Detrimental impact of acute and chronic glucose burden in wound-healing cells: fibroblasts, myofibroblasts and vascular precursor cells

Jorge Berlanga-Acosta1, Ernesto López-Mola1, Marianela García-Siverio1, Gerardo Guillén-Nieto1, Pedro López-Saura1, Calixto Valdez-Pérez2, Isabel Puentes-Madera2, William Savigne-Gutiérrez2, Héctor Álvarez-Duarte2, Norberto Miranda-Espinosa2, Ana J Mir-Benítez4, Diana García del Barco1, Yssel Mendoza-Mari1, María D Martinez-Espina1, Ariana García-Ojalvo1, Nelvys Subiros-Martínez1, Luis Herrera-Martínez1

1 Departamento de Cicatrización y Citoprotección. Dirección de Investigaciones Biomédicas, Centro de Ingeniería Genética y Biotecnología, CIGB Ave. 31 e/ 158 y 190, Cubanacán, Playa, CP 11 600, La Habana, Cuba

2 Servicio de Angiopatía Diabética, Instituto Nacional de Angiología y Cirugía Vascular, INACV, Calzada del Cerro s/n. Cerro, La Habana, Cuba

3 Departamento de Cirugía, Hospital Pediátrico Juan Manuel Márquez, Marianao, La Habana, Cuba

4 Departamento de Cirugía Plástica y Reconstructiva, Hospital Joaquín Albarrán, Cerro, La Habana Cuba

E-mail: jorge.berlanga@cigb.edu.cu

ABSTRACT

Type 2 diabetes mellitus comprises a group of non-communicable metabolic diseases with an expanding pandemic magnitude. Diabetes predisposes to lower extremities ulceration and impairs the healing process leading to wounds chronification. Diabetes also dismantles innate immunity favoring wound infection. Amputation is therefore acknowledged as one of the disease’s complications. Hyperglycemia appears as the proximal detonator of toxic effectors including pro-inflammation, spillover of reactive oxygen and nitrogen species. The systemic accumulation of advanced glycation end-products irreversibly impairs the entire physiology from cells-to-organs. Insulin axis deficiency weakens wounds’ anabolism and predisposes to inflammation. These factors converge to hamper fibroblasts and endothelial cells proliferation, migration, homing, secretion and organization of a productive granulation tissue. Diabetic wound bed may turn chronically inflamed, pro-catabolic and a superimposed source of circulating pro-inflammatory cytokines, establishing a self-perpetuating loop. Diabetic toxicity breadth includes mitochondrial damages in fibroblasts and endothelial cells becoming prone to apoptosis thus hindering granulation. Endothelial progenitor cells recruitment and tubulogenesis are also impaired. Failure of wound re-epithelialization remains as a clinical challenge while it appears to be biologically multifactorial. Novel medical interventions as the local intra-ulcer infiltration of epidermal growth factor have emerged to hopefully reduce the current worldwide amputation rates.

Keywords: diabetes, granulation, ulcer, re-epithelialization, growth factors

Impacto perjudicial del exceso agudo y crónico de glucosa en células involucradas en la cicatrización: fibroblastos, miofibroblastos y células precursoras vasculares. La diabetes mellitus tipo 2 implica desórdenes metabólicos no transmisibles, cuya incidencia va en aumento, con una extensión casi pandémica. Predispone a padecer úlceras en las extremidades inferiores y a su cronicidad, al afectar el proceso de cicatrización. También interfiere la inmunidad innata, lo que favorece la infección de posibles lesiones, y puede conducir a la amputación. La hiperglucemia desencadena los efectores tóxicos que incluyen la inflamación y la producción en exceso de especies reativas de oxígeno y nitrogénio. Los productos finales altamente glicosilados, que se acumulan sistemáticamente, desarticulan la estructura de células y órganos. La deficiencia en el eje insulinico debilita el anabolismo en las lesiones y predispone a la inflamación. Estos factores convergen y debilitan la proliferación, la migración, el direccionamiento, la secreción y la organización de los fibroblastos y células endoteliales, lo que interfiere en la formación de tejido de granulación útil. El lecho de las heridas puede convertirse en una fuente inflamatoria y pro-catabólica de citocinas, y constituir un ciclo de perpetuación. La toxicidad diabética provoca daños mitocondriales en los fibroblastos y las células endoteliales, que los hace susceptibles de apoptosis y dificulta la granulación del tejido. También afecta el reclutamiento de las células progenitoras endoteliales, e impide la tubulogénesis. La regeneración del tejido epitelial en las lesiones sigue siendo un desafío clínico que depende de múltiples factores biológicos. Nuevas intervenciones médicas, como la infiltración local intralesional del factor de crecimiento epidérmico recombinante, prometen la reducción de las tasas mundiales de amputación.

Palabras clave: diabetes, granulación, úlcera, reepitelización, factores de crecimiento

Corresponding author
Introduction

Diabetes mellitus represents today a worldwide pandemic disease under the most two common clinical forms identified as types 1 and 2. The former is a condition in which by autoimmun mechanisms pancreatic β-cells are eventually destroyed with an absolute insulin deficiency [1]. Type 2 diabetes mellitus (T2DM) is the most prevalent form of the disease and recently acknowledged not as a single clinical condition, but importantly, as a group of metabolic disorders. Diabetes in general, causes chronic hyperglycemia and a wide range of downstream metabolic disturbances and multi-organ complications [2]. It is notorious however, that although insulin secretion collapse, peripheral insulin resistance, and/or receptors’ activity failure play a definitive role for the onset of sustained hyperglycemia in T2DM, a large portion of body glucose is cleared by insulin-independent mechanisms, derived from the ability of plasma glucose to influence its own clearance by a mass action effect [3]. T2DM usually most common in adult subjects exhibits a slow, silent and insidious evolution. Hyperglycemia and its adjoining biochemical consequences undermine the whole tissues being sufficient to orchestrate irreversible systemic complications, from which the cells comprised in soft peripheral tissues and vascular structures do not escape. Lower extremities ulcerations and the potential for amputation are currently acknowledged as members of the list of diabetes complications [4].

Surgeon TD Pryce put forward as early as 1887 the link between diabetes and foot ulceration by writing in The Lancet that “...diabetes may of itself be a cause of perforating ulcer [...]” [5]. However, despite the years of efforts and research, the pathogenesis of impaired wound healing in diabetes remains incompletely elucidated [6]. This poor-healing condition appears to be a multifactorial process which includes the amalgamation of systemic and local factors that ensure a perpetual forward loop up to chronification. Along this path, the cells seem to progressively wipe out their ability to trigger evolutionarily imprinted mechanisms as migration, proliferation and transdifferentiation, becoming increasingly static. Thus, diabetic wounds do not only become chronic by a concept of aberrant healing trajectory within a physiological time frame, but also by the asynchrony on the sequence of overlapping events that make up the tissue repair mega-process. Broadly speaking, diabetes impairs most if not all these events. Thus, the challenge that represents the diabetic wound healing failure is the clinical gross expression of an outstanding array of biochemical and cellular disorders [7]. These ideas are supported in the clinical arena by the alarming statistics of amputations around the world every year [8].

The healing process in diabetes is also jeopardized by the patient’s susceptibility to infection due to deficiencies on the innate immunity. Although the diabetic wound bed may be adversely overwhelmed by inflammatory cells, it does not represent an overt anti-bacterial protection. On the contrary, the diversion of glucose to the polyol pathway affects bacterial killing by reducing neutrophil eosinophagocytosis. Furthermore, hyperglycemia-induced reactive oxygen species (ROS) deregulate the innate immunity via an overactivation of the NF-B transcription factor, thus amplifying the absurd inflammation and intoxicating the wound milieu [9, 10]. Peripheral arterial disease, leading to ischemia or lower limb hypoperfusion, is associated with the most severe outcomes, including lower probability of healing, longer healing times, higher probability of ulcer recurrence, greater risk of amputations, and potentially higher mortality [4].

Cells harvested and cultured from hyperperfused granulation tissue orchestrate a molecular program of arrest and senescence (Jorge Berlanga, manuscript in preparation). The outcome of the combination ‘healing failure’ and ‘infection susceptibility’ untowardly contributes to amputation.

Here we have reviewed the current evidences on the toxic resonance of acute and long term exposure to high glucose on the two main cells for the granulation tissue organization: fibroblasts and endothelial cells. We have included a characterization of the organizational disorders affecting diabetic granulation tissue and the challenge that represents its ultimate process, wound re-epithelialization. The literature search was based on English language articles downloaded from PubMed [11] and Bioline International [12] databases.

Consequences of glucose overload toxicity on fibroblasts and endothelial cells

Fibroblasts

Fibroblasts are central to the wound healing process by secreting, contracting and remodeling the extracellular matrix (ECM). They also secrete growth factors as important messengers for mesenchymal-to-mesenchymal and epithelial-mesenchymal communication, especially for establishing the emerging basement membrane and subsequent re-epithelialization. Therefore, any impediment to fibroblast function is detrimental for normal wound healing and may result in chronic, non-healing wounds. The fibroblast, when engaged in fibrogenesis, displays the highly activated phenotype characteristic of myofibroblasts. Although their origin has not yet been definitely elucidated, proliferation of preexisting adjacent dermal fibroblasts, and probably recruited from the bone marrow, has been documented [13]. Under the high glucose burden imposed by diabetes, cutaneous and extracutaneous fibroblasts appear perturbed, and for many years, in vitro models recreating ‘clinical hyperglycemia’ have proved to disrupt normal fibroblasts physiology and derange the secretion of ECM ingredients. These experiments have suggested that high glucose concentration is the proximal detonator of a downstream cascade of molecular disturbances for the skin fibroblasts [14].

Rowe et al., who pioneered the in vitro models, demonstrated that the synthetic, proliferative and secreting capabilities are reduced in diabetics’ cutaneous fibroblasts [15]. Other parallel studies in which high glucose concentrations were introduced, proved to inhibit fibroblast proliferation, while the cells turned resistant to proliferate in response to growth factors such as insulin-like growth factor type-I (IGF-I) and epidermal growth factor (EGF) [16]. Following these attractive targets, Goldstein’s findings allowed for...
establishing the hypothesis that diabetics fibroblasts replicative life span did proportionally decline with diabetics predisposition under normal glucose concentrations, concluding that a persistent, heritable abnormality is present in mesenchymal tissues of overt diabetics and genetically predisposed subjects [17]. Years later, Goldstein also announced that cells obtained from insulin-dependent or insulin-independent diabetics not only exhibit abnormal replicative capacity in vitro, but that the aging process appeared more precociously than in non-diabetic counterparts [18]. Other studies showed that the addition of conditioned media from non-insulin-dependent diabetes mellitus wound fibroblasts induced a dose-dependent inhibition in normal fibroblast proliferation which appeared related to elevated l-lactate levels [19]. This replicative refractoriness of diabetic fibroblasts has been reproduced by different groups in subsequent years [20], thus confirming the need for additional external supplements to ensure cell cycle progression [21]. Accordingly, Loots et al. demonstrated the need of the simultaneous rather than sequential addition of different growth factors combinations for diabetic ulcer fibroblasts in order to induce a proliferative response [22]. Diabetic wound fibroblasts develop a quiescent and senescent phenotype, and their ability for horizontal and vertical migration is also dramatically impaired when compared to normal donor cells in different migration assay as in the modified Boyden chamber haptotaxis assay [23]. Most of these attributes are reproduced under acute exposures to high glucose concentrations so that migration speed is reduced by ~40%. This is associated to a decrease in cell directionality and to non-productive protrusive events (e.g.: loss of cell polarization), consistent with the increased activity of Rac1 and the projection of multiple lamellipodia. This experiment concluded that the generation of ROS may lie behind these abnormalities as they were partially or completely rescued by treatment with N-Acetyl-Cysteine (NAC) [2]. In contrast to the cellular reactions when exposed to high glucose in vitro, full-thickness wounds induced in non-diabetic pigs exposed to a local hyperglycemic environment exhibited no difference in wound closure when compared with normoglycemic controls [24]. This suggested that delayed wound healing by diabetics is a far more complex phenomenon than circumscribed to the high-glucose concentration itself [24]. As a consequence of the cutaneous accumulation of advanced glycation-end products (AGE), the skin increases its chronological age. One of the AGE precursors is 3-deoxyglucosone (3DG). Fibroblasts cultured on 3DG-treated collagen reduce the ability to migrate efficiently since 3DG increases its adherence to the matrix. Additionally, the authors described a higher level of misfolded proteins [25]. Using the same experimental system, this group demonstrated two years later that the inhibition in fibroblast migration, proliferation, and collagen expression by exposure to 3DG-collagen was mediated via extracellular regulated kinase 1/2 (ERK1/2) and the protein kinase B (Akt) downregulation through activation of p38MAPK (mitogen-activated protein kinase). These findings indicated that p38 is a key signaling molecule that plays an opposite role during times of cellular growth and cellular stress [26]. Enriching the above findings, this group also demonstrated that 3DG-modified collagen induces oxidative stress, endoplasmic reticulum stress and apoptosis via caspase-3 activation. Oxidative stress appeared dependent on the upregulation of the NAD(P)H oxidase 4 (Nox4), a ROS Nox homologue, which appeared activated by p38MAPK. Proximal to this cascade is the effect caused by the interaction of the modified collagen with 3DG, which signals to the fibroblast by interacting with integrins alpha-1/beta-1 (α1β1) and not through the canonical AGE receptor (RAGE) [27]. Another group has also shown the induction of cutaneous fibroblasts apoptosis through cytoplasmic and mitochondrial pathways by plating the cells in an AGE-enriched environment made up by Nc-(carboxymethyl)lysine (CML)-collagen, which primarily activated the classic RAGE [28]. A subsequent study elegantly demonstrated that after AGE-RAGE interaction, ROS generation increases, activating both nitrogen reactive species and ceramides, which in turn activates p38 and the c-Jun N-terminal protein kinase (c-JNK). Activated p38 and c-JNK triggers a cascade leading to amplified caspase-3 activity, whereas activation of Forkhead box O class 1 (Foxo1) increases the likelihood of apoptosis through enhanced expression of proapoptotic genes [29]. Under a number of circumstances, Foxo transcription factors induce the expression of BIM and other pro-apoptotic genes.

In addition to the deleterious effects of glucose and its derivatives, diabetic fibroblasts exhibit particular features. Literature documents that diabetic mice fibroblasts show a severe impairment in VEGF production under normoxic and hypoxic conditions as well as a reduced pro-degradative activity due to the high expression of metalloproteinase type 9 (MMP-9) [30]. Similarly, diabetic pigs exhibited an impaired healing that was accompanied by a reduction of IGF-I in the wound milieu [24]. Studies with human fibroblasts have confirmed the pro-degradative phenotype by the increased MMP-2 and MMP-3 production and reduced collagens gene expression [31]. Human diabetic fibroblasts also showed a failure in nitric oxide (NO) production which was concomitant to elevations of nitric oxide synthase 26. The fact that these fibroblasts fail in secreting NO is particularly negative given its role for wound healing. Conversely, NO donors’ administration has shown to stimulate cell proliferation and restore the balance of MMPs [33].

It seems that amplification of oxidative stress acts as a primary culprit in harming fibroblasts biology in diabetes, involving electron transport in mitochondria. High intracellular glucose levels increase the electron transport chain in mitochondria during oxidative respiration, leading to formation of O₂⁻ and the generation of superoxide anion [34]. Chronic hyperglycaemia-induced mitochondrial ROS stimulate various signaling pathways that amplify inflammation and cell death. They include protein kinase C, c-JNK, and p38MAPK [33]. According to an excellent review by Duncanson et al. group [35], ROS leads to the activation of members of the Foxo family. This is a family of transcription factors that regulate gene expression for different functions in health and disease, including metabolism, aging, and stress response. In diabetes, the hyperglycaemic environment causes an increase in the production of ROS, which in turn activates these transcription factors, leading to downstream effects like increased inflammation and cell death.

24. Velander P, Theopold C, Hirsch T, Bleiziffer O, Zuhaili B, Fossum M, et al. Fibroblasts from chronic diabetic wounds show a severe impairment in VEGF production under normoxic and hypoxic conditions as well as a reduced pro-degradative activity due to the high expression of metalloproteinase type 9 (MMP-9) [30]. Similarly, diabetic pigs exhibited an impaired healing that was accompanied by a reduction of IGF-I in the wound milieu [24]. Studies with human fibroblasts have confirmed the pro-degradative phenotype by the increased MMP-2 and MMP-3 production and reduced collagens gene expression [31]. Human diabetic fibroblasts also showed a failure in nitric oxide (NO) production which was concomitant to elevations of nitric oxide synthase 26. The fact that these fibroblasts fail in secreting NO is particularly negative given its role for wound healing. Conversely, NO donors’ administration has shown to stimulate cell proliferation and restore the balance of MMPs [33].
with apparently opposing roles that may defend cells against oxidative stress but also promote cell-cycle arrest in G1 by inducing the cell cycle inhibitory protein p27kip1 [36]. Foxo1 activation appears elevated in diabetic connective tissue cells and mediates AGE and tumor necrosis factor-alpha (TNF-α)-induced apoptosis, both of which are abundant in diabetic connective tissue [37]. Foxo1 limits wound healing by inhibiting fibroblasts proliferation and enhancing their apoptosis [37, 38]. Interestingly, insulin inactivates Foxo1 via Akt leading to its nuclear export and degradation. Defective insulin action in the skin has been proposed as an important mechanism contributing to wound healing defects in diabetes. Perhaps the assorted constellation of the hormone’s pharmacological bounties (increased expression of endothelial nitric oxide synthase (eNOS), vascular endothelial growth factor (VEGF) and stromal-derived factor-1) observed in experimental and clinical wounds when insulin is topically administered, may be attributable to Foxo1 neutralization. Curiously, the acceleration of wound healing occurs in parallel to a local recovery in the expression of proteins involved in insulin signaling pathways [39]. Aside from the above arguments, these preclinical and clinical findings are not surprising in light of the potent anti-inflammatory, pro-anabolic and cytoprotective actions of insulin [40] which extend beyond the exclusive regulation of glucose homeostasis [3].

- Despite the prolific investigation conducted so far, still questions remain to be answered in relation to ex vivo diabetics’ fibroblasts behavior:
  - Why do diabetics’ fibroblasts evoke behavioral traits in culture, mirroring the donor’s tissue, even when grown under optimized oxygenation, nutrient, growth factors, and glucose supply?
  - Is there any sort of ‘behavioral imprinting’ so that they are reminiscent from a diabetic donor?
  - Why can cultured fibroblasts from both ischemic and neuropathic ulcers exhibit different ultrastructural morphology and organize the monolayer in a private manner?
  - Is there any epiphenomenon beyond the irreversible glycation sustaining the ‘impersonation’ of the in vivo traits?

**Endothelial cells**

Angiogenesis is a comprehensive term which indicates the physiological process involving the growth of new blood vessels or neovascularization. This is a vital process for embryological growth, tissue development, and wound healing. Different growth factors families regulate angiogenesis in collaboration with other proteins, by promoting endothelial cells recruitment, proliferation, migration, co-opting and collar stabilization. Among the growth factors are: VEGFs, fibroblast growth factor, angiopoietins, platelet-derived growth factor (PDGF) and transforming growth factor-β (TGF-β). Proteins comprise integrins, cadherins and ephrins. There is an enormous and ever-growing body of evidence indicating the close correlation between hyperglycemia and the abnormalities in endothelial function and morphology [41]. The UK Prospective Diabetes Study (UKPDS) [42] and Diabetes Control of Complications Trial (DCCT) [43] found microvascular disease and hyperglycemia to be intrinsically related. Thus, anomalous angiogenesis is a hallmark of both type forms of diabetes which is clearly and early observable during the process of granulation tissue growth; condition that has been successfully reproduced in animal models [44]. For subjects with macrovascular disease, the defective angiogenesis prolongs and disturbs the healing process. The concept of abnormal angiogenesis extends beyond the wound, given the inability of these patients to create collateral circuits due to a VEGF-dependent monocytes dysfunction [45]. Furthermore, insulin has a dramatic impact on the endothelial homeostasis by its ability to stimulate NO release via a cascade that involves activation of the phosphatidylinositol 3-kinase (PI3K)-Akt signaling and eNOS phosphorylation. The later is of paramount importance in angiogenesis and wound healing as described below [46].

As depicted for fibroblasts, high glucose and the glycated-by-products exert a dramatically toxic effect on endothelial cells and the vascular wall in general. In parallel, the endothelial cells per se seem to be a very sensitive target to high glucose. Endothelial dysfunction is intricately related to insulin resistance through the stimulatory effects of insulin on glucose disposal and NO production in the endothelium. Today, vascular dysfunction remains as a major cause of morbidity, amputation/disability and mortality in diabetic patients. Even after achieving the successful reperfusion of an ulcerated lower extremity, the healing process is slow and torpid. Therapeutic angiogenesis has been pursued for years but the clinical results have shown relatively limited outcomes [47-49].

High glucose concentrations have been associated with endothelial metabolic dysfunction in vitro and in vivo and as for multiple physiological processes; insulin and its downstream signaling regulate most of the endothelial cell functions [50]. High glucose ambient has been shown to disturb endothelial cells cycle, increase DNA damage, delay endothelial cells replication and induce excessive cell death [51]. In addition, high glucose also prevents NO-induced inhibition of vascular smooth muscle cells migration [29] thus contributing to Monckeberg’s media thickening.

In vitro models simulating ‘normoglycemia’ and hyperglycemia have demonstrated that under high glucose ambient, proliferation and tube formation of dermal microvascular endothelial cells appear impaired [52]. Furthermore, high glucose levels selectively trigger apoptosis in cultured endothelial cells as has been demonstrated by different laboratories [53]. High glucose induces the up-regulation of TNF-α level concomitant to the death receptors TNF-R1 and Fas in a variety of cultured endothelial cells [54]. Under this ambient, the expression of the proapoptotic Bax protein increases, cytochrome c is released, subsequently conjugating to the apoptotic protease activating factor-1 and triggering a caspase cascade-induced death [55].

Hyperglycemia-induced oxidative stress promotes inflammation through increased endothelial cells damage, microvascular permeability, and uncontrolled release of proinflammatory cytokines, including TNF-α, interleukin-1β (IL-1β), and interleukin-6 (IL-6), ultimately leading to decreased insulin sensitivity and...
diabetic vascular complications. Moreover, hyperglycemia-induced Foxo1 also plays an important role in the induction and amplification of proinflammatory cytokines production. Foxo1 directly binds to IL-1β promoter and increases its expression in macrophages [56].

Hyperglycemia and the accumulation of AGE disturb the role of angiogenic growth factors as VEGF, its receptor, its signaling pathway; thus disrupting endothelial proliferation, migration, and endothelial progenitor cells (EPCs) recruitment and release from bone marrow [57]. Insulin resistance disrupts the NO-mediated angiogenic positive regulation over angiogenic growth factors such as VEGF, fibroblast growth factor and TGF-β [57]. Studies using streptozotocin-induced diabetic mice with simultaneous hind-limb ischemia have suggested that the angiogenic responses remain preserved even under the diabetic state, and that 40 to 50% reduction of PDGF-BB expression is responsible for the induction of functional and morphological vascular abnormalities and pericyte apoptosis. Conversely, PDGF-BB external supplementation was sufficient to prevent limb autoamputation, also reproduced with a protein kinase C inhibitor that restored the expression of endogenous PDGF-BB [58].

The glycation of collagen and other proteins within the wound ECM and AGE accumulation bring catastrophic consequences for the angiogenic reaction with inhibition of angiogenesis in vivo. The fact that angiogenesis is restored by aminoxyanidine treatment reinforces the antiangiogenic role of AGE [59]. Angiogenesis is a multifaceted process demanding an appropriate, non-glycated extracellular substrate. This is clearly illustrated by the fact that PDGF-BB anchors to different components of the ECM under physiological conditions acting as a natural depot and slow release system for the growth factor. Local PDGF unavailability has proved to impair the coverage of newly formed vessels with mural cells and local pericytes [60]. These evidences reinforce the pathophysiological impact of high glucose toxicity, the release of pro-inflammatory cytokines and the activation of the intrinsic mitochondrial-mediated apoptotic signaling pathway on endothelial cells. In summary, endothelial cells exposed to excess glucose develop a pro-inflammatory profile, becoming a significant source of cytokines and ROS production. The agonistic stimulation of the RAGE is able to mount the same response leading to apoptosis and vascular run. The pathogenic effects of hyperglycemia on fibroblasts and endothelial cells are summarized in figure 1.

Compelling evidences indicate that at least a portion of the hyperglycemia-mediated endothelial damages and dysfunctions are associated with an impaired mitochondrial activity resulting in mutations of mitochondrial DNA, due to a disproportionate ROS production, leading to an inflammatory reaction and apoptosis [61]. As a matter of fact, mitochondrial DNA has a much higher mutation rate than nuclear DNA because it lacks histones and is exposed to the direct action of oxygen radicals while its repair system is limited. Therefore, ROS appear to play a pivotal role in systemic endothelial deterioration and biological aging [62]. As described, ROS generation enhances Foxo1 activation and induction of several classes of genes that regulate endothelial cell behavior, including pro-inflammatory factors, and eventually the execution of apoptosis of endothelial and adjacent cells [35]. ROS-mediated lipid peroxidation appears to impair most healing events, contributing to growth factors reduction, keratinocytes migration failure, slow or torpid fibroplasia, delayed contraction and matrix remodeling, for not to mention aberrant angiogenesis [63]. Under experimental conditions, the pharmacological intervention with a chemical inhibitor of lipid peroxidation proved to reduce the local edema and to stimulate re-epithelialization, neovascularization, proliferation of fibroblasts, and synthesis and maturation of the ECM. A parallel finding was the normalization of VEGF mRNA expression and secretion in those diabetic mice. This further supports the view that lipid peroxidation perturbs VEGF production [64]. An extraordinary background has accumulated about the role of NO in vascular biology in diverse horizons as ischemia, inflammation and neovascularization. Impaired endothelium-dependent NO-mediated relaxation occurs in both cellular and in vivo models [65]. Many of the metabolic conditions associated with diabetes are conditioned by failure in NO synthesis or its degradation. In this respect, the integrity of the Akt/eNOS coupling pathway for a normal endothelial function appears compulsory [66]. Hyperglycemia is also associated to a deficit in tetrahydrobiopterin (BH4) and to an increase in arginine expression, which attempt against NO synthesis and normal endothelial functions such as vascular remodeling responses [67]. The increased generation of peroxynitrite levels under high glucose conditions contributed to deplete cellular anti-oxidant reserves as to activate the NF-kB transcription factor and consequently the expression of the inducible form of nitric oxide synthase (iNOS), intercellular adhesion molecule-1 and other inflammatory mediators [68].

Figure 1. Pathogenic consequences of hyperglycemia on fibroblasts and endothelial cells. AGE-RAGE: glycation-end products (AGE) interaction with their receptors (RAGE); ROS: reactive oxygen species; MAPK: mitogen-activated protein kinase; c-JNK: c-Jun N-terminal protein kinase; NF-κB: nuclear factor kappa beta. Thick arrow indicates increased protein activity.

EPCs are active players for the maintenance and repair of endothelial cells. They participate in angiogenesis as they proliferate, migrate and differentiate, and are a source for proangiogenic factors and cytokines [69]. Multiple evidences indicate that the number of circulating EPCs is decreased under both clinical forms of diabetes, which is likely to be involved in the pathogenesis of vascular complications [70]. Under experimental diabetic conditions the EPCs number appears significantly decreased in the bone marrow as in the peripheral blood which was reverted by treating the mice with insulin [71]. In general the bone marrow derived EPCs in the diabetic patients are considered as dysfunctional, producing fewer endothelial cells with reduced replicative, and migratory potential [72]. Tamarat et al. have described a limited capacity of diabetic animals-derived bone marrow mononuclear cell to differentiate into EPCs in vitro as to organize tubulogenetic structures when subcutaneously implanted in a matrigel plug, thus hindering the revascularization of damaged areas [70]. Over again, the activation of p38/MAPK mediated by an excessive ROS generation has been identified as responsible for the EPCs impaired proangiogenic potential in vivo by limiting cell proliferation and differentiation [73]. To fully divert the physiological role of EPCs in tissue repair and angiogenesis, the dext hyperglycemia-ROS stimulates EPCs to produce pro-inflammatory cytokines and to shift NO production by elevating iNOS and decreasing eNOS [74]. As described for other cells, AGE treatment disrupts EPCs physiology thus leading to a downregulation of eNOS and the anti-apoptotic factor Bcl-2 expression, as well as an elevation in cyclooxygenase-2, proapoptotic factor Bax, NF-κB and caspase-3 in a MAPK (ERK/p38/c-JNK)-dependent manner [73].

Angiogenesis is of paramount importance for wound healing, but diabetes-mediated vascular cells damages are as varied and broad as it is the concept of systemic endothelial dysfunction. Diabetes distorts the angiogenic program so badly that angiogenic factors are defective whereby and when required (lower limbs), and overproduced in erroneous anatomical niches with fatal consequences for the patient (diabetic retinopathy).

Failure of granulation tissue onset and progression

Once described the main consequences of high glucose/hyperglycemia on the two principal architects of granulation tissue, fibroblasts and endothelial cells, the most distinguishing features for the onset of the granulation process in diabetic cutaneous wound healing will be recapitulated next.

Tissues’ regenerative capabilities have been neglected along the species evolution; thus, scarring process has emerged as an urgent alternative to favor the structural and functional restoration of a wounded zone. Within these events, the process of granulation tissue formation is pivotal as it constitutes a sort of living, temporary aggregate of cells and proteins, acting as a welding material until the tissue’s continuity is restored. However, the reluctance to trigger and sustain the out-growth of a productive granulation tissue with an appropriate ECM is typical in diabetic patients, and particularly if ischemia concurs. As mentioned, these wounds are characterized by a proliferative arrest, pro-inflamed, pro-oxidant and pro-degradative phenotype [75].

This stubbornness and slowness to heal in diabetics is conditioned by systemic and local factors that in complicity counteract intrinsic reparative mechanisms. In a broad systemic context, inflammation and anabolic deficit can be conceptually mentioned. Diabetic patients with foot ulceration bear a specific and ordered alteration of the immune status with an active upregulation of circulating levels of acute-phase proteins, cytokines, and chemokines that impose a chronic systemic inflammatory profile, and amplify local wound inflammatory networks [76].

The systemically elevated levels of pro-inflammatory response markers and the wound’s expression of cytokines and chemokines are among the culprits of the abnormal repair mechanism [77]. Another factor to be considered is that diabetes per se is a metabolic disease in which fuels metabolism is perturbed given the rupture of one of the most important anabolic axis of the organism: insulin/IGF-I. The role of insulin in wound healing is well known by its anabolic effect on wound protein balance, favoring synthesis and preventing degradation [78, 79]. IGF-I has a similar effect on stimulating wound tissue anabolism. Both insulin and IGF-I appear to act in part by the induction of the anabolic transcription factor ATF4 (CREB2), essential for the activation of the mammalian target of rapamycin complex 1 (m-TORC1) protein which in turn is required for protein synthesis via Foxo-dependent gene expression [80]. We do not rule out that the diabetes-concomitant deficit of incretins could participate in the negative anabolic balance observed in such wounds. Glucagon-like peptide-1 in addition to its anti-hyperglycemic actions is endowed with a vast number of multi-organ cytoprotective, trophic and anti-inflammatory effects [81]. In support to the glucagon-like peptide-1 action is the study by Ta et al. with alogliptin, a specific inhibitor of dipeptidyl peptidase-4, which was shown to inhibit macrophage-mediated inflammation response and was suggested as tissue remodeling promoter by inhibiting the expression of different MMPs [82].

Rapid formation and deposition of an appropriate ECM, in particular by fibroblasts, is required for an efficient cellular anchoring and homing at the wound bed. As mentioned above, the cutaneous fibroblast is a cell type sensitive to high glucose, AGE-precursors, AGE, ROS and TNF-α, rapidly undergoing premature senescence, arrest or apoptosis. Fibroblasts are the main source of collagen, and the number of fibroblasts can be taken as a measure of repair based on their collagen synthesis ability. It is very likely that the deficit of growth factors such as TGF-β1, IGF-I, and PDGF, that stimulate fibroblasts proliferation, transdifferentiation and the synthesis of matrix components, appear in shortage in diabetic foot ulcers (DFUs) and results in a scarce ECM formation. Numerous growth factors (TGF-β1, IGF-I, PDGF) are able to regulate the balanced expression of MMPs and tissue inhibitors of metalloproteases, while most of them exhibit an altered expression in DFUs [83]. Moreover, the imbalance in the DFU milieu between TGF-β1 and TGF-β3 in which the former appears downregulated, may
explain fibroblasts quiescence in terms of proliferation and secretion [84]. This phenomenon represents the deficit of one of the most potent pro-fibrogenic and fibroblasts-mitogenic growth factors, which at the same time is able to downregulate macrophage activation [85].

The ECM represents the granulation tissue dynamic stroma that provides support for inflammatory cells, fibroblasts, and endothelial cells and allows for the chemotaxis of epithelial cells, thus hosting the re-epithelialization process [86]. One of the main challenges for the diabetic wound healing is the structuring of a normal matrix in quality and quantity. In general, a poor ECM formation distinguishes DFUs which can result from: i) diminished ECM synthesis, ii) increased ECM degradation rate by proteolytic enzymes, iii) toxicity due to glycated by-products accumulation, and iv) toxicity by biofilm bacterial contaminants diffusion [87]. We deem that an important cause of the clinical dilemma of the high rate of re- ulcerations and ipsilateral amputations in DFU patients’ shortly after re-epithelialization [88] may be inherent to the qualitative composition of the scar ECM to tolerate tensile forces and mechanical stress.

The diabetic granulation process does not generally exhibit the sequential cascade of events that characterize normal wound healing. This has been confirmed through the histopathological analysis of granulation tissue biopsies by Loots et al., who described the lesions as ‘frozen’ in a chronic low-grade inflammatory state associated to a scarce provisional ECM [89]. Our group’s serial biopsies from both neuropathic and ischemic ulcers-derived granulation tissue have identified histological differences for both types of wounds in the absence of clinical infection. Polymorphonuclear cells (PMN) infiltration is intense and prolonged particularly in neuropathic wounds, co-existing with a scarce ECM accumulation in which collagen deposit is impoverished (Figure 2).

Under more mature stages, the neuropathies may also show an abnormal sprout of new small vessels and capillaries that may derive not from a normal angiogenic response but due to arterio-venous shunts. Our observations recall those of Black et al. who demonstrated that in neuropathic patients there exists a decrease in fibroblast proliferation and a scarce amount of collagen accumulation within the wound bed [90]. On the contrary, a broadly spread infiltration of round cells predominate in those patients suffering from wound bed ischemia, associated to a fibro-hyaline matrix of ‘hardened’ aspect and abnormal angiogenesis in which vascular wall cellular mosaicism, precocious media thickening, endothelial nuclei hypertrophy and many other defects can be identified (Figure 3). It is likely that the combination of arterial hypoperfusion and glucose toxic derivatives imprints a particular pattern of damage to the morphogenesis of vessels in the wound [91]. These observations invite to speculate that the biochemical microenvironment in ischemic and neuropathic diabetic wounds is different, and that the inflammatory ‘badge’ is in correspondence with the wound’s most prevalent pathogenic component [92]. In contrast to acute wounds in non-diabetic subjects, the inflammatory reaction in diabetics appears prolonged [93] and sharply delays granulation tissue formation and maturation [94]. Data derived from murine diabetic models indicate that the exaggerated inflammatory reaction is related to the prolonged expression of macrophage inflammatory protein-2 and macrophage chemoattractant protein-1 [95]. Furthermore, the downregulation of the anti-inflammatory cytokine IL-10 in DFUs environments represents the collapse of an important inflammatory restrainer [77]. Other evidences indicate that PMN are critical toward the acquisition and perpetuation of inflammation and a degradative phenotype. The granulocytes secrete TNF-α and IL-1β which act as a triggering signal for MMPs expression via the common NF-κB signaling pathway. Within the wound context, TNF-α stimulates its own secretion and that of IL-1β, contributing to a persistent inflammatory status [96]. TNF-α has proved to negatively impact the repair process as it is early secreted since the inflammatory phase. Its deregulation is not only associated with persistent inflammation but also to connective tissue degradation [97]. Concomitantly, TNF-α mediates its antagonistic effects on TGF-β through the c-JNK pathway via inhibition of Smad phosphorylation, consequently reducing the expression of TGF-β1, and that of several

Figure 2. Common histological aspect of a neuropathic granulation tissue. It exhibits a scarce deposition of extracellular matrix. Note a central blood vessel with abundant surrounding cellularity and fibrin, suggesting hyperpermeability. Hematoxylin/eosin staining, 10 × magnification.

Figure 3. Common histological aspects of ischemic granulation tissue. Note the presence of an intense inflammatory infiltrate of round cells spread all over the tissue area. The emerging vessels appear dysfunctional with thickened walls of fibrohyaline material, ‘hardened aspect’ and endothelial nuclei hypertrophy. Hematoxylin/eosin staining, 10 × magnification.
downstream matrix proteins [98]. In this highly pro-
teolytic milieu, fibronectin, collagens, growth factors and their receptors are degraded while the wound is way down to a catabolic state [99]. Importantly, the perpetuated homing of PMN within the wound bed is associated to high local levels of elastase secretion, ROS and reactive nitrogen species [100]. High circulating and PMN-associated elastase levels are attribut-
able to a poor glycemia control and are considered a risk marker for the development of diabetic angiopa-
thy [101]. Fibronectin degradation, for instance, is referred as one among the several causes of diabetic re-epithelialization failure. Epidermal keratinocytes require of the interaction between fibronectin and its surface receptor integrin α2β1 to effectively migrate [102]. Curiously, insulin-degrading activity has also been demonstrated in the fluid of diabetic experimental and human wounds which have been shown to correlate with the glycated hemoglobin levels [103]. The connection between NO metabolism and foot ulcer protease profile has been described. In contrast to elevated MMP-8 and -9 displayed by the non-healing diabetic foot wound, the concentration of NO appears significantly reduced. Diabetic skin fibroblasts treated with NO donor compounds selectively raised NO production, increased cell proliferation, and decreased the expression of MMP-8 and -9 in a dose-dependent manner. Thus, that NO resumes the cell proliferation program and promotes the reestablishment of an anti-proteases effect have emerged as argument in favor of the NO salutary effect in wound healing [32].

The link between wound cells and apoptosis was described above; we just wish to comment that in sharp contrast to wound-infiltrated inflammatory cells becoming refractory to apoptosis, granulation tissue-producing cells are sensitive to commit suicide where TNF-α stands as a major driving force. The negative impact of TNF-α levels on the sensitivity of tissues to insulin has been consistently documented. Skin cells are not excluded from this effect [104]. Conclusively, any therapeutic approach aimed to neutralize TNF-α or to increase the wound local availability of active TGF-β1 would be similarly effective for stimulating granulation tissue and wound closure [105].

Chronic wounds and especially diabetic foot ulcers exhibit a highly pro-oxidant microenvironment that amplifies the cytotoxic cascade. Endothelial cells and fibroblasts, in particular senescent fibroblasts, are a prominent source for oxygen radicals, but at the time they turn into these radicals targets which, by converging mechanisms, arrest cell proliferation and induce apoptosis [106]. Thus, the disturbed oxidant/ antioxidant balance as the AGE accumulation within the chronic wound microenvironment is considered a major factor amplifying the unrestrained and persist-
ent inflammatory, toxic and catabolic state of non-
healing wounds [100].

The failure of wound contraction is a clinical hall-
mark of diabetic granulation tissue. Fibroblast-to-
microfibroblast transdifferentiation represents a key event during wound healing and tissue repair. The contractile force generated by myofibroblasts as a highly specialized cell, speeds the healing process of dermal wounds in healthy humans, accounting for an 80-90% of scar tissue reduction [107]. In addition, the contraction process reduces the area to be resur-
fac'd by re-epithelialization which represents a sort of ergonomic response. In diabetic subjects however, contraction is impaired and deep ulcers heal by the combination of granulation and re-epithelialization. The classical view on dermal wound healing implies recruitment of local fibroblasts [108], followed by a subsequent process of transdifferentiation in which the fibroblasts gains a definitive phenotype of differenti-
ated myofibroblasts by neo-expressing the α-smooth muscle actin (α-SMA). Nevertheless, α-SMA ex-
pression is precisely controlled by the joint action of growth factors like TGF-β1 and ECM proteins like the fibronectin splice variant ED-A, as by the local mechanical microenvironment [108]. It should be noted however, that indwelling fibroblasts in diabetic wounds are refractory to proliferate, adopt a senes-
cent phenotype, and that TGF-β1, fibroblast and other matrix proteins may appear in deficit. Hence, all these factors may contribute to the poor contrac-
tile activity. Furthermore, Goldberg et al. showed that among the deleterious activities of TNF-α within the wound is to suppress α-SMA expression in hu-
man dermal fibroblasts [98]. Figure 4 integrates the cascade of deleterious factors that impact on diabetic granulation tissue onset.

If the animals-derived evidence that a high fraction of the wound myofibroblasts potentially derives from bone marrow fibrocytes is valid for humans [109]; then we have already learned that diabetes negatively impacts on the general bone marrow physiology [110]. Beyond this stromal-derived factor-1α (acting as re-
cruting factor) together with its CXCR4 chemokine receptor being also impaired [111]. Finally, it has been documented that the circulating acute inflam-
matory reactants involved in insulin resistance inhibit

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Figure 4. Impact of hyperglycemia on granulation tissue biology. Thick arrow represents the unles-
shment of a novel amplification pathogenic loop in which the tumor necrosis factor α (TNF-α) and the Fas system pulls the trigger against endothelial cell, fibroblasts and myofibroblasts, hampering the granulation tissue growth. AGE-RAGE: glycation-end products (AGE) interaction with their receptors (RAGE); ROS: reactive oxygen species; TGF-β1: transforming growth factor-β1.
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fibrocytes differentiation [112]. There are numerous cellular and molecular aspects unknown and that remain to be answered on the granularity tissue biology, such as: i) What are the molecular and cellular driving forces supporting the microscopic structural differences between neuropathic and ischemic ulcers beds?; ii) What is the explanation for the ‘inherence’ of vascular changes as a dramatic Monckeberg media thickening in nascent arteries within an early hatchling granulation tissue?; iii) Why granularity tissue is histomorphologically abnormal even in metabolically compensated patients?

Re-epithelialization at the clinical level is not a less important problem, as most of the diabetic wounds may granulate in time, while re-epithelialization is even far slower, arrhythmic, and torpid. Re-epithelialization is accomplished through the combined actions of keratinocytes’ dedifferentiation, proliferation and migration, requiring a complex basement membrane which emerges from the mutual interaction between mesenchymal and epithelial cells. Re-epithelialization failure is therefore one of the landmarks of diabetic and other chronic wounds. The epidermal edge of a chronic wound is thick and hyperproliferative, with mitotic pairs the integrin-mediated adhesion of keratinocytes through an akt-dependent on EGFR catalytic activity. Lack of insulin receptor at both the transcriptional and translational levels [120]. Lack of insulin receptor expression derives in reduced skin proliferation and abnormal differentiation in vivo [121]. Furthermore, TNF-α has also been implicated in epithelial cells arrest by deeply perturbing critical elements of keratinocytes’ physiology, including insulin sensitivity [122]. A notorious study provided evidences on the roles of c-myc and β-catenin in impairing epithelial edges migration [123]. Nuclear β-catenin stabilization inhibits keratinocytes migration by blocking EGF response via c-myc induction, and repressing keratins 6 and 16 expression, ultimately depleting the pool of epidermal stem cells at the non-healing edge [123]. It is therefore evident that keratinocytes migration incapability plays an important role in re-epithelialization failure, since cytoskeletal keratins K2, K6 and K10 have been found diminished in DFUs [124]. Moreover, the observation that EGF response was blocked may have further deleterious impact. Many peptide growth factors, including members of the EGF family, accelerate wound re-epithelialization “in vitro” and “in vivo” [125]. Among them, the activation of the EGF family of ligands and its receptor (EGFR) is of physiological significance. Furthermore, EGFR expression is transiently increased at wound margins, suggesting its active role in wound repair. EGF stimulates both cell proliferation and motility [126], the later being dependent on EGFR autophosphorylation and the subsequent activation of phospholipase Cγ-1. On the other hand, EGFR activation also leads to membrane ruffling and focal adhesions through activation of members of the Rho subfamily of GTP-binding proteins [127]. Recent experiments document the negative effect of null mutation of the Slug transcription factor on its function as a downstream EGFR catalytic mediator for wound re-epithelialization. Thus, any interference with the EGFR cascade will hamper epidermal resurfacing [128]. Classic experiments provide illustrative examples on the relevance of the epithelial-mesenchymal cross-talk and on the irreplaceable role of growth factor as a networking bridge [129] for re-epithelialization. Skin-reconstitution studies have shown that bone marrow stromal cells, in addition to dermis-localized preadipocytes and fibroblasts, distinctively promote epidermal regeneration [130]. As diabetes courses with a deficient secretion of growth factors and other chemotactic mediators in areas of tissue repair, recruitment of circulating stromal cells appears reduced, which may turn into an additional hit to that of high glucose-associated toxicity [131]. At the end, there are so many factors which may interact as to obstruct chronic wounds re-epithelialization that it may turn into a puzzle. Above all, two questions from the clinical practice remain: Why do keratinocytes become stunted and arrested again soon after resuming migration following wound contours debridement? Why do the biopises invariably show a hypertrophic lip of cells in vertical downward growth in spite of a horizontal polarization?

Concluding remarks
Our current understanding on the molecular mechanisms impairing wound healing in diabetes simply sizes the tip of the iceberg. Diabetes per se is a complex disease. Even more complex is the group of hard-to-name members that constitutes the type 2 form. This is ethiopathogenically multifactorial and
behaves as individual as the affected subject is. So, it is the pattern of clinical complications, including the wound itself. To date, all the evidences aim to high glucose burden as the proximal trigger to unleash acute and chronic self-perpetuating loops, which include but are not limited to ROS-lipid peroxidation, hyperinflammation/disimmunity, AGE-RAGE toxicity and mitochondrial dysfunction. All these factors enforce precocious senescence, arrest and apoptosis.

At both, experimental and clinical levels, the diabetic wound phenotype is the expression of countless molecular factors that orchestrate a complex biochemistry and an aberrant cellular behavior. The pathway to chronicization has not been fully elucidated but by all means it represents a form of cells’ biological disobedience and entails the need of continuous surgical “cutting” in order to transiently restore an acute behavior by ‘refreshing’ the cellular environment. Despite scientific and clinical efforts, and resources investments, amputation rates have not attained a significant reduction since the emergence of first line technology pharmaceuticals or sophisticated devices. Not to mention the dismal rates of local re-ulceration once the lesion was re-epithelialized. Since limb salvage is always a worthwhile goal and diabetic foot ulceration involves predisposing factors that can be diagnosed and clinically graded; primary care ulcer prevention plans, podiatric assistance, optimal glycemic control and educational programs, will be always far more socially rewarding.