1. Introduction

Plasminogen receptors (Plg-Rs) are a broadly distributed and heterogeneous group of cell surface proteins that share a common feature, the ability to interact with plasminogen (Plg) and plasmin. The list in Table 1, not necessarily all inclusive, identifies 12 different Plg-Rs. Many of these Plg-Rs are expressed by many different cell types, and many are present on the same cell type. Indeed, the number of Plg binding sites on any particular cell type can be extraordinarily high (range from $10^5$ to $10^7$ Plg binding sites per cell). The similarities among these Plg-Rs are very limited and appear to rest only on their ability to be expressed at cell surfaces where they can display their Plg and Plm binding function. Nevertheless, this binding function allows many different Plg-Rs to orchestrate diverse biological responses including fibrinolysis, inflammation, wound healing, and angiogenesis. The question then arises as to why there are so many Plg-Rs and whether there is a plausible explanation for this extensive functional redundancy? This paper will consider these basic questions. As a forewarning, we do not purport to provide clear answers to these questions but hopefully our speculations will be challenging and stimulating.

2. So Many Plg-Rs: Do Different Plg-Rs Bind Plg Differently?

Almost all of the Plg-Rs listed in Table 1 engage the lysine binding sites (LBS) of Plg and Plm by virtue of a C-terminal lysine or by presenting an internal amino acid residue in a context that mimics a C-terminal lysine. As a consequence of a common mechanism of engagement, Plg-Rs are projected to enhance Plg activation by either urokinase plasminogen activator (uPA) or tissue plasminogen activator (tPA), to enhance the catalytic activity of plasmin and to protect bound plasmin from inactivation by inhibitors [1–3]. Indeed, several Plg-Rs have been reported to have one or more of these functional attributes [4–6]. Also, with a similar mechanism of binding, the affinities of the various Plg-Rs for Plg should be similar. The context of the LBS binding residue within a Plg-R might be influenced by adjacent amino acids or local conformation and thereby influence the affinity of specific subset of Plg-Rs for Plg. However, even for Plg-Rs that utilize an internal residue rather than a C-terminal lysine to engage Plg, affinities for the ligand appear to be similar ($\sim 1 \mu M$), [7]. One potential exception to this assertion could be the annexin A2/p11
Table 1: Plg-Rs on various cell types.

<table>
<thead>
<tr>
<th>Plg-Rs</th>
<th>Cell types</th>
<th>C-terminal lysine</th>
<th>Major cellular localization</th>
<th>Secretory pathways</th>
</tr>
</thead>
<tbody>
<tr>
<td>(1) Annexin A2</td>
<td>Endothelial cells, monocytoid lineage</td>
<td>Absent*</td>
<td>Cytosol and or nucleus</td>
<td>Translocation depends on p11 and phosphorylation; activity of L-type like Ca(^{2+}) channels and intracellular Ca(^{2+}); associates with plasma membrane via phosphatidylserine.</td>
</tr>
<tr>
<td>(2) Actin</td>
<td>Endothelial cells, carcinoma, catecholaminergic cells, PC-3, HT1080</td>
<td>Absent</td>
<td>Cytoskeleton</td>
<td>Not known</td>
</tr>
<tr>
<td>(3) Amphoterin</td>
<td>Neuronal cells</td>
<td>Absent</td>
<td>Cytoplasmic and extracellular</td>
<td>Not known</td>
</tr>
<tr>
<td>(4) αVβ3</td>
<td>Endothelial cells</td>
<td>Absent</td>
<td>Integral membrane protein</td>
<td>Classical endoplasmic reticulum and Golgi pathway</td>
</tr>
<tr>
<td>(5) αMβ2</td>
<td>Neutrophils, monocytes, macrophages</td>
<td>Absent</td>
<td>Integral membrane protein</td>
<td>Classical endoplasmic reticulum and Golgi pathway</td>
</tr>
<tr>
<td>(6) αIIbβ3</td>
<td>Platelets, RA synovial fibroblasts</td>
<td>Absent</td>
<td>Integral membrane protein</td>
<td>Classical endoplasmic reticulum and Golgi pathway</td>
</tr>
<tr>
<td>(7) Cytokeratin 8</td>
<td>Hepatocellular, breast carcinoma, Monocytes, neutrophils, carcinoma, lymphoid, myoblast neurons</td>
<td>Present</td>
<td>Cytoskeleton</td>
<td>Not known</td>
</tr>
<tr>
<td>(8) α-Enolase</td>
<td>Neutrophils, monocytes, monocytoid cells, Endothelial cells</td>
<td>Present</td>
<td>Cytosol</td>
<td>L-type-like Ca(^{2+}) channel and intracellular Ca(^{2+})</td>
</tr>
<tr>
<td>(9) Histone 2B</td>
<td>Neutrophils, monocytoid cells, Endothelial cells</td>
<td>Present</td>
<td>Nucleus</td>
<td>L-type-like Ca(^{2+}) channel and intracellular Ca(^{2+}), associates with plasma membrane via phosphatidylserine and heparin sulfate</td>
</tr>
<tr>
<td>(10) P11</td>
<td>Endothelial cells, HT1080 cells</td>
<td>Present</td>
<td>Cytosol and or nucleus</td>
<td>L-type-like Ca(^{2+}) channel and intracellular Ca(^{2+}), associates with multiple plasma membrane binding partners, including annexin 2</td>
</tr>
<tr>
<td>(11) Plg-RKT</td>
<td>Monocytes, macrophages, neuronal cells</td>
<td>Present</td>
<td>Integral membrane protein</td>
<td>Classical endoplasmic reticulum and Golgi pathway</td>
</tr>
<tr>
<td>(12) TATA-binding protein-interacting protein</td>
<td>Monocytoid cells</td>
<td>Present</td>
<td>Nucleus</td>
<td>Not known</td>
</tr>
</tbody>
</table>

* requires cleavage to bind Plg [8].

heterotetramer, where the proximity of multiple Plg binding sites within a single molecular species could enhance affinity substantially. To support this possibility or other reports of higher-affinity Plg-Rs, variability in ligand preparations used (e.g., presence of Lys-Plg in Glu-Plg preparation) must be controlled. Furthermore, since ligand availability seems not to be limiting (Plg is present at high concentrations), differences in apparent affinity may have less impact than anticipated.

### 3. So Many Plg-Rs: Do Different Cell Types Use Different Plg-Rs?

Not all Plg-Rs are expressed on all cell types. As an example of a Plg-R with a restricted cellular distribution, integrin αMβ2 is a Plg-R [7] and its expression is confined to leukocytes. However, leukocytes express many other Plg-Rs, including annexin A2/p11, which has long been promulgated as the major Plg-R on endothelial cells (ECs). Indeed, inactivation of either the annexin A2 or p11 genes does affect EC-dependent responses, including angiogenesis, tumorogenesis, fibrinolysis, and inflammation [9–12]. However, ECs do express other Plg-Rs. As an illustrative example, histone H2B, a high-abundance Plg-R on monocyteid cells, is also readily detected on the surface of HUVEC (Figure 1). In Figure 1, H2B was detected on the surface of HUVEC by a cell-surface biotinylation approach [13] in which the cells were surface labeled with biotin, lysed, and the biotinylated proteins were isolated on streptavidin beads and then identified by western blotting with specific antibodies (see Figure 1 and its legend for details). H2B was labeled with biotin, whereas p65, a control intracellular protein, was not even though both H2B and p65 were readily detected in the whole cell lysates of HUVEC. H2B associates with the surface of monocyteid cells by binding to phosphatidylserine (PS) [14]. Annexin V, another PS binding protein displaces the H2B from the surface of monocyteid cells [14] and also chases biotin labeled H2B from the surface of HUVEC (see
Plg activation. However, recent data have suggested that one Plg-R is particularly proficient in the leading edge of migrating retinal glial cells and malignant glioma. Furthermore, annexin 2 has been localized to the leading edge of migrating cells [17], and uPAR also localizes to the leading edge of migrating cells [16]. Thus, in addition to annexin A2/p11, several Plg-Rs have been detected on endothelial cells. Hence, the notion of the preeminence of a specific Plg-R on a particular cell type does not seem tenable.

The compartmentalization of specific Plg-Rs to select locations on the cell surface could provide a mechanism to distinguish the function of one Plg-R from another. Several integrins serve as Plg-Rs (Table 1) and integrins do localize to the leading edge of migrating cells [16], and uPAR also localizes to the leading edge of migrating cells [17]. Furthermore, annexin 2 has been localized to the leading edge of migrating retinal glial cells and malignant glioma cell [18, 19]. Thus, an advantageous microenvironment may be created in which one Plg-R is particularly proficient in Plg activation. However, recent data have suggested that cell-surface-bound Plg can be efficiently activated or even more efficiently activated by uPA bound to another cell than that on the same cell [20]. The boost in efficiency of Plg activation gained by localization on a single cell may be offset by the restricted diffusion or orientation of the Plg activator on the cell surface. Thus, localization of certain Plg-Rs to a specific microdomain on the cell surface and the functional advantage of such localization remain a possibility. We did note a uniform distribution of several Plg-Rs, as well as bound Plg, on monocyteoid cells by confocal microscopy although changes in distribution under stimulated conditions were not tested [13]. A common mechanism dependent on L-type like calcium channels has been implicated in translocation of several Plg-Rs to the surface of monocyteoid cells [21], but the mechanisms by which these Plg-R tether to the cell surface are distinct [14]. Hence, Plg-Rs could compartmentalize on the cell surface.

4. So Many Plg-Rs: Are Plg-Rs Differentially Regulated on Cells?

It is well established that Plg binding to cells can be markedly modulated; changes in Plg binding capacity of specific cell types can increase 3- to 20-fold in response to specific stimuli. Cellular events and responses that can induce such changes include oncogenic transformation (breast and adenocarcinoma cancer) [22, 23] differentiation (monocytes, adipocytes) [21, 24], agonist stimulation, (leukocytes, endothelial cells, platelets) [7, 25–27], adhesion (monocyteoid cells) [28], and apoptosis (monocyteoid cells) [14]. In addition, Plg binding can be enhanced by proteolysis of existing cell surface proteins to generate N-terminal lysines [29, 30]. This latter mechanism for exposing new Plg-Rs can be triggered by plasmin itself and depends on the availability of uPA on the cell surface [31]. Thus, even though a cell type can express multiple Plg-Rs, a subset of Plg-Rs may be differentially upregulated and utilized to mediate a specific cellular response.

The data in Figure 2 provides an illustrative example of how different Plg-Rs maybe utilized by the same cell in responding to different stimuli. THP-1 monocyteoid cells were either stimulated to undergo differentiation using vitamin D3 + IFNγ or apoptosis using camptothecin. Consistent with our prior report [13, 14], the cells respond to these stimuli by markedly upregulating their Plg binding capacity. In association with differentiation, Plg binding increased by 3.3-fold. Of the Plg-Rs analyzed by FACS, enolase, annexin2, p11, and H2B, surface expression increased most markedly for H2B (4.7-fold) in response to differentiation. In response to apoptosis induced by camptothecin, Plg binding increased by 10-fold. While surface localization of H2B did increase significantly (4.6-fold), much more striking was the 20-fold upregulation of p11 in the camptothecin-treated THP-1 cells. This pattern of enhanced p11 expression was also observed in U937 monocyteoid cells treated with camptothecin, where 5.8-fold increase of Plg binding was associated with 6.3-fold increase in p11 expression. Of note, these increases in p11 expression on apoptotic cells were not paralleled by substantial increases of the annexinA2 subunit. In the camptothecin-treated THP-1 cells, surface expression of the annexinA2 subunit increased by 2.8-fold and for U937 cells, the increase was 2.3-fold. As explanations for this disproportional upregulation of p11, the subpopulation of annexinA2 molecules that escort p11 to the cell surface may not react with the antibody used in this analysis, or the anti-p11 may selectively penetrate apoptotic cells, which are known to be leaky [32]. A more interesting possibility is that a portion of the p11 that becomes surface expressed is in a free form or is associated with other binding partners.

Figure 1: H2B exposure on the surface of endothelial cells. Human umbilical vein endothelial cells (HUVECs) were either untreated or treated with annexin V (250 nM) for 48 hr. Cells were surface labeled with biotin, and the biotinylated proteins were isolated using streptavidin-conjugated beads. H2B and p65 (a transcription factor with a cytosolic and nuclear localization) that were bound and eluted from the streptavidin beads were detected by western blotting with a rabbit anti-H2B or rabbit anti-p65. The absence of biotinylated p65 serves as a control for surface labeling of H2B. Band intensities of the western blots were analyzed using Kodak ID 3.6 software, and net intensity (NI) of each band is indicated below each lane. In each set of two lanes, the right-hand lane is in the presence of annexin V and the left-hand lane in its absence. (WCL: whole cell lysates).
Besides annexin2, other plasma membrane proteins, NaV1.8 sodium channel, TASK1 potassium channel, TRPV5/TRPV6 channels, and cathepsin B [33] have been shown to interact with p11, could assist in its transport to the cell surface, and may still further extend the repertoire of Plg-Rs expressed by monocyteid cells.

*In vivo* data also support the proposition that different Plg-Rs mediate the response of the same cell type to different stimuli. In a thioglycollate-induced peritonitis model, an antibody to H2B that blocks Plg binding inhibited macrophage recruitment by ~50% while an antibody to α-enolase that also blocks Plg binding to its target produced less than 25% inhibition of macrophage recruitment [13]. In contrast, in an LPS-induced lung inflammation model, Plg binding to α-enolase overexpressing U937 cells produced a substantial enhancement of macrophage migration [34].

**5. So, Why So Many Plg-Rs?**

While the utilization of different Plg-Rs to orchestrate different cellular responses is supported by data cited above, blocking of several different Plg-Rs has been shown to markedly suppress what appears to be the same inflammatory response thioglycollate-induced peritonitis. The contribution of H2B (45% [13]), p11 (53% [12]), and Plg-RKT (58% [35]), either with antibodies or gene inactivation, exceeds 100%. Such extensive inhibition becomes even more incomprehensible since macrophage recruitment is decreased by only 65% in Plg<sup>−/−</sup> mice compared to wild-type littermates [36]. At least four explanations can be considered to explain such observations. First, these various Plg-Rs may exert an effect on macrophage recruitment unrelated to Plg. The effect of blockade of individual Plg-Rs in a Plg<sup>−/−</sup> background could be used to identify such functions. Second, a threshold of bound Plg must be attained in order for Plg to facilitate cell migration. No single Plg-R may harness sufficient Plg to reach this threshold, and, hence, cooperation among several Plg-Rs is required. Third, while many different Plg-Rs enhance Plg activation, the intracellular signaling responses that they elicit may be distinct. Cellular recruitment is a complex response requiring activation of many different intracellular signaling pathways. Different Plg-Rs may trigger distinct signaling events, and these pathways may need to cooperate to yield efficient migration. Blunting the signaling
response elicited by occupancy of any one Plg-R may lead to suppressed signaling and diminished migration. Fourth, recruitment into the peritoneum is a temporally extended and multi-step response, and different Plg-Rs may come into play at different times and stages during the response. Hence, difference Plg-Rs may be utilized to achieve specific steps in the recruitment cascade.

6. Summary and Concluding Remarks

In this brief discussion, we have raised the question as to why there are so many Plg-Rs. With so many different receptors frequently, with many of them expressed on the same cell type, it is difficult to envision how the cell would prioritize its utilization among these multiple Plg-R. Affinity differences between Plg-Rs for their Plg and plasmin ligands could distinguish one receptor from another, but this can only be tested by direct comparisons among Plg-Rs. Utilization of specific Plg-Rs to mediate tissue specific or cell specific responses can also be envisioned, but such analyses again mandate comparative studies. In fact, in each of the explanations suggested above, to account for the profound role of many different Plg-Rs in what is globally visualized as a single cellular response, macrophage recruitment into the peritoneum, comparative studies are again needed. The goal of such comparative studies is not to prove that one particular Plg-R is better than another, but rather to help dissect the ways in which Plg orchestrates cell migration and other cellular responses in vivo.

Acknowledgments

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References

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