RAGE in the pathophysiology of skeletal muscle

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Abstract

Emerging evidence suggests that the signalling of the Receptor for Advanced Glycation End products (RAGE) is critical for skeletal muscle physiology controlling both the activity of muscle precursors during skeletal muscle development and the correct time of muscle regeneration after acute injury. On the other hand, the aberrant re-expression/activity of RAGE in adult skeletal muscle is a hallmark of muscle wasting that occurs in response to ageing, genetic disorders, inflammatory conditions, cancer, and metabolic alterations. In this review, we discuss the mechanisms of action and the ligands of RAGE involved in myoblast differentiation, muscle regeneration, and muscle pathological conditions. We highlight potential therapeutic strategies for targeting RAGE to improve skeletal muscle function.

Keywords RAGE; AGES; S100B; HMGB1; myogenesis; muscle wasting

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Introduction

Skeletal muscle is the most abundant tissue and endocrine organ of the human body1,2 playing vital roles in breathing, posture maintenance, locomotion, whole-body metabolism, and reservoir of glucose and amino acids that can support energy demand during extreme metabolic perturbation. Skeletal muscle is a highly plastic tissue that can adapt its mass and functionality in response to different conditions, including ageing, diabetes, chronic and metabolic disorders, immobilization, inflammatory myopathies, and various genetic muscle disorders, such as muscular dystrophy.3,4 Due to its essential role, skeletal muscle wasting accelerates disability and mortality in various disease conditions.

RAGE (Receptor for Advanced Glycation End products) was purified from bovine lung homogenates in 1992, thanks to its ability to bind and mediate the effects of Advanced Glycation End products (AGEs), which are produced by the non-enzymatic glycation of lipids or proteins upon exposure to reducing sugars.5 RAGE is a multiligand receptor considered a key mediator of several physiological (e.g. tissue differentiation and regeneration/repair and resolution of inflammation) and pathological (e.g. inflammation, diabetes, cardiovascular diseases, neurodegeneration, and cancer) processes through the activation of multiple cellular signalling cascades, depending on the cell type, the ligands involved, and the density of the receptor on the cell surface.6,7

The intracellular signalling cascades activated by RAGE change in response to environmental cues (i.e. the presence and concentration of RAGE ligands) in many systems, including skeletal muscle tissue. Indeed, an appropriate recruitment of RAGE concurs to skeletal muscle development and restoration of muscle homeostasis in physiological conditions and upon acute muscle injury, respectively.8,9 On the other hand, over-stimulation of RAGE concurs to muscle damage and altered muscle metabolism in ageing and pathological conditions, such as metabolic perturbations and myopathies, all characterized by chronic inflammation and increased production of reactive oxygen species (ROS). Therefore, the fate (i.e. proliferation, differentiation, or death) of muscle precursor cells, myofibre trophism, and the success of muscle regeneration appears to be strongly dependent on the extent of RAGE activity and the availability of specific ligands. In the current review, we summarize information about the role of...
RAGE as foe or friend in muscle tissue and raise questions about RAGE as a potential target in the prevention and treatment of muscle wasting.

The receptor RAGE

RAGE structure

RAGE is a member of the immunoglobulin (Ig) superfamily, which includes Igs, cell surface receptors, and adhesion molecules. In humans and mice, RAGE is encoded by \textit{AGER} and \textit{Ager}, respectively, which localizes in both species next to the HLA (human leukocyte antigen) locus on chromosome 6, in close proximity of the MHC class III complex. RAGE is highly conserved across different mammalian species at both DNA and protein level. Despite its name, however, RAGE is a multiligand receptor recognizing high mobility group B1 (HMGB1), certain S100 proteins, and other compounds besides a number of AGEs (see below). In humans, RAGE is a 55-kDa cell surface receptor whose mature form consists of three regions: an extracellular region of 321 amino acids, a single hydrophobic transmembrane-spanning domain of 19 amino acids, and a short cytosolic tail of 41 amino acids. The extracellular region is divided into three Ig-like domains: one N-terminal V-type (variable) and two C-type (constant, C1 and C2) domains (Figure 1). V and C1 domains form a structural and functional unit (VC1) connected with C2 domain by a short (seven amino acid long) and flexible linker region. Although the VC1 unit binds directly the majority of RAGE ligands, the C2 domain is important for the stabilization of ligands interaction.

Figure 1 Schematic representation of fl-RAGE structure and its of ligands. The dominat negative isoform of RAGE (RAGEΔcyto) lacking of cytoplasmatic tail is reported.
Box 1. RAGE isoforms
Twenty-two different variants occurring at mRNA and protein level have been identified in 18 different tissues by real-time PCR assays, 15 of which were named HaspRAGEv1 to v15. The most frequent splice variant, transcript variant 1 (Tv1-RAGE), is the membrane-bound form of RAGE consisting of the three classical domains (fl-RAGE). The other variants are characterized by different exon/intron structure, causing changes in the sequence/function of the corresponding proteins, including changes in amino acid sequence of the extracellular ligand-binding V domain (RAGE_v4 and RAGE_v5); the presence of a potential starting codon at the end of exon 7 producing two short variants lacking a large part of the extracellular domain of RAGE (RAGE_v11 and RAGE_v12); the presence of potential initiating codon in exon 3 producing a transmembrane protein lacking the V domain (RAGE_v2); the inclusion of the intron 1, 4, 6, or 9 translating into RAGE_v2, v6, v14, and v1 variants, respectively; and the deletion of the entire or part of exon 8 resulting in loss of the transmembrane or cytosolic domain (RAGE_v3 and RAGE_v7, respectively). In human brain, a slightly truncated version of fl-RAGE characterized by a partial deletion of exons 10 and 11, resulting in a predicted deletion of 16 amino acids, was also identified. This unique form of RAGE, named RAGEA, possesses an identical extracellular domain to fl-RAGE but lacks a significant amount of the intracellular signal transduction domain. Truncation of the cytoplasmic tail confers a dominant negative effect on RAGE function, impairing its activation by blocking key signalling pathways downstream of RAGE. Several RAGE splice variants are potential targets of the nonsense-mediated mRNA decay pathway and thus are likely to be degraded before protein expression. On the other hand, the mRNA isoforms lacking the signal sequence on exon 1 might be able to undergo protein expression, but the expressed protein could be subject to premature degradation.

The only human variants that have been detected at the protein level in vivo are fl-RAGE, sRAGE, and eSAGE.

Human AGER is composed of 11 exons and 10 introns of variable length and a 3′UTR (untranslated) region. Several splice variants of RAGE have been identified (see Box 1). Those variants that lack both the transmembrane-spanning domain and the intracellular tail likely generate circulating soluble isoforms of RAGE, collectively known as soluble RAGE (sRAGE). Alternative mRNA splicing and proteolytic cleavage of full-length RAGE (fl-RAGE) are the two primary mechanisms giving rise to sRAGE variants, including endogenous secretory RAGE (esRAGE) and cleaved RAGE (cRAGE). esRAGE (or RAGE_v1) is the C-truncated form of RAGE originating from alternative splicing at exon 9 that causes inclusion of part of intron 9 and deletion of exon 10, whereas cRAGE arises from proteolytic cleavage by the membrane metalloprotease, ADAM10, of fl-RAGE at the cell surface (see Box 1).

RAGE ectodomain cleavage (shedding) occurs constitutively, but it can be induced by ligand binding, cell stimulation with phorbol 12-myristate 13-acetate, ionomycin, or cholesterol depletion. sRAGE can act as a decoy receptor that intercepts RAGE ligands, thereby blocking them. sRAGE can be detected in human blood at picomolar levels, and a strong correlation exists between circulating sRAGE levels and several pathological conditions, such as cardiovascular disease, diabetes, hypertension, and dementia/Alzheimer’s disease (AD) (see Box 1).

RAGE ligands
RAGE mediates several physiological and pathological processes through the interaction with a large variety of ligands (Figure 1). AGEs are a group of non-enzymatic adducts of proteins, lipids, and nucleic acids, which form in a time-dependent manner under conditions of enhanced oxidative stress, especially when target molecules turnover slowly and aldose (such as glucose and ribose) levels are elevated. Over 20 different AGE modifications have been characterized including N-carboxymethyl-lysine (CML), pentosidine, N-carboxyethyl-lysine (CEL) adducts, methylglyoxal-derived AGEs, argpyrimidine, and others, which makes AGEs a heterogeneous class of compounds. Formation of AGEs, which occurs naturally during ageing or in conditions of hyperglycaemia and oxidative stress, leads to the activation of different signalling pathways mediated by a series of cell surface receptors, the most studied of which is RAGE. As AGEs are heterogeneous, it is likely that not all of them interact with RAGE. Although CML is recognized as a major ligand of RAGE, other AGE compounds were reported to be RAGE ligands. Indeed, CEL (N-carboxyethyl-lysine), argpyrimidine, imidazoles, pentosidine, and methylglyoxal (MGL)-derived AGEs have been reported to interact with RAGE and to activate intracellular signal transduction cascades suggesting that they might play a role in the complications of diabetes, cancer, and chronic inflammation (see Boxes 2 and 3).

In addition to AGEs, RAGE is a signal transduction receptor for amyloid-β peptide (Aβ) and β-sheets fibrils; β2-integrin Mac/CD11b; complement C3a; lipopolysaccharides (LPS); phosphatidylinerse on the surface of apoptotic cells; S100/calgranulins, a family of closely related calcium-binding proteins that accumulate extracellularly at sites of chronic inflammation; and HMGB1 (amphoterin), a DNA-binding protein released by cells undergoing necrosis that mediates inflammatory reactions and is involved in the pathophysiology of several diseases. Besides binding ligands actively participating in chronic inflammatory and immune responses, RAGE interacts with molecules on the surface of leukocytes and bacteria, and prions (Figure 1). Yet RAGE activity is not restricted to the promotion of inflammatory responses, having a role in tissue regeneration/repair as well (see Box 2).
RAGE ligands are also of endogenous origin and typically involved in the recognition of microbial-associated antigens, expressed on the cells of the innate immune system and are TLRs in immune response.37,38 In contrast to TLRs, which are receptors (TLR), pointing to cooperation between RAGE and considered as a pattern recognition receptor (PRR), like toll-like molecular patterns (DAMPs) such as HMGB1, it can be considered in case of acute inflammation.4 Most RAGE ligands (e.g. HMGB1, S100 proteins, and AGEs) are involved in acute and chronic inflammatory events, and many signalling cascades triggered by RAGE engagement determine the intracellular activation of the pro-inflammatory NF-κB (Figure 1).190 The ‘alarmin’ HMGB1 is a recognized DAMP, since it is secreted in case of inflammation, tissue injury, or infectious events, and it is released by necrotic cells, macrophages, NK cells, and dendritic cells. By interacting with RAGE, HMGB1 causes the activation of NF-κB, thereby inducing the production of several cytokines, including interleukin (IL)-6, IL-1α, and TNF-α (tumour necrosis factor). Macrophages from Agertm−/− mice produce lower amounts of pro-inflammatory cytokines after the stimulation with HMGB1, confirming the important role of RAGE in this process.191 S100–RAGE interactions might have important consequences in inflammatory and degenerative events, since they determine the activation of endothelial and vascular smooth muscle cells, neurons, astrocytes, microglia/macrophages, and other cell types, inducing the activation of different signalling pathways leading to production of pro-inflammatory adhesion molecules and cytokines. In general, S100 proteins have to be present at high levels to generate these effects, except for S100B, which activates RAGE in neurons at low and high concentrations with trophic and toxic effects, respectively,192–194 and in macrophages with anti-inflammatory and pro-inflammatory effects at low and high concentrations, respectively.9 By impinging on NF-κB, AGE–RAGE interactions also are involved in inflammatory processes.195 RAGE stimulation by AGEs increases levels of the pro-inflammatory (M1) macrophage markers, iNOS and CD86, and the pro-inflammatory cytokines, IL-6 and TNF-α.196 A positive feedback regulation of the RAGE promoter following activation of NF-κB causes increase in RAGE expression levels, thus amplifying and prolonging the inflammatory process.20 A positive feedback regulation of the RAGE promoter following activation of NF-κB causes increase in RAGE expression levels, thus amplifying and prolonging the inflammatory process.20

Since RAGE can interact with pathogen associated molecular patterns (PAMPs) such as LPS, and damage-associated molecular patterns (DAMPs) such as HMGB1, it can be considered as a pattern recognition receptor (PRR), like toll-like receptors (TLR), pointing to cooperation between RAGE and TLRs in immune response.37,34 In contrast to TLRs, which are expressed on the cells of the innate immune system and are involved in the recognition of microbial-associated antigens, RAGE ligands are also of endogenous origin and typically accumulate in tissues during ageing, inflammation, or in response to other tissue stresses39,40 (see Boxes 2 and 3).

Ligand domains interacting with RAGE display a negative charge on the surface. This acidic character of the RAGE ligands is complementary to the highly basic, electropositive surface of the RAGE ligand-binding domain. Thus, a strong electrostatic interaction drives the formation of a tight receptor-ligand complex, leading to sustained activation of the downstream signalling pathways.28,41 Binding of ligands to RAGE does not accelerate its clearance or degradation but rather begins a sustained period of cellular activation mediated by receptor-dependent signalling. Moreover, the enhanced surface expression of RAGE in environments rich in its ligands explains how upregulation of this receptor can contribute to an ascending spiral of RAGE-dependent cellular perturbation. Taken together, these features of RAGE allow the receptor to propagate cellular dysfunction in a number of pathophysiological relevant situations, most often dictated by the presence and persistence of RAGE ligands in tissues.14,42

**RAGE expression**

RAGE is constitutively or inducibly expressed depending on the cell type and the developmental stage. RAGE is extensively expressed in a constitutive manner during embryonic and early post-natal life, especially in the brain, but its expression level decreases after the accomplishment of development. This feature suggests that RAGE might play a role...
during tissue development, even if Ager−/− mice develop normally and do not show any overt phenotype and reproduce normally.45 This is probably because in Ager−/− mice, compensatory mechanisms are induced to replace RAGE activities. During the adult life, and in physiological conditions, RAGE is expressed at low levels in a wide range of differentiated cells, such as vascular endothelial cells, smooth muscle cells, almost all cells of the central nervous system, several cells of the immune system (such as monocytes, macrophages, neutrophils, dendritic cells, leukocytes, and CD4+ and CD8+ T cells), and cardiomyocytes.44 However, RAGE is upregulated under pathological conditions in which its ligands accumulate and/or transcription factors regulating RAGE are activated, such as cancer, retinal disease, atherosclerosis, cardiovascular disease, AD, respiratory disorders, liver diseases, and diabetes6,44 (see Boxes 2 and 3). Known exceptions are represented by the skin and lung, which express constitutively RAGE at high levels throughout life.39

RAGE not only localizes in the cytoplasm and to plasma membrane but also is found in mitochondria and the nucleus of specific cells, where it might have different functions from that of surface RAGE. In pancreatic tumour cells, released HMGB1 induces mitochondrial translocation of RAGE by promoting its ERK1/2 (mitogen-activated protein kinases)-dependent phosphorylation. Mitochondrial RAGE is suggested to promote the first step in the process of ATP production to sustain the increased metabolic needs and growth of tumour cells. Although localization of RAGE in mitochondria has been observed in normal mouse pancreas, lung, heart, and liver cells,45 the physiological relevance of mitochondrial RAGE is unknown. Recently, nuclear accumulation of RAGE has been shown in pathological settings that challenge the genomic integrity such as Helicobacter pylori infection,46 diabetes,47 cancer, and under severe oxidative stress that affects the nuclear integrity.48,49 RAGE, which accumulates in the nucleus after induction of DNA damage in an ATM serine/threonine kinase-dependent manner,50 has been identified as a homeostatic regulator of DNA repair, suggesting an interesting role of the receptor in the prevention of cellular senescence, inflammation, pulmonary fibrosis, and cancer.

**RAGE Signalling**

Homodimerization is a crucial event for signal transduction in many classes of receptors. Dimerization or oligomerization through self-association of V-V or C1–C1 domains is an important step for RAGE signalling after binding its ligands (Figures 1 and 2), especially if these are in an oligomeric configuration (such as S100B and AGEs).51,52 RAGE C-terminal region is not homologous to any known kinase domain, and receptor dimerization and its enhancement by ligand binding affect the association of RAGE cytosolic tail with its interacting adaptor proteins to trigger signal transduction.53 Also, upon binding by certain ligands, the phosphorylation of RAGE at Ser391 in the cytoplasmic domain by PKCζ (protein kinase C-ζ) promoted the recruitment of the adaptor protein, TIRAP (toll-interleukin 1 receptor adaptor protein) and RAGE’s downstream signalling24 (Figure 2).

Since the cytosolic domain of RAGE lacks intrinsic tyrosine kinase activity, over the past years, several studies have addressed the question how RAGE can activate signal transduction. Some cytoplasmic or transmembrane adaptor proteins have been identified whose interaction with the cytoplasmic domain of RAGE translates into the activation of specific pathways leading to sometimes opposite cellular events (such as growth, migration, invasion, apoptosis, differentiation, and/or inflammatory cytokine and chemokine secretion)55 (see Boxes 2 and 3).

Most signalling cascades activated by RAGE, such as cdc42/Rac1 (cell division control protein 42/Ras-related C3 botulinum toxin substrate 1), Src (Sarcoma family of proteins), Egr-1 (early growth response protein-1), PI3K/Akt (phosphoinositol-3-kinase/protein kinase B), and GSK3β (glycogen synthase kinase 3β), are regulated by the adaptor molecule mDia-1 (diaphanous related formin 1), a member of the GEF (guanine nucleotide exchange factor) family. Recruitment of Dia-1 by RAGE results in increased cell motility and/or chemokine secretion through activation of the Rac1/Cdc42 pathway.7,56,57 In U-87MG human glioma cells, DOCK7 (dedicator of cytokinesis 7), another member of the GEF family, is recruited to the cytoplasmic tail of RAGE to activate Rac1/Cdc42, resulting in an enhancement of cell migration.58 Moreover, RAGE cytosolic domain is connected to the tyrosine kinase protein, Src, which is related to several downstream signal factors, such as caveolin-1, ERK1/2, p38 MAPK, JNK (c-Jun N-terminal kinase), AP-1 (activating protein 1), and NF-κB (nuclear factor kappa B) and JAK/STAT3 (janus kinase/signal transducer and activator of transcription)7,57 (Figure 2).

Two additional adaptor proteins, TIRAP and MyD88 (myeloid differentiation primary response gene 88), which are involved in TLR2/4 signalling, have been shown to be connected with the phosphorylated form of RAGE. Recruitment of these two proteins translates into the activation of Akt, IRAK4 (interleukin-1 receptor-associated kinase 4), p38 MAPK, JNK, and the IKKs (IκB kinases), leading to regulation of cell survival, apoptosis, and secretion of inflammatory cytokines.7,54 Recently, DAP10 (daxx-activating protein 10) has been identified as a transmembrane adaptor for RAGE necessary to regulate cell survival and proliferation via activation of the PI3K-Akt and Ras-ERK1/2 pathways in human keratinocytes.59

In addition, RAGE interacts with novel co-receptors, FPRs (Formyl Peptide Receptors), and BLT1 (Leukotriene B4 Receptor 1), which are members of the GPCR (G-protein-coupled receptor) family. In glial cells, RAGE association
with FPRs leads to ERK1/2 activation and accelerated consumption of intracellular cAMP resulting in increased inflammatory response in the neurodegenerative AD.\textsuperscript{55,60}

The RAGE/BLT1 interaction in neutrophils results in ERK1/2-mediated reduction of NF-κB activity and increased chemotaxis, whereas in the absence of RAGE, BLT1 activates ERK1/2 and NF-κB signalling, inducing inflammatory cytokine production.\textsuperscript{61} Moreover, stimulation of RAGE in smooth muscle cells, tubular epithelial myofibroblasts, myoblasts, osteoblasts, and monocyctic cells translates into the activation of ERK1/2, whereas p38 and SAPK/JNK MAPKs are activated in monocytes/macrophages and tumour cells.\textsuperscript{6,62,63} Other pathways connected to RAGE are rho-GTPases, PI3K, and the JAK/STAT3 pathways. Activation of PI3K and ERK1/2 signalling cascades has also been observed in microglial cell lines.\textsuperscript{64,57}

**RAGE and skeletal muscle physiology**

The process of skeletal muscle formation during embryonic development, known as myogenesis, is a highly regulated process that involves the determination of multipotential mesodermal cells to myoblasts, the exit of myoblasts from the cell cycle, and their differentiation and fusion into muscle fibres.\textsuperscript{64} During myogenesis, a distinct subpopulation of myoblasts do not differentiate and remain closely associated with myofibres.
as satellite cells (SCs) in a quiescent state. In case of muscle damage consequent to trauma, intense physical exercise, chronic inflammation, cancer, or muscular dystrophies, quiescent SCs activate to repair muscle tissue. Muscle regeneration is also known as adult myogenesis, since it reproduces the events occurring during embryonic development, with the involvement of the same transcription factors as those controlling SCs proliferation and differentiation.

Myogenesis is regulated by the sequential expression of PAX3 and PAX7 (paired-domain transcription factors) that operate upstream of basic helix-loop-helix transcription factors known as MRFs (myogenic regulatory factors). PAX7 is expressed in and marks quiescent SCs and is essential for the proliferation and survival of myogenic progenitors, preventing their precocious differentiation. The MRFs (namely, Myf5, MyoD, myogenin, and MRF4) are timely activated in myoblasts to regulate downstream targets such as MyHC (myosin heavy chain), MCK (muscle creatine kinase), and troponin T, necessary for terminal differentiation and generation of functional myofibres. PAX7 becomes repressed in differentiating myoblasts/myocytes, which are characterized by the expression of the terminal differentiation marker myogenin. Indeed, the expression of PAX7 and myogenin are mutually exclusive in myoblasts, and downregulation of PAX7 is required for myoblast terminal differentiation to occur. In myoblasts committed to differentiation, PAX7 expression is downregulated post-transcriptionally by microRNA-1, -206, and -486, and post-translationally by MyoD and myogenin. PAX7 is also essential for SC self-renewal. In fact, during muscle regeneration, some myoblasts reacquire a quiescent state through asymmetric division, thus reconstituting the SC reserve pool. Many steps of myogenesis can be recapitulated in vitro by culturing primary myogenic cells or myoblast cell lines in low serum conditions, in which myoblasts can form multinucleated myotubes. An essential feature of skeletal muscle formation is selective apoptosis that eliminates differentiation-incompetent myoblasts during myogenesis.

In skeletal muscle tissue, RAGE expression is developmentally regulated. RAGE can be detected in immature, nearly mature, and some mature myofibres up to 11 days after birth in rodents, with RAGE expression being restricted to the sarcolemma. However, RAGE is absent in adult muscle tissue. The presence of both positive and negative muscle fibres in 11-day-old rats suggests that repression of RAGE expression occurs around this time. This pattern of expression is typical of RAGE, which is expressed during development in several cell types and is repressed in their adult counterparts, suggesting that RAGE might play a role during muscle development. Accordingly, proliferating and differentiating myoblasts express nearly constant levels of RAGE mRNA, whereas protein levels significantly decline during late phases of differentiation in vitro. RAGE engagement by serum factors, such as HMGB1, promotes and sustains myoblast differentiation and myotube formation. The density of RAGE molecules on their surface and the relative concentration of HMGB1 and, likely, other RAGE ligands appear to determine myoblasts’ final fate. HMGB1–RAGE interaction in myoblasts results in activation of p38 MAPK via Cdc42–Rac1–MKK6, triggering myogenin and MyHC expression and MCK induction. Besides promoting differentiation, RAGE engagement by HMGB1 in myoblasts induces cell proliferation arrest through p38 MAPK-mediated inhibition of the Raf-MEK-ERK1/2 pathway with ensuing downregulation of cyclin D1 expression and Rb (retinoblastoma suppressor protein) phosphorylation and upregulation of the proliferation inhibitor, p21WAF1, and JNK inactivation. Moreover, HMGB1–RAGE interaction in myoblasts promotes apoptosis via reduction of the anti-apoptotic factor Bcl-2 expression (Figure 3).

RAGE signalling also modifies myoblast morphology and motility. Functional inactivation of RAGE in myoblasts results in decreased cell size along with a less organized cytoskeletal F-actin and increased filopodia formation, hallmarks of proliferating, migrating, and poorly adherent cells. On the other hand, fl-RAGE-expressing myoblasts exhibit high levels of adhesion molecules, such as β1-integrin, NCAM, VCAM, and caveolin-3, which play an important role in myoblast adhesiveness and differentiation. Thus, RAGE activity contributes to the increased adhesion properties and reduced motility of differentiating myoblasts. Further, RAGE activation and signalling in myoblasts result in reduced activity of matrix metalloproteinases (MMPs) 1 and 2, important for myoblast migration and invasiveness during early phases of embryonic skeletal muscle formation, when proliferating myoblasts have to reach the place of muscle formation, and during muscle regeneration, when activated SCs migrate towards damaged myofibres.

However, different RAGE ligands elicit different effects by interacting with the receptor on the myoblast surface. Indeed, S100B can bind RAGE without activating it and block the RAGE promyogenic signalling by preventing its interaction with HMGB1. S100B is able to differentially activate RAGE or bFGF (basic fibroblast growth factor)/FGFR1 (fibroblast growth factor receptor 1) depending on its own concentration, the presence or absence of bFGF, and myoblast density. In low-density myoblast cultures, which better reflect the beginning of the regeneration process in vivo, low S100B engages RAGE, thereby simultaneously stimulating cell proliferation via MEK-ERK1/2 and activating the myogenic programme via p38 MAPK. S100B oligomers present in the extracellular space bind to RAGE V domain and promote RAGE oligomerization and stabilization, resulting in its activation. At high S100B concentrations, higher-order S100B oligomers might additionally interact with FGFR1-bound bFGF resulting in the formation of a
RAGE/S100B/bFGF/FGFR1 complex on the same cell. Here, RAGE cannot efficiently oligomerize and signal, while bFGF- FGFR1 signalling is enhanced. However, in the absence of bFGF, either low or high S100B activates RAGE in low-density myoblast cultures, resulting in the simultaneous stimulation of proliferation and the myogenic programme. Similar to RAGE, other cell surface receptors belonging to the Ig superfamily such as NCAM (neural cell adhesion molecule), CDO (cell adhesion associated oncogene), and BOC (brother of CDO) are expressed during embryonal skeletal muscle development and sustain myoblast differentiation.

Although RAGE is not expressed in healthy adult skeletal muscles, activated/proliferating SCs and differentiating myoblasts re-express RAGE upon muscle injury, and RAGE expression is repressed at the completion of the regeneration process. ROS produced following muscle damage induce NF-κB (p65)-dependent expression of RAGE in activated SCs. Further accumulation of RAGE in differentiating myoblasts is dependent on myogenin. During muscle regeneration, RAGE activity is driven by the presence of the RAGE ligands, S100B and HMGB1, which are released with different kinetics upon injury. S100B released by skeletal muscles early after...
injury stimulates RAGE signalling and enhances RAGE expression in activated SCs, inducing myoblast proliferation and activating the myogenic programme. However, during the proliferation phase, S100B and bFGF are present at the same time in the extracellular space, and S100B promotes myoblast expansion by enhancing bFGF/FGFR1 signalling. At this time, RAGE is recruited into a RAGE/S100B/bFGF/FGFR1 tetradecorplex that impairs RAGE signalling. HMGB1 is passively released from muscle tissue following damage and progressively accumulates during the regeneration process. By binding RAGE on the myoblast surface, HMGB1 activates p38 MAPK, thus upregulating the expression of myogenin, which in turn upregulates RAGE expression. This positive feedback loop leads to myogenin accumulation culminating in the repression of PAX7 expression through binding of myogenin to specific consensus sequences in the PAX7 gene promoter. This mechanism strongly contributes to the mutually exclusive presence of PAX7 and myogenin in muscle precursor cells. Besides translating into reduced myoblast proliferation rate, the downregulation of PAX7 subsequent to HMGB1–RAGE interaction promotes symmetric division in activated SCs, thereby limiting SC self-renewal (Figure 4). Therefore, RAGE might accelerate myogenic differentiation and the regeneration process and optimize myofibre size and the SC number. Consistent with these results, acutely injured skeletal muscles of Ager−/− mice are characterized by excess myoblast proliferation and elevated myoblast asymmetric division leading to delayed regeneration. Also, during early regeneration phases, the number of infiltrating macrophages in Ager−/− injured muscles is smaller than

**Figure 4** Role of RAGE in acute or chronic muscle injury. Upon acute skeletal muscle injury, re-expressed RAGE in activated satellite cells results in their proliferation and differentiation ultimately leading to regeneration. Under the action of S100B, RAGE signalling promotes proliferation and simultaneously activates the myogenic programme. RAGE activated by HMGB1 causes proliferation arrest and stimulates terminal differentiation of myoblasts into myocytes that build up new myofibres and/or repair damaged myofibres. RAGE-mediated signalling in infiltrating macrophages modulates the inflammatory response in injured muscles, contributing significantly to accelerate regeneration. S100B/RAGE axis also promotes macrophage infiltration of acutely damaged skeletal muscle and macrophage polarization to M2 phenotype (left). On the contrary, in the course of muscle diseases characterized by chronic inflammation, high glucose conditions, or oxidative stress, an accumulation of RAGE ligands occurs in the serum and damaged myofibres. The overexpressed and overstimulated RAGE amplifies the inflammatory response by stimulating proinflammatory cytokines secretion, induces apoptosis of myocytes, and causes lethal accumulation of oxidative stress contributing to chronic myofibre damage (right).
that of controls, in accordance with the role of RAGE in macrophage migration,26 thus conditioning the timing of muscle regeneration78 (Figure 4).

Infiltrating M1 (classically activated) macrophages contribute significantly to muscle regeneration by removing cell debris and providing important factors for the activation, migration, and proliferation of SCs.87 The transition of macrophages from a pro-inflammatory (M1) to an anti-inflammatory (M2) phenotype is crucial for efficient tissue repair,88 since M2 (alternatively activated) macrophages dampen the inflammatory response and promote muscle regeneration.29,88 RAGE-mediated signalling in infiltrating macrophages modulates the inflammatory response in injured muscles, contributing significantly to accelerate regeneration. Indeed, absence of RAGE results in a delayed and amplified inflammatory response9,78 (Figure 4).

As mentioned earlier, complement C3a interacts with RAGE,29 and complement C3a has been recently shown to promote muscle regeneration following acute injury by activation of the so-called alternative complement’s pathway required for initiating recruitment of circulating monocytes into injured muscle.89 It can be hypothesized that RAGE might participate in the muscle regeneration process by transducing complement C3a effects. Besides, the S100B/RAGE axis promotes macrophage infiltration of acutely damaged skeletal muscle and macrophage polarization to M2 (anti-inflammatory and pro-regenerative) phenotype.9

Thus, in acutely injured muscles, RAGE affects muscle regeneration by a dual mechanism, promoting macrophage infiltration and polarization to M2 phenotype and regulating SC/myoblast activity, thus allowing proper myoblast proliferation and differentiation phases (Figure 4). However, excess RAGE signalling consequent to accumulation of S100B (e.g. in muscular dystrophy) results in excess macrophage infiltration and prolongation of the inflammatory phase, ultimately leading to disturbed regeneration9 (Figure 4).

HMGB1 levels increased in muscles after ischaemia induced by femoral artery ligation, and blockade of endogenous HMGB1 by administration of its truncated form, BoxA, resulted in reduction of vessel density and delayed revascularization of the so-called alternative complement pathway reparation,88 since M2 (alternatively activated) macrophages dampen the inflammatory response and promote muscle regeneration. RAGE-mediated signalling in infiltrating macrophages modulates the inflammatory response in injured muscles, contributing significantly to accelerate regeneration. Indeed, absence of RAGE results in a delayed and amplified inflammatory response9,78 (Figure 4).

As mentioned earlier, complement C3a interacts with RAGE,29 and complement C3a has been recently shown to promote muscle regeneration following acute injury by activation of the so-called alternative complement’s pathway required for initiating recruitment of circulating monocytes into injured muscle.89 It can be hypothesized that RAGE might participate in the muscle regeneration process by transducing complement C3a effects. Besides, the S100B/RAGE axis promotes macrophage infiltration of acutely damaged skeletal muscle and macrophage polarization to M2 (anti-inflammatory and pro-regenerative) phenotype.9

Thus, in acutely injured muscles, RAGE affects muscle regeneration by a dual mechanism, promoting macrophage infiltration and polarization to M2 phenotype and regulating SC/myoblast activity, thus allowing proper myoblast proliferation and differentiation phases (Figure 4). However, excess RAGE signalling consequent to accumulation of S100B (e.g. in muscular dystrophy) results in excess macrophage infiltration and prolongation of the inflammatory phase, ultimately leading to disturbed regeneration9 (Figure 4).

HMGB1 levels increased in muscles after ischaemia induced by femoral artery ligation, and blockade of endogenous HMGB1 by administration of its truncated form, BoxA, resulted in reduction of vessel density and delayed regeneration, the opposite being observed with intramuscular administration of HMGB1.90 The ability of HMGB1 to sustain skeletal muscle regeneration was attributed to a direct effect on myoblasts, enhancing their recruitment to the site of injury, as well as to the arteriogenic action of HMGB1. Although myoblasts and endothelial cells were found to express the HMGB1 receptors, RAGE and TLR4, the authors did not establish which receptor mediates the HMGB1’s effects in these conditions. However, others reported that, despite HMGB1–RAGE interaction is important for the recovery of the skeletal muscle following acute injury consequent to hind limb ischaemia, RAGE was proposed not to play a significant role in muscle regeneration.71 In these conditions, neutralization of HMGB1 resulted in prolonged muscle necrosis, suggesting that HMGB1 released in the setting of ischaemia is critical for promoting muscle recovery. However, while TLR4 KO mice showed impaired regeneration, RAGE KO mice exhibited muscle histology that was indistinguishable from control animals, suggesting that HMGB1 effects are mediated by TLR4 rather than RAGE in these experimental conditions.91 It is possible that strain differences of RAGE and TRL4 KO mice affect muscle regeneration after ischaemia. Another possibility stands on the fact that HMGB1 can exist into two redox forms, an oxidized one and a reduced one, that are able to differentially activate RAGE and TLR4. The HMGB1’s ability to activate RAGE is restricted to the oxidized form of the protein.92,93 Further studies are required to determine the exact role of RAGE and TLR4 in the HMGB1’s effects in muscle tissue after ischaemic injury.

**RAGE and skeletal muscle atrophy**

Skeletal muscle wasting (atrophy) is a complex and highly regulated process, characterized by a substantial decrease in muscle mass, myofibre area, protein content, regenerative capacity, and muscle strength, together with increased myocyte apoptosis and fatigability.4,94 In addition to cancer, muscle wasting is associated with ageing, disuse, diabetes, denervation, and many chronic diseases characterized by a variable degree of systemic chronic inflammation, in which muscle atrophy contributes to morbidity and mortality.3 In healthy young individuals, skeletal muscle maintains mass and function through the balance of intricate signalling networks that regulate muscle protein synthesis (hypertrophy) and degradation (atrophy),95,96 leading to physiological levels of muscle proteins. In muscle atrophy, this balance shifts towards a catabolic state resulting in the breakdown of myofibrillar proteins due to activation of the two major intracellular proteolytic systems (i.e. ubiquitin-proteasome [UPP] and autophagy) and decrease in protein synthesis.3,95,96 Myogenic precursors, including SCs, also might contribute to the maintenance of skeletal mass in health and diseases,97 and recent work has shown that atrophy factors inhibit their regenerative and fusion potential.98,99

Systemic atrophy factors affect muscle mass by interfering with catabolic and anabolic pathways. In particular, inflammatory cytokines deactivate the PI3K/AKT/mTOR (mammalian target of rapamycin) pathway, leading to the reduction of protein synthesis and activation of the FOXO3 (forkhead box O3) transcription factor. FOXO3 is sufficient to induce the expression of the muscle-specific ubiquitin ligases atrogin-1/MAFbx (muscle atrophy F-box protein) and MuRF-1 (muscle RING finger-1), collectively called atrogenes, and at the same time autophagy-associated proteins (Atgs), leading to selective proteolysis of MyHC in skeletal muscle.100–102 On the other side, inflammatory cytokines activate NF-κB and p38

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MAPK, resulting in a direct induction of MuRF-1 and atrogin-1, respectively. Many other pathways such as AMPK (AMP-activated kinase), ERK, and JNK are involved in the activation of UPP system by regulation of FOXO3 activity and atrogene expression.

**RAGE in diabetes-induced atrophy**

Diabetic myopathy is characterized by impaired muscle growth and development, loss of muscle mass, a switch to a glycolytic myofibre phenotype, muscle weakness, and a reduced physical functional capacity that influence the rate of co-morbidity development. Also, the ability of SCs to repair damaged myofibres is adversely affected by diabetes environment probably due to prolonged exposure to high glucose.

Early evidences showed that CML accumulates in myofibres of the plantaris muscle of diabetic rats and that the sites where CML accumulates are associated with fibre-type transformation, which in turn is associated with muscular endurance. Increased blood levels of different AGEs, including CML, and increased levels of unspecified muscle AGEs and RAGE were found in diabetic patients and mice. Recently, the role of AGEs and RAGE as molecular players in diabetic myopathy has been examined in some depth using different approaches, that is, diabetic mice, cultured muscle cells, and patient samples. In a cell culture model mimicking diabetic conditions, high glucose levels promote the production and accumulation of AGE–BSA, which in turn are responsible for the membrane repair defect. In particular, AGE–BSA, even in the absence of high glucose, might cause repair defects by RAGE engagement, resulting in cell death and loss of muscle mass. Consistently, RAGE-deficient cells are resistant to glucose-induced repair defects.

It is intriguing that the alagebrum chloride, an AGE crosslink breaker with chelating proprieties, used as a therapeutic compound in diabetic complications in various experimental animal models, is a potential therapeutic candidate to improve diabetic myopathy, including muscle wasting, muscle atrophy, and muscle regenerative capacity, by reducing plasma and muscle AGE levels in diabetic mice. Accordingly, AGE–BSA induced atrophy in myotube cultures and strongly inhibited myoblast differentiation through a RAGE-mediated mechanism. In particular, AGE–BSA activated AMPK, a negative mediator of muscle mass, and downregulated Akt signalling, leading to activation of the atrophy marker Atrogin-1 and reduction of myotube size. Moreover, AGE–BSA treatment showed detrimental effects in myoblast differentiation and fusion by reducing myogenin and MHC expression via phosphorylation of AMPK and dephosphorylation of Akt. The detailed mechanism acting in the negative regulation of muscle SC/stem cell homeostasis by AGE/RAGE in vivo remains to be elucidated.

A decrease in insulin action on target tissues, defined as insulin resistance (IR), is characteristic of type 2 diabetes. Abnormal accumulation of MGL, a highly reactive AGE precursor, occurs in diabetic patients, and MGL-derived AGEs were shown to activate RAGE signalling. When MGL was reduced in type 2 diabetic patients, IR was improved. Recently, an interesting study proposed a novel MGL-derived AGE inhibitor, MK-I81, as a potential therapeutic compound to alleviate AGE-mediated downregulation of insulin signal transduction and insulin action in skeletal muscle cells restoring insulin sensitivity.

In addition to those formed endogenously, AGEs are abundant in exogenous sources such as food prepared under elevated temperatures. Dietary AGEs have been shown to induce IR in mice also in absence of hyperglycaemia. Accordingly, reduced intake of AGEs has been shown to decrease the incidence of type I diabetes in non-obese diabetic mice and improved insulin sensitivity in db/db mice. Also, similar to exposure to glycated albumin, chronic exposure of L6 myoblasts to dietary AGEs induces the formation of a complex including RAGE-PKCα-Src that leads to phosphorylation of IRS (insulin receptor substrate) and inhibition of insulin action. Lastly, muscle tissue from weight gainers and elderly subjects show a marked increase in immunostaining intensity for CML and RAGE in association with immunostaining of markers of oxidative stress and inflammation. Future studies would hopefully provide a complete picture of the role of RAGE and its ligands in the pathophysiology of skeletal muscle disorders in diabetes (Figure 4).

**RAGE in sarcopenia**

Sarcopenia defines an age-related, continuous decline in muscle mass, quality, and strength. Sarcopenia is characterized by an overall decrease in size and number of skeletal muscle fibres, mostly the type 2 or fast-twitch muscle fibres, and a marked infiltration of skeletal muscle with fibrous and adipose tissue. Many factors contribute to sarcopenia, including changes in skeletal muscle metabolism, hormonal changes, compromised regeneration, elevated levels of inflammatory cytokines, changes in vasculature, inactivity, and oxidative stress. Besides the activation of autophagy/lysosome systems leading to protein degradation, there is consensus regarding an age-related decline in the performance of SCs as a primary cause of sarcopenia. In this regard, both extrinsic factors of the extracellular environment (the so-called SC 'niche') and intrinsic properties of the SCs have been proposed to contribute to sarcopenia. Interestingly, early work showed that compared with their young counterparts, SCs isolated from aged (>72 years old) human subjects (i) exhibit higher intracellular levels of the RAGE ligand, S100B, known to inhibit myoblast...
differentiation via NF-κB-dependent repression of MyoD (and myogenin) expression; (ii) release lower amounts of mitogenic factors such as S100B and bFGF; and (iii) express a truncated form of RAGE in plasma membrane. The altered levels of S100B and RAGE correlated positively with the defective proliferation and differentiation potential exhibited by aged SCs, a notion reinforced by the observation that either transient transfection with fl-RAGE or knockdown of S100B switched aged SCs to a young phenotype and conversely, functional inactivation of RAGE or transient transfection with S100B switched young SCs to an aged phenotype. Lastly, treatment of aged SCs with S100B, bFGF, or young SC-conditioned medium enhanced aged SC proliferation and their subsequent differentiation. Also interestingly, chronic oxidative conditions lead to accumulation of S100B in an NF-κB-dependent manner in myoblasts and S100B-stimulated conversion of myoblasts into brown adipocytes via an NF-κB/YY1/miR-133 axis and NF-κB/YY1/BMP-7 axis, events prevented by S100B knockdown. In this same experimental setting, interstitial cells and, unexpectedly, a subpopulation of myofibres in muscles of geriatric but not young mice were found to co-express S100B and the brown adipocyte marker, uncoupling protein-1. However, RAGE expression and activity were not studied in this experimental setting, thus whether RAGE has a role in S100B-induced myoblast-brown adipocyte conversion and/or RAGE transduces effects of chronic oxidative stress in myofibres remain elusive.

AGE-modified proteins are known to become resistant to degradation by the UPP machinery, and the accumulation of modified proteins in several tissues, including skeletal muscle, has been considered as one of several potential mechanisms producing age-related functional decline of diverse organs. Several evidences point to a correlation between the decline of muscular function in healthy aged subjects and circulating or muscle CML, CEL, and pentosidine. In this respect, the AGE product, pentosidine, is increased by more than 200% in skeletal muscle of older adults compared with younger adults, suggesting that glycate-related cross-linking of intramuscular connective tissue might cause the reduction of viscoelastic properties of muscle, thus contributing to the decline in muscle function with ageing. CML may also play a role in sarcopenia through upregulation of inflammation and endothelial dysfunction in the microcirculation of skeletal muscle via RAGE by inducing the loss of myocytes and loss of muscle mass and strength (see Figure 4).

RAGE and skeletal muscle diseases

Myopathies are a large group of clinical disorder of skeletal muscles characterized by progressive weakness due to abnormalities of myofibre structure and metabolism, which ultimately leads to severe disability of the patient, pain, and death. Based on aetiology, myopathies can be divided into two main categories: inherited (i.e. muscular dystrophies, congenital myopathies, mitochondrial myopathies, and metabolic myopathies) and acquired (i.e. myositis or idopathic inflammatory myopathies, toxic myopathies, and myopathies associated with systemic conditions). RAGE in inherited myopathies

Muscular dystrophies (MDs) are a group of disorders that share clinical and pathological features, characterized by progressive muscle degeneration, affecting principally limb, axial, and facial muscles. There are different forms of MDs depending on the severity, age of onset, rate of progression, consequent implications, and prognosis. The actual classification of MDs into seven major groups is based on the links between protein mutations, their localization and functions, and the final clinical phenotype and include Duchenne muscular dystrophy (DMD), limb girdle muscular dystrophies (LGMDs), and facioscapulohumeral muscular dystrophy (FSHD). RAGE is detectable and co-localizes with CML and NF-κB in regenerating myofibres and less frequently in degenerating or necrotic myofibres of patients with LGMDs, even if with a lower staining intensity than in inflammatory myopathies. Moreover, RAGE is overexpressed in FSHD muscles when compared with normal and DMD muscles likely consequent to increased production of AGEs due to elevated ROS levels. AGE/RAGE interaction is known to increase ROS production via NF-κB activation, thus perpetuating the oxidative cellular damage involved in the FSHD pathogenesis.

RAGE in acquired myopathies

Myositis is a heterogeneous group of disorders that are characterized clinically by chronic muscle weakness, fatigue, and damaged myofibres due to infiltration of inflammatory cell in muscle tissue. Based on differences in clinical and histopathological features, myositis can be subdivided into polymyositis (PM) and dermatomyositis (DM), which are chronic autoimmune diseases, and inclusion body myositis (IBM), the most common degenerative muscle disease in adults over the age of 50. In PM and DM, an overexpression of RAGE and its ligands, HMGBl and CML, has been demonstrated in immune cells as well as in regenerating muscle fibres. Treatment with glucocorticoids, conventionally used in patients with PM and DM, was associated with a reduced expression of HMGBl,
sustaining muscle in nutrient homeostasis. Interestingly, HMGB1 and RAGE levels increase in skeletal muscle of patients with IBM, and the HMGB1–RAGE axis induces intracellular protein accumulation in IBM as well as cell death. Thus, similar to AD where the β-amyloid–RAGE interaction is crucial for the cellular damage and cognitive neuronal dysfunction, RAGE may be a relevant mediator in the degenerative mechanism of muscle in IBM. Furthermore, there is evidence of significantly elevated serum levels of pentosidine and CML and accumulation of CML in the muscle tissue of patients with fibromyalgia (FM), a frequent rheumatic disorder of unknown aetiology and pathogenesis. In this same study, a more intensive staining for RAGE, NF-κB, and CD68-positive monocytes/macrophages was found in the interstitial connective tissue of muscle of FM patients suggesting that AGEs’ binding to RAGE might prolong NF-κB activity leading to the secretion of pro-inflammatory cytokines, which in turn might contribute to pain generation and perpetuation of inflammation.

Myasthenia gravis (MG) is an autoimmune disease caused by the production of antibodies against receptors at the neuromuscular junction such as nicotinic acetylcholine receptor and is clinically characterized by weakness of facial, ocular, and limb muscles. An analysis of a broad inflammatory circulating protein and cytokine profile in patients affected by different MG subtypes revealed the presence of EN-RAGE (also known as S100A12 protein) in the sera of MG patients. Besides being a potential biomarker in MG, EN-RAGE/S100A12 might contribute to MG pathogenesis by binding to RAGE to activate NF-κB and induce secretion of pro-inflammatory cytokines, a property shared with several other S100 proteins.

**RAGE and skeletal muscle metabolism**

Skeletal muscle can be considered as the largest metabolic organ in the human body, and metabolic dysfunctions occurring in skeletal muscle in diabetes, obesity, sarcopenia, cancer cachexia, and muscular dystrophies are characterized by a ‘metabolic reprogramming’ and can affect the whole-body nutrient homeostasis. Recently, various authors have advanced the idea that the presence of tumour promotes metabolic reprogramming of the skeletal muscle tissue in order to counter the high energy demand of tumour tissue and sustain proliferation and cell-death resistance of tumour cells. In this scenario, the dramatic muscle wasting occurring in cancer, known as cancer cachexia, may be considered as a part of the metabolic reprogramming in cancer patients. A metabolic crosstalk has been identified in colorectal cancers between tumour masses and skeletal muscle. During the development and progression of colorectal cancer, expression of the autophagy-inducing stress protein HMGB1 increased in the muscle of tumour-bearing animals. This effect was associated with decreased expression and activity of the pyruvate kinase PKM1, which mediates the linker reaction between glycolysis and the TCA cycle in normal tissue muscle via RAGE. Colon cancer cells release HMGB1 in the serum, which decreases mTOR levels in muscle through RAGE-dependent p38 phosphorylation, thus increasing autophagy-associated proteins (beclin-1 and LC3II). This results in increased plasma glutamate and free amino acids that supply energy to cancer cells.

**RAGE and rhabdomyosarcoma**

Rhabdomyosarcomas (RMSs) are highly malignant tumours of skeletal muscle origin accounting for more than 50% of soft tissue sarcomas in childhood. It is widely accepted that RMSs represent aberrant stages of normal muscle development: like normal muscle precursor cells, RMS cells express PAX3, PAX7, and MRFs such as myogenin, all of which are hallmarks of RMS. Alveolar RMSs (ARMSs) and embryonal RMSs (ERMSs) are the most frequent variants, with ERMSs accounting for 60–70% of RMSs. ERMSs show the phenotypic and biological features of embryonic muscles, suggesting that they may arise from muscle progenitor cells, such as activated SCs or poorly differentiated myoblasts. The tumour properties of ARMSs and ERMSs have been associated to PAX genes because of the transcriptional hyperactivity of PAX3/FOX01A (forkhead box protein O1) or PAX7/FOX01A fusion transcripts in ARMSs and the high level of PAX7 expression in ERMSs. As mentioned earlier, the persistence of PAX7 in myoblasts results in prolonged proliferation and defective differentiation, implying that conditions leading to accumulation of PAX7 in myoblasts can potentially promote ERMS formation. Moreover, PAX activity regulates the expression of cell surface molecules involved in cell adhesion, migration, and invasiveness, particularly HGFIR (hepatocyte growth factor receptor), also called Met, ephrins (EFNs) and Eph receptors (EPHs), and NCAM in various cell types. Studies of the role of RAGE in RMS have shown that high PAX7 levels in ERMS cells are partly dependent on reduced and/or absent RAGE signalling and are major cause of their uncontrolled proliferation and the acquisition of a metastatic behaviour. As mentioned earlier, HMGB1–RAGE interaction on myoblasts results in enhanced myogenic differentiation with simultaneous cell proliferation arrest, repression of PAX7 transcription, and enhanced PAX7 degradation by a proteasomal-dependent mechanism and apoptosis induction. Indeed, an inverse relationship exists between RAGE and myogenin protein levels and those of PAX7 in
muscle precursor cells\textsuperscript{170} (Figures 3 and 5). Interestingly, inoculation of myoblasts stably overexpressing RAGE\textsubscript{Δcyto} into immunocompromised mice results in much larger and more aggressive tumour formation compared with inoculation of myoblasts stably overexpressing fl-RAGE.\textsuperscript{8} Tumour masses derived from RAGE-defective myoblasts appeared earlier and were characterized by large central areas of necrosis and peripheral neovascularization compared to masses formed by myoblasts harbouring fl-RAGE. Injected cells invaded the neighbouring skeletal muscle tissue in the case of RAGE\textsubscript{Δcyto} myoblasts only, and the masses obtained in this case showed hallmarks of ERMSs.\textsuperscript{8}

Studies of several RMS cell lines showed that the ability of RMS cells to undergo myogenic differentiation appears to be dependent on the amount of functionally active RAGE.\textsuperscript{171} Indeed, TE671 cells stably transfected with RAGE showed Figure 5: Schematic representation of the effects of enforced expression of RAGE in TE671 ERMS cells. RAGE engaged by HMGB1 stimulates myogenic differentiation through activation of MKK6/p38 MAPK and myotube hypertrophy through PI3-K/Akt. HMGB1/RAGE-dependent activation of p38 MAPK also causes inactivation of ERK1/2 and JNK with consequent inhibition of proliferation and decrease in cell survival. Induction of RAGE in ERMS cells also results in the expression of myogenin and MyoD so as to cause repression of PAX7 expression and PAX7 proteasomal degradation, ultimately leading to reduction of proliferation, enhancement of apoptosis, and myogenic differentiation. Finally, the reduction of Pax7 expression by the RAGE/myogenin axis causes a reduction of cell migration and invasiveness in ERMS.
RAGE inhibitors

The involvement of RAGE in the pathophysiology of several diseased conditions has prompted studies aimed to identify endogenous inhibitors and synthesize chemical inhibitors. Indeed, a variety of RAGE antagonists are now available for preclinical and clinical studies.\(^{176}\)

To date, several small-molecule inhibitors have been developed to contrast RAGE activity in several diseases that bind either the extracellular or intracellular domain. Among the extracellular inhibitors of RAGE, TTP488 ([3-(4-{2-butyl-1-[4-(4-chlorophenoxy)-phenyl]-1H-imidazole-4-yl}-phenoxy)-propyl]-diethylamine 1), also known as azeliragon or PF-04494700, is an orally bioavailable small molecule able to cross the blood–brain barrier (BBB).\(^{177}\) TTP488 is able to inhibit the interaction between RAGE and Aβ, S100B, HMGBl, and AGEs and preclinical studies have demonstrated that TTP488 efficiently contrasts complications due to RAGE activity in diabetes, cardiovascular disease, cancer, and AD.\(^{52,178}\)

On the basis of the results obtained in a mouse model of AD, in which TTP488 administration inhibited inflammation, neuronal Aβ accumulation, and neurocognitive decline, a phase III clinical trial of TTP488 is currently in progress in an AD cohort of 800 human patients.\(^{179}\) Several derivatives of TTP488, set up to reduce the toxic proprieties of this inhibitor when used at elevated doses, have been developed and demonstrated able to inhibit the disease in mouse model of AD.\(^{180,181}\) However, none of these derivatives have progressed to human clinical trial yet.

Similarly, the compound FPS-ZM1 (N-benzyl-N-cyclohexyl-4-chlorobenamide 12) was selected through the screening of 5000 compounds in the search for inhibitors of RAGE/Aβ interaction. FPS-ZM1 is able to cross the BBB and blocks inflammatory signalling in the mouse brain, reduces Aβ accumulation, and improves cognitive performance without causing toxic side effects in mice, even at high doses.\(^{182}\) The efficacy of FPS-ZM1 was explored in other animal models of neuropathology (such as hippocampal inflammation and brain stroke), heart failure, asthma, and cancer progression/metastasis of lung and liver tumours.\(^{183,184}\) The results demonstrated that the administration of FPS-ZM1 reduces or abrogates the pathogenic effects caused by RAGE activity by reducing AGE–RAGE interactions. Further, small-molecule inhibitors have been identified, which interfere with the RAGE–mDia-1 interaction. In vitro and in vivo RAGE–Dia-1 competitive inhibitors displayed inhibitory effects on AGE–RAGE signal transduction by reducing inflammation, vascular cell migration, and pro-inflammatory cytokines.\(^{185}\) Although these results are promising, further studies are needed to establish chemical refinement, tolerance, selectivity, and pharmacokinetics of these compounds and to test whether intracellular and extracellular inhibitors are effective therapeutics in several diseases.
Conclusions

Increasing evidence suggests that RAGE signalling plays an important role in skeletal muscle during the development, dictating the fate of myogenic precursor cells. Once activated by S100B or HMGB1, RAGE regulates the fine balance between apoptosis, proliferation, and differentiation in myoblasts so that absent or reduced RAGE signalling in activated muscle SCs might concur to RMS genesis. Accordingly, restoring RAGE signalling can be a potential therapeutic tool in ERMSs by promoting terminal differentiation, reducing proliferation, and repressing the expression of molecules responsible for cell migration and invasiveness. Also, the timely expression of RAGE during skeletal muscle development and regeneration, together with its absence in healthy adult muscle tissue, highlights the role of this receptor in building muscle mass. The fact that Ager−/− mice do not show overt muscle defects and do not spontaneously develop RMS suggests that other receptors belonging to the same superfamily (e.g. NCAM, CDO, and BOC) might compensate for the lack of RAGE activity.

Several studies demonstrated that RAGE is re-expressed in muscle under acute or chronic inflammatory conditions, and the local environment appears to determine the outcome of RAGE signalling. In acutely injured myofibres, re-expressed RAGE supports the restoration of tissue homeostasis and concurs to tissue repair by promoting myoblast proliferation and differentiation and the resolution of inflammation. On the contrary, in the course of muscle diseases characterized by chronic inflammation, RAGE signalling appears to concur to the pathogenesis by amplifying and perpetuating the inflammatory response, inducing apoptosis of myocytes, and causing lethal accumulation of oxidative stress leading to myofibre damage. Thus, RAGE is likely to represent one main factor intervening in the fine regulation of the balance between injury and repair: in acute muscle injury, short-term RAGE signalling triggered by relatively low concentrations of RAGE ligands results in beneficial effects, whereas in chronic muscle injury conditions, long-term RAGE signalling triggered by high concentrations of RAGE ligands is deleterious. Although additional studies are required to better elucidate the role of RAGE and its ligands in myopathies, blockade of ligand–RAGE interaction results in reduction of inflammation and ROS-induced myofibre damage, thus representing a potential therapeutic strategy.

Although several studies suggest that AGE–RAGE is able to activate well-known signalling pathways and transcriptional factors inducing muscle atrophy, membrane damage, and insulin resistance in diabetes, these findings suffer from some limitations. The techniques used in in vivo studies are not able to differentiate the type of AGEs capable of recruiting RAGE, and supraphysiological doses of albumin were used in in vitro studies to provide a heterogeneous level of AGEs. Additional studies are needed to discern the true physiological relevance of such findings in the context of RAGE activation in diabetes.

Interestingly, CML–RAGE interactions have been reported to stimulate ROS formation determining systemic oxidative stress and inflammation and loss of muscle mass in obesity and during ageing. However, the causal relationship between high levels of specific AGEs and sarcopenia/obesity is still elusive. The data obtained so far suggest that RAGE is involved in the pathogenesis of these two important medical problems through common mechanisms that await elucidation, though. Moreover, the opposite behaviour of RAGE in skeletal muscle might be explained by the presence and expression levels of different types of RAGE-binding proteins in physiological and pathological conditions. We can hypothesize that the different effects of RAGE in muscle precursor cells in acute injury versus chronic inflammation/elevated ROS conditions might be explained by a differential availability of RAGE adaptor proteins and/or physical/functional interactions between RAGE and receptors shown to share with RAGE certain ligands. The discovery of these binding partners in skeletal muscle might hopefully elucidate the role of RAGE in maintaining or altering muscle homeostasis. The behaviour of RAGE in skeletal muscle may be also affected by the state of RAGE phosphorylation, which has been demonstrated to dictate both the downstream signalling of RAGE and the translocation of the receptor to the nucleus or mitochondria, where RAGE might exert yet unidentified function(s) in skeletal muscle.

A few studies have shown that RAGE signalling contributes to the remodelling of skeletal muscle metabolism by reducing protein reserve in chronic stress conditions, such as cancer. Considering that the majority of metabolic disorders induce muscle degenerative diseases and that high serum levels of RAGE ligands correlate positively with cancer severity in a variety of tumour types, it will be important to explore RAGE’s activity in tumour-induced metabolic disturbances and RAGE’s role(s) in the crosstalk between metabolic diseases and cancer.

In this respect, several reports suggest that modulation of RAGE activity might be of therapeutic importance for skeletal muscle in various disease states that have chronic inflammation and oxidative stress as common stigmata. Interestingly, a number of RAGE inhibitors (such as TTP488 and FPS-ZM1) have been developed, and some of them resulted well tolerated in human clinical trials. Potentially, these molecules might be useful for skeletal muscle diseases as well. However, caution should be exerted with the use of pharmacological blockers of RAGE signalling, since the complete absence of RAGE activity might severely compromise SC function and skeletal muscle regeneration. Modulation rather than inhibition of RAGE activity might represent a successful approach to restore muscle homeostasis in different pathological conditions affecting skeletal muscle tissue.
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Conflict of interest

The authors declare no conflict of interest.

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