells. Our previous study found that exosomes derived from autologous DFs could reverse...
the damage caused by high glucose levels in human umbilical vein endothelial cells (HUVECs) in vitro and promote diabetic cutaneous wound healing through the Akt/β-catenin pathway [102]. Deng et al. demonstrated that both MSC-EVs and DFs-EVs showed antioxidant activity that could reduce the intracellular levels of reactive oxygen species (ROS) induced by UVB irradiation, although DFs-EVs were initially used as a negative control [96]. Interestingly, Wang et al. indicated that exosomes derived from ESCs (ESC-Exo) accelerated wound healing when compared to fibroblast exosomes (FB-Exo) and phosphate buffered saline (PBS) controls [103]. From the experimental results, FB-Exo also has an effect but is not as effective as ESC-Exo. Due to the conditions and standardization of research, it is difficult to conclude which origin of exosomes works the best. Furthermore, EVs derived from three-dimensional cultured DFs are also able to induce efficient collagen biosynthesis and ameliorate inflammation in UVB-induced skin injury, even better than those isolated from two-dimensional cultured DFs [104]. Changes in growth conditions and the culture environment can alter the state of cells, including their paracrine function. By changing the growth environment or adding stimulation factors, cells can transform to a better state and secrete EVs with higher yields and functions to achieve a better therapeutic effect, which is also a research hotspot.

EVs facilitate intercellular communication and enable cells to exchange proteins, lipids and nucleic acids [105]. The dermis and epidermis are in close contact with each other; therefore, what is the role of EVs as a means of communication? Choi et al. investigated whether human dermal fibroblast-derived EVs (HDFs-EVs) affect the homeostatic regulation of keratinocytes and found that senescent fibroblasts produced more EVs than young HDFs; however, these EVs attenuate the dermal effect on keratinocyte differentiation and evoke the pro-inflammatory cytokine IL-6 [106]. Similarly, senescent fibroblasts-derived EVs accelerated the scratch/gap closure of keratinocytes by delivering miR-23a-3p [107]. In addition, HDF-EVs can increase the ability of human DP and ORS cells, thereby promoting hair growth in cultured human hair follicles [108]. EVs from activated DFs stimulate hair follicle growth via DP-secreted norrin [109]. During wound healing, myofibroblasts shrink the wound by contracting the wound edges and secreting a large quantity of extracellular matrix to connect with the surroundings [110]. Myofibroblasts, particularly those existing in the wound, also produce MVs as mediators of intercellular communication to facilitate ECM formation and neovascularization [71]. The release of MVs by wound myofibroblasts brings new perspectives to the field of communication between cells during the normal healing process [111].

**EVs derived from dermal papilla cells**

Hair is a defining feature of mammals and has critical functions, including protection, production of sebum, apocrine sweat and pheromones, social and sexual interactions, thermoregulation, and the provision of stem cells for skin homeostasis, regeneration and repair [112]. Hair follicles and hair growth are critical indicators of skin recovery after injury. Hair DPCs are specialized mesenchymal cells that reside in the DP located at the bottom of the hair follicles [113]. DPCs serve as signaling centers in hair follicles and regulate hair formation and cycling via paracrine secretion. DPC-derived EVs (DPC-EVs) are critical for hair regeneration.

Numerous studies have demonstrated that DPC-EVs contribute to hair growth. For instance, research has revealed that the sustained release of DPC-EVs from injectable microgel promotes hair growth, which may be due to the upregulation of hair growth-promoting signaling molecules, such as Wnt3a and β-catenin, and downregulation of the inhibitory molecule BMP2 [114]. Another finding indicated that DPC-EVs encourage hair growth and regeneration by modulating the activity of follicular dermal and epidermal cells and augmenting the hair-inductive capacity of cultured DP spheres [115]. In vitro, DPC-EV treatment enhances the ORS proliferation and migration and stimulates the expression of β-catenin and sonic hedgehog [116]. Correspondingly, Chen et al. also demonstrated that EVs containing miR-140-5p from human papilla cells stimulate hair growth by promoting the proliferation of ORS and hair matrix cells [117]. DP also regulate stem cells; for example, Yan et al. found that miR-22-5p derived from DP-EVs mediates hair follicle stem cell proliferation and differentiation [118]. Additionally, DPC-EVs also increase hair-inductive gene expression in adipose stem cells via β-catenin activation [119]. Overall, DPC-EVs represent a therapeutic target for damage repair, especially for hair growth after injury.

**EVs derived from macrophages**

In wound healing, inflammation is a major physiological response and a necessary period of transition, which is orchestrated by cells (such as leukocytes, stem cells and tissue-resident cells) and the molecular messengers they secrete, including cytokines, growth factors and exosomes [120]. Immune cells play a prominent regulatory role in the inflammatory and proliferative phases of skin healing. A moderate inflammatory response promotes damage repair; however, excessive inflammation may cause further damage. Among the immune cells in the wound, macrophages are key players that facilitate the inflammatory proliferation phase transition. Nevertheless, macrophages show the opposite effect due to their subtypes: classically activated M1 with pro-inflammatory properties and alternatively activated M2 exhibiting anti-inflammatory and wound healing functions [121]. In the early stages of wound healing, macrophages are usually polarized to M1 macrophages, which are microbicidal and pro-inflammatory [122]. With the resolution of inflammation, macrophages transform into M2 macrophages, which secrete vascular endothelial growth factor (VEGF) to support angiogenesis [123,124]. During the proliferation stage, macrophages also interact with DFs and induce them to myofibroblast transformation, increasing ECM deposition [125,126]. An overreactive
Figure 3. Quadrants reflecting inflammation and cutaneous injury repair. Moderate inflammation contributes to repair, but excessive inflammation has the opposite effect. The first quadrant represents the early stages of inflammation. Neutrophils play an antibacterial and threatening role, dendritic cells (DCs) are antigen presenting cells that are involved in priming T-cell responses. The second quadrant represents the late stage of inflammation, in which M2 macrophages play a dominant role. The third quadrant represents the normal homeostasis of the skin when there is no injury or slight inflammation. The fourth quadrant represents excessive inflammation in pathological conditions, where the action of a range of inflammatory cells and proinflammatory factors makes wound healing difficult. ECM extracellular matrix, VEGF vascular endothelial growth factor

Inflammatory response during the inflammatory phase or excessive deposition of ECM during the proliferation phase causes abnormal scar formation [127]. Therefore, the study of inflammatory cells, particularly macrophages, is indispensable (Figure 3). Macrophage-derived exosomes have been shown to accelerate wound healing through anti-inflammatory effects and by inducing endothelial cell proliferation and migration to improve angiogenesis and re-epithelialization in diabetic wounds [128]. Another study also indicated that macrophage-derived EVs (MAC-EVs) promote angiogenesis in vitro and accelerate new vasculature formation in vivo and found that VEGF, Wnt3a and miR-130a were abundant in MAC-EVs, even more than in parental cells [129]. M2 macrophage-derived exosomal miR-590-3p reduces inflammatory signals and promotes epithelial regeneration [130]. Macrophages also stimulate DFs activity, and activated macrophages (M2) manipulate fibroblasts to differentiate into myofibroblasts with active biological functions and proliferation, which may cause a hypertrophic scar. Chen et al. demonstrated that lncRNA-ASLNCS5088 derived from M2 macrophage exosomes orchestrates fibroblast activation, and its blockade by
Figure 4. Diagrammatic illustration of EVs derived from different types of cells, including keratinocyte, epidermal stem cells, fibroblast, macrophages, dermal papilla cells and endothelial progenitor cells. These cells are related to cutaneous injury and their cross-talk plays significant roles in wound healing. ECM extracellular matrix, VEGF vascular endothelial growth factor, EV extracellular vesicle, KC keratinocyte, MAC macrophage, DF dermal fibroblasts

DW4869 dampens this effect [131]. Similarly, another study indicated that the inhibition of linc01605-enriched exosome generation in M2 macrophages impairs M2 macrophage-induced proliferation, migration and invasion of HDFs [132]. This finding further supports the dual role of macrophages. Furthermore, engineered EV-mimetics from macrophages facilitate hair growth in mice and encourage human hair follicle growth [133]. The complexity of macrophages in wound healing makes them more difficult to study, and the research is not sufficiently in-depth, which also provides a possibility for future research.

EVs derived from endothelial progenitor cells
Endothelial progenitor cells (EPCs), a particular type of stem cells, represent a heterogeneous population of resident mononuclear cells that originate from the bone marrow (BM) [134]. Although EPCs do not originate in the skin, they influence skin wound healing as precursors of vascular endothelial cells [135]. Vascular formation is an essential part of skin wound healing, which provides oxygen for the cells and microenvironment at the wound site, and is conducive to the formation of granulation tissue. EPCs can secrete paracrine factors such as growth factors, cytokines, chemokines and bioactive lipids that influence cell biology in damaged tissue, among which EVs are crucial bioactive factors [136,137]. EPC-EVs have potential therapeutic applications in tissue repair and regeneration, such as diabetic foot ulcers, kidney disease and bone healing [138,139]. Studies have shown that exosomes derived from EPCs attenuate vascular repair and accelerate re-endothelialization by enhancing endothelial function [140]. EPC-derived exosomes have been reported to facilitate skin wound healing by positively modulating endothelial cell function [141]. After local injection of EPC-EVs into the skin wounds of diabetic rats, Zhang et al. [142] discovered that EPC-EVs could significantly improve wound healing and accelerate angiogenesis in damaged areas by modulating extracellular signal-regulated kinase 1/2 signaling. Xu et al. [143] also identified exosomes isolated from murine BM EPCs that accelerate skin wound healing in both control and diabetic mice by transferring miRNA-221-3p. Hassanpour et al. found out that the diabetic state could affect the CD63-Alix-Rab27a signaling pathway, thereby reducing the formation, transportation and fusion of EVs in EPCs [144]. Those results underline the role of EPC-exosomes and suggest new potential approaches to the therapy for diabetic skin wounds (Figure 4).

Open questions and future directions
Accordingly, EVs derived from both MSCs and specific skin tissue cells are involved in wound healing and can be used
as promising agents for damage repair. However, several challenges and questions remain to be addressed.

**Skin cell-EVs vs MSC-EVs** MSCs have diverse sources, including BM, adipose tissue, umbilical cord, umbilical cord blood, dental pulp, synovial fluid, amniotic fluid, placenta, Wharton’s jelly and body fluids [145]. In addition, MSCs have immunomodulatory properties to regulate various cells involved in immune responses [146]. Clinical trials have indicated that MSC-based therapy could effectively improve burn wound healing, and treat hypertrophic scars [147,148]. Urine-derived stem cells have been successfully used for skin, bone and articular cartilage repair [149]. EVs are important effectors of MSCs in paracrine activity, emerging as the most promising candidates for tissue repair and regenerative medicine. Our previous research found that hucMSC-exosomes containing Wnt4 promoted cutaneous wound healing in a rat skin burn model [150]. Exosomes originating from both human umbilical cord blood mesenchymal stem cells and human BM-MSC effectively stimulate cutaneous wound healing by inhibiting the TGF-β signaling pathway [151,152]. Other studies indicated that adipose mesenchymal stem cell-derived exosomes accelerate skin wound healing via the lncRNA H19/miR-19b/SRY-box transcription factor 9 axis [153] and alleviate the inflammatory response [154]. Human amniotic fluid stem cell- or human amniotic epithelial cell-derived exosomes accelerate wound healing [155,156]. Moreover, human decidua MSC-derived exosomes can improve high-glucose senescent HDFs via intracellular miR-145-5p and miR-498 [157].

However, in the healing process, multiple internal cell types, including neutrophils, monocytes, macrophages, keratinocytes, fibroblasts, T cells, B cells, mast cells and endothelial cells are involved. These cells are actively associated with the production and regulation of various cytokines and growth factors [158], fight infection at the site of injury, form new tissues, and ultimately lead to wound closure [159]. MSC is ‘external assistance’, and these cells are ‘real participants’. EVs are also crucial in disease progression and anti-inflammatory repair, but the role of EVs is not fully understood and clarified. In the complex microenvironment of wound healing, skin tissue-EVs are more likely to express receptors for intercellular communication. The study developed a method of isolating keratinocyte-derived exosomes with genetically labeled GFP-reporter (Exoₖ-GFP) from wound-edge tissue. Moreover, surface N-glycans of Exoₖ-GFP were also characterized and glycan ions with high mannose were only detected in wound-edge Exoₖ. Wound-edge KC exosomes revealed characteristic N-glycan ions with an abundance of low-base-pair RNA and were selectively engulfed by wound macrophages (ommp) during granulation [87]. The Exoₖ-GFP may be a regulator of macrophage trafficking and epithelial barrier post-injury. Additionally, skin tissue cell-EVs can act directly on homogeneous cells. These findings reveal that miR-125b is transferred via exosomes from young fibroblasts to old fibroblasts, contributing to the promotion of fibroblast migration and transition to counter-act aging [160]. Growth factor-mediated cross-talk between keratinocytes and fibroblasts reveals the double paracrine signaling [104]. Some early research suggested that EVs have different characteristic molecular expression profiles due to their different sources and can act on different target cells with certain selectivity. For example, exosomes that express the transmembrane surface protein network tSPAN8-Alpha4 are more likely to enter endothelial cells and pancreatic cells [161]. Exosomes that express CD47 are less susceptible to uptake by monocytes [96]. Skin cell-EVs and MSC-EVs may have various molecular expression profiles and it may be speculated that specific skin cells-derived EVs could selectively function on target cells; however, the explicit difference between the two remains unclear and related research is lacking. Is there a unique advantage of the skin microenvironment for the self-repair effects of skin tissue cells? Recent studies have shown that EVs derived from ESCs exhibit more potent repair effects than other stromal cell exosomes do. Both ESC-Exo and FB-Exo promote diabetic wound closure, but ESC-Exo show better effects [162]. Various cells have special or common functions; for example, re-epithelialization is dependent on the ability of keratinocytes to form holoclones [163]. Melanocytes and fibroblasts may contribute to re-pigmentation [164]. Cellular interactions at wound sites promote fibroblast attachment and produce a new connective tissue matrix and double paracrine interaction between keratinocytes and fibroblasts [165]. These findings suggest that EVs from skin tissues play unique roles in cutaneous wound repair. They can better reflect the cross-talk between cells in the wound healing environment and reveal the mechanism of injury repair. Further research will aid in the understanding of their contribution to wound healing in more detail and may explain the different clinical outcomes. Both skin cell-EVs and MSC-EVs possess great clinical application potential, but currently clinical trials mainly focus on MSC-EVs (Table 2). More potential and applications of skin-EVs require further research.

**Complexity and risks of EV application in pathological conditions** The regenerative effects of EVs are at least in part attributed to the transfer of functional messages, which are highly dependent on specific conditions. The normal wound healing cascade is well coordinated and synchronized by growth factors, different MMPs, cytokines, inflammatory cells, keratinocytes, fibroblasts and endothelial cells. Chronic injury, such as diabetic wounds, exhibits a persistent inflammatory phase and is mainly characterized by the presence of chronic low-grade inflammation [166,167]. In pathological conditions, such as inflammation, aging and different living statuses, EVs may carry ‘undesired’ messages that contribute to the spread of diseases. Do EVs secreted by specific skin tissue cells consistently exhibit the supporting effect? Macrophages are pivotal contributors to healing; however, inflammatory cells are also a double-edged sword, and EVs as a medium of communication may have the opposite effect in
the context of persistent inflammatory infiltration. Research has indicated that defective MV secretion/function may cause aberrant wound healing and chronic wounds that fail to re-epithelialize [168]. Evidence indicates that mast cell exosomes can produce cytosolic PLA2 and contribute to a CD1α-reactive T cell response in psoriasis, a chronic inflammatory skin disease [86]. The study discovered that keratinocyte exosomes activate neutrophils and enhance skin inflammation [169]. Similarly, neutrophil exosomes can also stimulate keratinocytes and increase skin autoinflammation in generalized pustular psoriasis [170]. This reveals that exosomes not only work as a means of keratinocyte-neutrophil communication but also show detrimental effects. In allergic inflammation, exosomes mediate cellular interactions by regulating the ROS levels [171,172].

Aging is accompanied by chronic low-grade systemic inflammation [173]. Senescent cells accumulate in the skin during aging and alter the skin’s microenvironment [174,175]. Research has proved that fibroblasts from aged skin exhibit a senescence-associated secretory phenotype rich in pro-inflammatory cytokines, chemokines and proteases, which can have a detrimental impact on nearby cells and the local tissue environment [105]. Choi et al. also demonstrated that EVs derived from senescent fibroblasts attenuated the dermal effect on keratinocyte differentiation [176]. EVs in the aging microenvironment may have detrimental effects on downstream targets at the levels of immunity, inflammation, gene expression, and metabolism [75]. Additionally, keratinocytes are divided into differentiated (diff-K) and undifferentiated keratinocytes (undiff-K). The EVs produced by them also differ in their contents and effects. The study found exosomes containing 14-3-3s from diff-K had a higher MMP-1 stimulatory effect in fibroblasts than exosomes isolated from undiff-K [177].

**Intrinsic challenges of EVs** At present, EVs derived from stem cells have been developed into products and are gradually being applied from the bench to the bedside. For example, exosome repair fluid is used for scars or in the form of dressings. Although exosomes have broad prospects in the field of skin repair and cosmetic medicine, there are still several challenges in their clinical application. For instance, a lack of appropriate methodology, inadequate cell number availability, insufficient targeting and poor stability in circulation [178–180].

**Technical challenges** Currently, we do not yet have the tools needed to properly study EVs. The heterogeneity of sEVs and the presence of nonvesicular extracellular substances have led to debate about the contents and functional properties of exosomes [181]. Several commonly used EV purification methods have been developed, but all have limitations, such as producing EVs with different yields and purities [18,182]. To improve the purity of separation, the first problem to be addressed is to clarify the classification of EVs and the specific differences between sEV, other EVs, and nonvesicular in physical and chemical aspects, such as size, density, surface protein, sugar, lipid components and surface charge [180]. High-resolution density gradient fractionation and direct immunoaffinity capture were employed to precisely characterize the RNA, DNA, and protein constituents of exosomes and other nonvesicular materials. They found that extracellular RNA, RNA-binding proteins, and other cellular proteins were differentially expressed in exosomes and nonvesicle compartments [183], allowing for more precise determination of the molecular composition of classical exosomes.

### Table 2. Clinical trials with extracellular vesicles for cutaneous injury

| NCT number | Phase | Status | Condition | Source | Therapeutic | Reference |
|------------|-------|--------|-----------|--------|-------------|-----------|
| 05243368   | Not yet applicable | Not yet recruiting | Diabetic foot | MSC | AGLE 102 | / |
| 04173650   | Phase I/II | Not yet recruiting | Dystrophic epidermolysis bullosa | MSC | AGLE 102 | / |
| 05078385   | Phase I | Not yet recruiting | Burns | BM-MSC | AGLE-102 | / |
| 04134676   | Phase I | Completed | Chronic ulcer | MSC | Conditioned medium | / |
| 04235296   | Phase I | Recruiting | Residual burn wound | MSC | Conditioned medium | / |
| 03686449   | Not yet applicable | Recruiting | Post-burn raw area | Autologous KC ADSC-KC | Suspension | / |
| 02563264   | Early Phase I | Unknown | Ulcer | Plasma | Exosome | / |
| 04134676   | Phase I | Completed | Chronic ulcer | WJ-MSC | Conditioned medium | / |
| /          | Preclinical | / | Skin lesions, inflammatory and autoimmune disorders | Not indicated | Exosome-based product | [166] |
| /          | Preclinical | / | Skin lesions | ADSCs, placental MSCs, cord blood serum | / |
| /          | / | / | Acne scars | Adipose tissue stem cell | Exosomes gel | [18] |

**Notes:** MSC mesenchymal stem cells, BM-MSC bone marrow mesenchymal stem cells, ADSCs adipose-derived stem cells, KC keratinocyte, WJ-MSC Wharton’s jelly mesenchymal stem cells.
Inherent defects in natural EVs To overcome the inherent shortcomings of EVs, researchers have enhanced their capabilities by designing or generating biomimetic EVs. Wang et al. developed an injectable, self-healing and antibacterial polypeptide-based FHE hydrogel, which is named from its composition of pluronic F127 (F127), oxidative hyaluronic acid (OHA), and ε-poly-l-lysine, polypeptide (EPL) with exosomes to accelerate diabetic wound healing [184]. Shen et al. designed bilayered thiolated alginate/polyethylene glycol diacrylate hydrogels for the sequential release of EVs to achieve rapid and scarless wound healing [183]. The hydrogel itself provided a moist environment, which was conducive to debridement, and the hydrogel-loaded EVs enhanced their stability in vivo and played a role in sustained release [186]. Cutaneous injury is often accompanied by infection, so the combination of antibacterial materials and EVs is also a research hotspot, to play a dual role. Dressing loaded-exosomes and AgNPs have antimicrobial activity and promote wound healing [187]. Similarly, exosome-laden oxygen-releasing antioxidant and antibacterial cryogel wound dressing OxOBand could support diabetic wound closure [188]. Membrane-targeting modification of EVs is also a strategy. Su et al. demonstrated that PD-L1 overexpression in exosomes could be specifically bound to PD-1 on the T cell surface and suppressed T cell activation to facilitate tissue repair [189]. Immunosuppression of excessive or persistent inflammation during wound healing is another effective approach. To enhance the organ-targeting ability of EVs, Li et al. fabricated iron oxide nanoparticle (NP)-labeled exosomes (Exo + NPs) and found that they were readily internalized by HUVECs [190]. Whether natural or engineered, EVs show a strong ability to repair damage and engineered EVs constitute a new strategy and may soon be developed as smart drugs for clinical treatment.

Clinical translational opportunities To achieve the application of EVs from the bench to the bedside, research has focused on the basic biology of EVs, including their biogenesis, dynamics in extracellular space, transit and uptake by target cells. To completely capitalize on their potential, a better understanding of the cellular and molecular mechanisms that govern the many observed functions of EVs is required. Numerous issues need to be acknowledged; for example, how do EVs interact with different matrices? How are EVs maintained in (or cleared from) biological fluid? How do they cross the biological barriers? How are they determined in specific tissues? How are they taken up into cells, and how are cargo functionally delivered? The dose and safety should be evaluated in clinical applications. The systematic application of EVs may induce apoptosis in some cells, including immune regulation [191]. Generally, the wound is used locally, but whether systemic or local, EVs are easily cleared rapidly [192]. Topical EVs are, in the form of liquid dressings, sprays or solid plasters. Different forms of EVs may affect the state of EVs and have different effects. Additionally, the injection dose of EVs must be considered, as different doses can produce various biological effects. When EVs are applied for the treatment of neurodegenerative diseases, it has been shown in vitro that low doses of EVs have neuroprotective effects, whereas high doses may be harmful to neurons [193]. EVs may exert similar effects on wound repair. Moreover, injection of EVs into the body may induce an immune response, and there are differences in individual responses [194]. EVs have exhibited superior therapeutic effects in regenerative medicine; however, most applications are still in the experimental stage. A great deal of clinical validation is needed for the regulatory approval of EV-based diagnostics and therapeutics. An easier, efficient, safe and high-yield method for EVs production must be developed to provide better EVs for future clinical use. In summary, further studies on EVs and their gradual clinical trials in skin injury repair have broad application prospects.

Conclusions

This review discusses the effects of EVs derived from specific epidermal cells, such as dermis and inflammatory cells, on cutaneous repair and regeneration. EVs have high stability, biocompatibility, low toxicity, and immunogenicity. Therefore, they are potential candidates for novel cell-free based therapies for different diseases, including skin tissue repair. While not intended to substitute completely for classical forms of intercellular communication (hormones, chemokines, cytokines and mitogens), EVs provide a local environment potentially and modify the physiology and functional activities of cells. The study of special skin tissue cell-derived EVs in wound healing will be beneficial for revealing the dynamic process of wound healing and the specific mechanism of cell function. Taken together, this study provides novel insights into EVs derived from specific skin tissue cells and the importance of cross-talk between different skin cell types in wound healing. However, many open questions remain regarding the signaling molecules and uptake of EVs that participate in the EVs-induced cell response. Little is known about the dynamics of EVs in wound healing. Mechanistically, it would be interesting to further elucidate how cells interact with and influence each other. Therefore, further research is required to clarify the regulation of EVs during the wound healing process and to translate these results to clinical settings.

Abbreviations

BM: Bone marrow; DFs-EVs: Dermal fibroblasts-derived EVs; diff-K: Differentiated keratinocytes; DPCs: Dermal papilla cells; DPC-EVs: Dermal papilla cells-derived EVs; ECM: Extracellular matrix; EMT: Epithelial–mesenchymal transition; EPC-EVs: Endothelial progenitor cells-derived EVs; ESC-EVs: Epidermal stem cells-derived EVs; ESC-Exo: Epidermal stem cells-derived exosomes; ESCRT: Endosomal sorting complex required for transport; EV: extracellular vesicle; FB-Exo: Fibroblast exosomes; HDFs: Human dermal fibroblasts; HDFs-EVs: Human dermal fibroblasts-derived EVs; HUVECs: Human umbilical vein endothelial cells; ILVs:
Intraluminal vesicles; iPSCs: Induced pluripotent stem cells; KC-EVs: Keratinocyte-derived EVs; MAC-EVs: Macrophage-derived EVs; MITF: Microphthalmia-associated transcription factor; MMP: Matrix metalloproteinase; MSC-EVs: Mesenchymal stem cell-derived EVs; MVs: Microvesicles; MVBs: Multivesicular bodies; NETs: Neutrophil extracellular traps; ORS: Outer root sheath; ROS: Reactive oxygen species; sEVs: Small EVs; undiff-K: Undifferentiated keratinocytes; VEGF: Vascular endothelial growth factor; FHE: Pluronic F127 (F127); OHA: oxidative hyaluronic acid; EPL: polypeptide ε-poly-L-lysine

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Conflicts of interest

None declared.

Authors’ contributions

Conceptualization: HS, MW, PW; writing: MW; data collection: MW and WL; manuscript revision: MW and JH; review and editing: HS, YS, HQ.

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