Epidermal growth factor, latrophilin, and seven transmembrane domain-containing protein 1 marker, a novel angiogenesis marker

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Abstract: Epidermal growth factor, latrophilin, and seven transmembrane domain-containing protein 1 on chromosome 1 (ELTD1), an orphan adhesion G-protein coupled receptor, was reported as a regulator of angiogenesis, also involved in cancer progression and development. More recently, ELTD1 was identified as a potential new tumor marker for high-grade glioma. ELTD1, belongs to the G-protein coupled receptor superfamily that comprises the biggest receptor family in the human genome. Following the discovery of ELTD1 almost a decade ago, only a few research groups have attempted to find its role in normal and tumor cells, important information about this receptor remaining still unknown. The ELTD1 ligand has not currently been identified and intracellular signaling studies have not yet been performed in normal or tumor cells. Although the current published data on ELTD1 function and structure are rather limited, this receptor seems to be very important, not only as biomarker, but also as molecular target in glioblastoma. This review summarizes and discusses the current knowledge on ELTD1 structure, function, and its role in both physiological and tumoral angiogenesis.

Keywords: ELTD1, angiogenesis, glioma, biomarker

Introduction

Since the early ages of oncology, medical researchers have strived to discover new biomarkers, which can be correlated with early stages of cancer development, tumor progression, and resistance to therapy. With the discovery of targeted therapy, some of these markers became interesting therapeutic targets due to their implication in intricate pathophysiological processes involved in cancer growth, invasion, or resistance to curative approaches. One of the most studied pathophysiological events involved in cancer development and progression is angiogenesis. Depriving nascent cancer cells of their nutrients provided by aberrant, newly formed blood vessels became an attractive approach that, in theory, could deter or even reverse neoplastic growth. However, both early and more recent trials have yielded limited results due to the complex nature of microvasculature in each cancer type, the multiple pathways implicated in cell signaling, and the vast number of genes responsible for both normal and cancer angiogenesis.\(^1\)\(^-\)\(^5\) Together, these variables can confer both natural and acquired resistance to anti-angiogenic therapy, leading to mixed clinical results.\(^6\)\(^-\)\(^8\) Many angiogenic inhibitors, designated for cancer treatment, are currently in preclinical and clinical use. The vascular endothelial growth factor receptor family is a well-studied angiogenic class of receptors\(^9\)\(^-\)\(^11\) that have been constantly used to develop new cancer therapies for quite a while now.\(^12\)\(^-\)\(^14\) The therapeutic results, however, for many malignant diseases, in which vascular endothelial growth factor receptor dysfunction is believed
to drive tumor angiogenesis, have been ineffective. Future approaches should include inhibition of multiple pathways involved in cancer microvasculature formation or the discovery of new angiogenesis markers, more specific for each cancer type.15–18

One such possible candidate is epidermal growth factor, latrophilin, and seven transmembrane domain-containing protein 1 on chromosome 1 (ELTD1), a biomarker recently linked to both normal and pathological angiogenesis.19,20 An editorial article linked ELTD1 levels to how the human organism responds to hematopoietic stem cell transplant by developing graft-versus-host disease.21 Using complementary DNA microarray, another study also found that ELTD1 was significantly upregulated in patients with ulcerative colitis with dysplastic lesions when compared with normal, nondysplastic, and colic mucosa.22 Upregulation of the ELTD1 gene was observed in studies describing genomic adaption to high altitudes in the Andean population,23 cannabis use disorders,24 genetic predisposition for obesity in both humans and pigs,25 and even resistance to therapy in cattle afflicted by ticks.26

Initially mentioned in 2001, as a receptor normally regulated in the human heart,27 ELTD1 has been recently linked to the progression and development of glioblastoma multiforme (GBM),28,29 a particularly aggressive brain tumor with a very bleak prognosis.

Although it was discovered 14 years ago,30 in the scientific literature, only a few published works have dealt directly with this topic. The current review is an analysis of the information currently available on ELTD1.

**Origin and structure of the ELTD1 receptor**

Ontogenetically, ELTD1 belongs to the G-protein coupled receptor (GPCR) superfamily, which comprises the biggest receptor family in the human genome. Also called the 7α-helices transmembrane receptors, the GPCR superfamily transduces signals from the extracellular to the intracellular domain using guanine-binding proteins. The GPCR superfamily is involved in a multitude of human physiological processes, pertaining to different areas such as chemotaxis, smell, hormone secretion, taste, vision, or inflammation.30–32

Containing over 900 members, the GPCR superfamily comprises five major families: glutamate (22 members), rhodopsin (701 members), adhesion (33 members), frizzled (11 members), and secretin (15 members).33

The second biggest of the five families forming the GPCR super group, the adhesion family was initially considered as part of the secretin family, but was later revealed to be a distinct group. As a particular feature, this family presents an unusually elongated N-terminal ectodomain, which is often perceived as combinations of different adhesion-linked motifs: cadherins,34 thrombospondins,35,36 leucine-rich repeats,37 and epithelial growth factor (EGF)-like domain.38,39 Members of the adhesion family are implicated in multiple cellular functions such as cytoskeletal regulation,40–43 planar cell polarity,42,44,45 cell adhesion and migration,46–48 cellular proliferation and death,35,49–50 hematopoiesis, immunity,51–53 and angiogenesis.19,54–57 The adhesion family includes the latrophilin-like subfamily, with its four members: the orphan ELTD1 receptor, as well as the latrophilin 1, 2, and 3 receptors.58

Nechiporuk et al27 mapped the ELTD1 gene on the 1p33–p32 band, belonging to chromosome 1. The gene contains 15 exons having two splice variants: a full-length version and a shorter variant represented by a fraction of a transmembrane domain.39 The gene encodes a 3,527 nucleotide transcript, which further translates into a 690 amino acid protein. The ELTD1 receptor is formed of a large extracellular domain, a seven transmembrane domain coupled with a short cytoplasmic tail. The extracellular domain contains an N-terminal region, an EGF-like domain, a calcium-binding EGF domain, and a GPCR proteolysis site. The cytoplasmic domain contains a tyrosine kinase phosphorylation site, suggested to be involved in receptor intracellular signaling (Figure 1).27 Unfortunately, no information is yet available in current literature on the downstream signaling pathways implicated in signal transduction or the ligands that bind to the ELTD1 receptor.

**ELTD1 implication in angiogenesis**

Due to the low number of studies concerning ELTD1, its roles in both physiological and pathological angiogenesis are quite unknown. In the first study published in this area, Nechiporuk et al27 asserted that the ELTD1 receptor was highly expressed in cardiomycocytes and smooth muscle cells of both blood vessels and bronchi, in murine models. ELTD1 regulation was also linked to the switch between hyperplastic and hypertrophic evolution in the development of cardiomyocytes during the physiological differentiation process, in both human and rat models. The receptor’s presence in the coronary arteries also raises further question marks regarding its implication in coronary angiogenesis.27 More recently, another study mentioned that ELTD1 was actively implicated in cardiac hypertrophy, in rat models.63 This study also pointed out that ELTD1 impairment was linked to the faulty cardiomyocytic remodeling process, produced by
ELTD1 roles in cancer

GBM has long been considered an immense obstacle for modern medicine, being both the most common and the most aggressive brain tumor to date. Considered a grade IV glioma by the World Health Organization, GBM has an extremely poor prognosis with a vast majority of patients dying within 2 years of diagnosis, with less than 5% of them surviving more than 3 years.

Therapeutic approaches involving chemotherapy, radiotherapy, targeted therapy, and/or combined regimes have so far yielded very limited results with a poor contribution to GBM patient survival. So far, no viable biomarker has been selected as a strong candidate among the numerous gene alterations, protein overexpressions/suppressions, and abnormal signaling pathways in glioblastoma. One of the most well-known markers for GBM is the epithelial growth factor receptor (EGFR). EGFR amplification, along with mutations such as EGFRvIII/AEGFR, has long been considered as poor prognostic markers for GBM patients, but also viable therapeutic targets. Other biomarkers include mutations of the isocitrate dehydrogenase enzyme (IDH) with its two isoforms IDH1 and IDH2, considered the genetic signature for secondary GBMs. O°-methylguanine DNA methyltransferase protein expression and methylation status, which are considered markers for slower tumor progression, PTEN gene mutation. Circulating biomarkers such as extracellular vesicles were recently mentioned as valuable biomarkers for GBM. Extracellular vesicles are membrane-surrounded structures, secreted by cells in the biological fluids and consist of a mixture of several proteins, lipids, nucleic acids, glycans, and other biomolecules. These metabolites are suggested to be tumor-specific, thus considered to be good circulating biomarkers for GBM progression and response to the therapy. In a study by Towner et al conducted on human and rodent glioma samples, higher levels of ELTD1 were found in high-grade gliomas compared with low-grade gliomas. Higher levels of ELTD1 were also associated with the mesenchymal subtype compared with the proneural subtype. However, the authors didn’t find any correlations between ELTD1 levels and survival rates in both the human and murine models.

To analyze things from a broader perspective, an online database has also compared various tumor cell lines to see how ELTD1 levels correlate with different types of cancer (Figure 3). Surprisingly, in contrast with the previous studies we mentioned, the results placed ELTD1 expression in GBM lines midway through the list. More surprisingly, immunohistochemical staining did not find...
any ELTD1 levels in glioma tumor samples (http://www.proteinatlas.org/ENSG00000162618-ELTD1/cancer, accessed October 17, 2015). However, it should be noted that the database refers only to staining in glioma, without mentioning if the samples were taken from high- or low-grade gliomas. As Towner et al29 pointed out, higher levels of ELTD1 were associated with the more aggressive, high-grade gliomas in contrast to low-grade ones.

In order to better understand how the receptor works, alterations to the ELTD1 genetic sequence in different cancers were also taken into consideration (Figure 4). By comparison with other cancer types, ELTD1 alterations were almost nonexistent in malignant gliomas. These findings suggest that while a high level of ELTD1 expression may be indicative of the presence or aggressiveness of a malignant glioma, mutation levels are not linked to any aspect of tumor evolution.

ELTD1 has also been implicated in the angiogenesis of other types of cancer. A study of various tumor samples (renal, colorectal, ovarian, and head and neck) showed a match between the upregulation of the gene encoding ELTD1 and tumor development.19 Tissues of tumoral origin presented higher levels of protein expression when compared with normal tissue samples, while vascular smooth muscle
cells showed higher ELTD1 staining when compared with endothelial cells. Increased ELTD1 staining was also correlated with higher microvascular density levels and reduced hypoxia signature in some forms of cancer/tumor.19 The article also points out that ELTD1 impairment in endothelial cells halved the tumor cells’ capacity to sprout new microvessels. Furthermore, Masiero et al demonstrated, using both in vivo and in vitro models, that the two main antagonistic pathways involved in angiogenesis (VEGF/bFGF and DLL4-Notch)60–62 had an opposing effect on ELTD1 levels (the former increased ELTD1 expression while the latter determined the exact opposite)19 (Figure 1). A study by Dieterich et al also demonstrated that in human glioblastomas, increased VEGF-A signaling produced higher levels of ELTD1 gene expression in endothelial cells. Perhaps with a better understanding of the processes behind how ELTD1 works, which are currently unknown, such as ligand–receptor interaction and signaling pathways or the effect mutations have on the

Figure 3 ELTD1 protein levels in human cancer cell lines. The results show the intensity of antibody staining (protein levels) for each cell line. Results were obtained from the Human Protein Atlas Database (http://www.proteinatlas.org/ENSG00000162618-eLTD1/cell/HPA025229, accessed October 17, 2015).

Abbreviation: ELTD1, epidermal growth factor, latrophilin, and seven transmembrane domain-containing protein 1 on chromosome 1.
receptor function, we may draw a parallel between ELTD1 and the well-studied VEGF and in terms of involvement in both normal and neoplastic angiogenesis. However, because of the scarcity of information regarding ELTD1 due to the relatively recent discovery of the receptor, further studies are required to confirm its involvement in these mechanisms.

**Conclusion**

Recent studies have linked ELTD1 levels to both normal and pathological angiogenesis, such as cardiac hypertrophy in mice and even the development of nouvelle microvessels in various cancer types. One association that stands out is the link between ELTD1 and the evolution of GBM, recently pointed out in various studies. If proven feasible, this association may provide not only a viable biomarker for this highly aggressive cancer, but also a target for future therapeutic agents. These therapeutic approaches should focus on either silencing the ELTD1 gene using methods such as small interfering RNA or blocking the receptor with antibodies. However, with little knowledge about the ligands binding to the receptor or the pathways involved in downstream signal transduction, targeting ELTD1 might still prove problematic.

With several other roles besides vascular proliferation in cancer such as graft-versus-host disease, cannabis addiction, obesity or adaption to altitude in certain populations and notably the presence in a large number of normal tissues, ELTD1 may play a larger role than expected in the human body. Hopefully, increased attention to this receptor may shed some light on its multiple functions leading to a better understanding on how some pathological mechanisms are interrelated and how they can be tackled.

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**Author contributions**

All authors contributed toward the literature search, data analysis, drafting and critically revising the paper and agree to be accountable for all aspects of the work.

**Disclosure**

The authors report no conflicts of interest in this work.
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