PLOD2 Is Essential to Functional Activation of Integrin \(\beta_1\) for Invasion/Metastasis in Head and Neck Squamous Cell Carcinomas

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HIGHLIGHTS

- PLOD2 specifically regulates intracellular localization of integrin \(\beta_1\)
- The hydroxylation on integrin \(\beta_1\) by PLOD2 is critical for its stability
- PLOD2 and integrin \(\beta_1\) expression colocalized at the invasive front of SCC tissues
- PLOD2 is necessary for tumor invasion/metastasis through the integrin \(\beta_1\) maturation

Stability and membrane translocation of integrin \(\beta_1\), Progression of cell invasion / metastasis

Loss of translocation and degradation of integrin \(\beta_1\), Reduction of cell migration

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PLOD2 Is Essential to Functional Activation of Integrin β1 for Invasion/Metastasis in Head and Neck Squamous Cell Carcinomas

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SUMMARY

Identifying the specific functional regulator of integrin family molecules in cancer cells is critical because they are directly involved in tumor invasion and metastasis. Here we report high expression of PLOD2 in oropharyngeal squamous cell carcinomas (SCCs) and its critical role as a stabilizer of integrin β1, enabling integrin β1 to initiate tumor invasion/metastasis. Integrin β1 stabilized by PLOD2-mediated hydroxylation was recruited to the plasma membrane, its functional site, and accelerated tumour cell motility, leading to tumor metastasis in vivo, whereas loss of PLOD2 expression abrogated it. In accordance with molecular analysis, examination of oropharyngeal SCC tissues from patients corroborated PLOD2 expression associated with integrin β1 at the invasive front of tumor nests. PLOD2 is thus implicated as the key regulator of integrin β1 that prominently regulates tumor invasion and metastasis, and it provides important clues engendering novel therapeutics for these intractable cancers.

INTRODUCTION

Head and neck cancers are the sixth most common malignant tumors worldwide. They are mainly treated with extended resection or radiation/chemotherapy, but functional deteriorations in swallowing, vocalization, and respiration lead to clinical decline in quality of life (QOL). Even in cases of combined surgical treatment with radiation therapy and chemotherapy, local recurrence including primary lesion and surrounding lymph nodes, is usually observed in approximately 30% of the patients. As distant metastases, as in the lung, can be seen in 25% of cases, recurrence and therapeutic resistance are very conspicuous. The 5-year survival rate in advanced cases of stage III and above is about 40%; thus the patients still present a poor prognosis (Laramore et al., 1992). Recently, cetuximab, a molecular targeting agent, has been administered in combination with cisplatin to patients with head and neck malignancies, showing some benefit to outcomes. However, it remains that subjects suffer from serious side effects such as dermatitis and mucositis, and novel avenues for cancer therapeutics are needed for improved antitumor properties and patient QOL (Bonner et al., 2006).

Previous studies revealed that 2-oxoglutarate and the iron-dependent dioxygenases superfamily function as a hydroxylase/demethylase and that they hydroxylate or demethylate molecules such as transcription factor, histones, and DNA as substrates. Indeed, it has been reported that these enzymes play various roles in cell cycle and gene expression and control of invasion/metastasis of cancer cells in multiple cell lines via modified molecules (Markolovic et al., 2015).

There are some reports that procollagen lysyl hydroxylase (PLOD), which belongs to this superfamily, is involved in the hydroxylation of collagen molecules as the only target molecule identified to date, and it is considered to be an essential enzyme for cross-linking reaction between collagen molecules outside the cell (Wu et al., 2006). PLOD is composed of an N-terminal endoplasmic reticulum-body translocation signal, a hydroxyl-catalyzed domain at the C-terminus, and hydroxylates lysine residues located in the terminal region of the collagen precursor in the ER.

Currently, three members of the PLOD family molecular group are found to share high homology: PLOD1, PLOD2, and PLOD3 (Valtavaara et al., 1997; Passoja et al., 1998a). Tissue expression analysis by Northern blot showed that PLOD2 and PLOD3 had tissue-specific expression unlike the broad expression of PLOD1.
Specific Upregulation of PLOD2 and Its Effect on Cellular Motility of Oral SCC Cells

Since the 2-oxoglutarate and iron-dependent dioxygenases have been classified into several groups according to the target amino acid residue, such as proline hydroxylase, asparagine/aspartate hydroxylase, etc., we first examined their expression in each category to identify specific hydroxylases highly expressed only in SCC lines from the oral cavity, in contrast to the non-neoplastic cellular counterpart (non-tumorigenic immortalized keratinocyte, HaCaT). Three representative human oral SCC lines (HSC-2, HSC-3, Ca9-22) derived from different patients were examined in comparison with HaCaT, and gene expression of typical hydroxylases for each category was examined by a semiquantitative RT-PCR method (Figure S1A). The expression of JMJ2A, JMJ2D, EGLN2, MINAS3, NOA6, HIF1AN, FTO, and TET1 did not show a significant difference between the tumor cells and HaCaT. On the other hand, the expression of PLOD2, EGLN1, EGLN3, OGFOD1, ASPH, and ALKBH3 revealed different differences in expression among these cell lines, especially in that the mRNA level of PLOD2 in the SCC lines showed the most prominent and consistent increase at four to eight times compared with HaCaT (Figures S1A and 1A). This upregulation of PLOD2 mRNA in tumor cells was also corroborated by protein levels in immunoblot analysis even among the lysyl hydroxylase family including PLOD1 and PLOD3 (Figure 1B). Immunofluorescence analysis with anti-PLOD2 antibody revealed that endogenous PLOD2 was predominantly localized to the ER which was confirmed by GFP-labeled ER marker (Figure 1C). Thus, tumor-specific upregulation of the expression was observed only in PLOD2 among the examined hydroxylase and demethylase family, and the mRNA and protein levels of PLOD1 and PLOD3 were not specifically elevated in the tumor cells although they are categorized to the same family.

RESULTS

Specific Upregulation of PLOD2 and Its Effect on Cellular Motility of Oral SCC Cells

In this study, apart from tumor microenvironmental factors, we focused on biological function of PLOD2 on cancer cell itself using oral squamous cell carcinoma (SCC) lines as a model because they are high in PLOD2 expression, especially in tumor invasion/metastasis highlighting specific interaction between PLOD2 and integrin β1. We have found that PLOD2 induces the hydroxylation of integrin β1, which plays an essential role in the stability and cellular localization of integrin β1 for its functional activation. The integrin β1 requires specific interaction with PLOD2, which will lead to greater understanding of molecular mechanisms for cancer invasion/metastasis and foster innovation in therapies against refractory malignancies.

In particular, PLOD2 has been reported to be induced in response to differentiation of bone marrow stromal cells (Veltavaara et al., 1998; Uzawa et al., 1999). Gene mutations in PLOD1 and PLOD3 have been identified in the case of Ehlers-Danlos syndrome (fragile skin, hematomas, and joint hypermobility), whereas PLOD2 mutations are clinically reported as involved in the Bruck syndrome characterized by imperfect bone formation (Walker et al., 2004; van der Slot et al., 2003).

Furthermore, based on the previous studies, PLOD2 has been reported to be specifically induced by activation of transcription factor HIF-1α in response to hypoxia and TGF-β1 stimulation on tumor stroma, cancer cells (breast cancer, hepatocellular carcinoma), and sarcoma (mouse sarcoma model) (Chen et al., 2015; Gilkes et al., 2013a, 2013b; Eisinger-Mathason et al., 2013).

It is known that collagen secretion and remodeling of extracellular matrix (ECM) are accelerated especially in cancer stromal cells, leading to invasion and metastases of cancer cells (Gilkes et al., 2013a). The association of HIF-1α-dependent increase in expression of PLOD2 and epithelial mesenchymal transition (EMT) has been established in human glioma cells, and glioma patients with high expression of PLOD2 showed a poor prognosis (Xu et al., 2017). However, the intracellular substrate and tumorigenic activity of PLOD2 expressed in cancer cells remains unclear, especially in refractory cancers, such as oral, head, and neck cancers.

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Figure 2. PLOD2 Is Essential for Stabilization of Integrin β1

(A) Immunofluorescence revealed expression, and localization of CDH1 was not affected by siPLOD2-treatment in SCC cells.
(B) Expression and intracellular localization of integrin β1 of the SCC cells was examined at 48 h after treatment with siPLOD2. Cytoskeleton and nuclei were stained with phallolidin and Hoechst, respectively. Scale bar = 20 μm.
(C) Expression of integrin β1, CDH1, and SNAIL in the siPLOD2-transfected cells by immunoblotting using anti-PLOD2, anti-integrin β1, anti-CDH1, and anti-SNAIL Ab, respectively.
(D) Semi-quantitative expression of integrin β1 mRNA by RT-PCR with or without siPLOD2-treatment.
(E) Comparative ratio of integrin β1 mRNA in siPLOD2-treated cells based on the quantitative PCR results. Quantitative results are mean ± s.d. from three biological replicates (n.s. = not significant, Student’s t-test).
(F) Restoration of integrin β1 by treatment with MG132 and chloroquine (CHQ). HSC-2 cells pretreated with siPLOD2 were examined for integrin β1 expression 18 h after treatment with MG132 (1 nM) or CHQ (50 μM), respectively. Expression of integrin β1 protein by immunoblotting (upper panel), intracellular localization of integrin β1 by immunofluorescence using anti-integrin β1 Ab (lower panel). Integrin β1 (red) was merged with lysosome marker (Lyso-GFP). Scale bar = 20 μm.
(G) Effect of PLOD2 mutant lacking the catalytic domain (ΔPKHD) to integrin β1. Integrin β1 of the HSC-2 transfected with myc-tagged PLOD2 lacking the hydroxylase domain (ΔPKHD) compared with that of the cells transfected with the WT. Reduction of integrin β1 detected by immunoblotting (upper panel) and the loss of plasma membrane localization indicated by arrowhead with immunofluorescence (lower panel). Scale bar = 20 μm.
(H) Wound healing assay revealed cell migration was affected in the ΔPKHD-transfected cells as shown in the graph (upper panel) and migratory images (lower panel). Each symbol in the graph represents empty vector (circle, black), PLOD2 WT (square, blue), and PLOD2 ΔPKHD mutant (triangle, red). Data are means ± s.d. from three technical replicates for one biological replicate (*p < 0.05, Student’s t-test as compared with empty vector).

Based on these results, we next studied the cellular motility of the SCC cells, HSC-2, HSC-3, and Ca9-22, treated with specific siRNA against PLOD2 (siPLOD2) in comparison with the cells treated with siPLOD1 or siPLOD3. Specificity of the siRNA for each PLOD isoform (siPLOD1, siPLOD2, and siPLOD3) was confirmed by immunoblotting, which demonstrated that knockdown by these siRNAs did not affect other isotypes (Figures S1B and 1D). In the wound-healing assay, each stable GFP-expressing clone established from parental HSC-2, HSC-3, and Ca9-22 was employed for evaluation. Mobilization of each clone, visualized by fluorescence microscope 24 h after treatment with the siRNA, revealed that only attenuated expression of PLOD2 significantly affected cellular migration among these three SCC cells, whereas neither PLOD1-knockdown nor PLOD3-knockdown critically disturbed migration of the tumor cells (Figure 1E, upper images and Figure S1B). Inhibition of wound closure was over 50% in siPLOD2-treated cells at 24 h following siRNA treatment (Figure 1E, lower graphs); however, the MTT assay indicated that tumor cell proliferation was not affected by siRNA-treatment in all PLOD isoforms (Figure S1C). These data implied that PLOD2 might be deeply involved in regulating tumor cell motility.

Crosstalk between PLOD 2 and Integrin β1 in Cellular Motility

On the basis of these findings, we focused on the specific role of PLOD2 in tumor cell motility. Generally, acceleration of cell mobility is closely related to invasive properties of tumor cells, and we examined whether expression of E-cadherin (CDH1) as a marker of epithelial-mesenchymal transition (EMT) was altered with or without siPLOD2-treatment of the SCC cells. We found that PLOD2-knockdown did not affect the expression and membrane localization of CDH1 for all SCC cells (Figures 2A and S2A). Because all of these three SCC lines highly express integrin β1 as one of the critical motor molecules for invasion, we next examined the effect of PLOD2 on integrin β1 therein. In siPLOD2-treated tumor cells, lack of filopodia development was observed as a morphological change as confirmed by phalloidin staining, and a marked decrease in integrin β1 was revealed, in contrast to the cells treated with control siRNA (Figures 2B, S2B, S2C, and S3). This phenomenon was not seen in siPLOD1- or siPLOD3-treated cells (Figure S1D). The siPLOD2-treated tumor cells with decreased integrin β1 showed no altered expression of CDH1 or SNAIL, suggesting the PLOD2 seemed not to be involved in EMT at least in these SCC cells (Figures 2C and S4A). Involvement of integrin β1 in migration of these SCC cells was demonstrated by knockdown assay using silintegrin β1 (Figures S4B and S4C). Taken together, our data indicate that integrin β1 appears directly regulated by PLOD2 for these tumor cells in an EMT-independent manner.

Next, to clarify whether PLOD2 affects induction of integrin β1 mRNA, or directly modifies the integrin β1 protein, RT-PCR was first performed to examine fluctuations in mRNA levels. Ultimately, no significant alteration in integrin β1 mRNA expression with or without siPLOD2 introduction was detected in SCC cells, which was further confirmed by qPCR (Figures 2D and 2E). Therefore, siPLOD2 did not affect induction of integrin β1 mRNA, i.e. the result suggested that integrin β1 protein might be persistently produced in tumor cells but may require a certain modification by PLOD2 for stabilization.
Figure 3. Specific Coupling of Integrin β1 with PLOD2

(A) The construction of PLOD2 expression fused Monti-red tag (PLOD2-Red) and integrin β1 expression fused Ash tag (Integrin β1-Ash) (left panel). Schematic representation of mechanism for intracellular fluorescent dot formation in response to specific protein-protein interaction (right panel).

(B) Marked aggregation of fluorescent dots was detected only in the HeLa cells cotransfected with PLOD2-Red and Integrin β1-Ash. Scale bar = 20 μm.

(C) The fluorescent dots were merged with GFP-labeled ER marker (ER-GFP). Arrowhead indicated each PLOD2, ER-GFP, and colocalized proteins in the cotransfected HeLa cells, respectively. Scale bar = 20 μm.

(D) The fluorescent dots were observed at lamellipodia of the PLOD2-Red-Integrin β1-Ash cotransfected cells or the Integrin β1-Red-PLOD2-Ash cotransfected cells. Hatched box indicated hyperview field in the right. Scale bar = 20 μm.

(E) Domain organization of the FLAG epitope tagged-Integrin β1 expressor, Myc epitope-tagged PLOD2 mutant expressor lacking PKHD domain (ΔPKHD) used in the immunoprecipitation assay.

(F) Integrin β1 was coprecipitated both with WT-PLOD2 and ΔPKHD-PLOD2 in HEK293T cells. Expression of each PLOD2 expressor and EGFP expressor as a control in whole cell lysates (left), immunoprecipitated PLOD2, and its variant (probed with anti-Myc Ab) using anti-Flag Ab for precipitation (right). IgG heavy chain (H.C.) and light chain (L.C.) are shown.

Following the recent study reporting degradation of integrin α at lysosome post-ubiquitination (Lobert et al., 2010), we examined the effects of proteasome inhibitor, MG132, and chloroquine (CHQ)—which is known to interfere with endocytosis processes and with ligand delivery to the lysosome (Erbacher et al., 1996)—as to whether they might restore the integrin β1 protein in siPLOD2-treated tumor cells. Immunoblotting demonstrated that both MG132 and CHQ successfully restore integrin β1 to the level of the control siRNA-treated cell as shown in Figure 2F (upper panel) and Figure S5A and that intracellular localization of integrin β1 was observed in the siPLOD2-introduced cells treated with MG132 and with CHQ, whereas membranous integrin β1 was only restored in MG132-treated cells (Figure S5B). The incidence of lysosomal localization of integrin β1 remains at the same level in all cases (Figure S5C).

PLOD2 Regulates Intracellular Localization of Integrin β1 Through Their Coupling

To examine whether PLOD2 specifically regulates intracellular localization of integrin β1, “Fluorescent based technology detecting Protein-Protein Interaction in living cells” (the Fluoppi system) was employed to visualize the movement of integrin β1 in the presence of PLOD2 (Figure 3A). When the two proteins of interest specifically bind to each other, it yields aggregated fluorescence foci (red colored dots) inside cells under the Fluoppi system. As shown in Figure 3B, large fluorescence foci in HeLa cotransfected with PLOD2-Red and integrin β1-Ash demonstrated specific binding of PLOD2 to integrin β1 within cells. Further we did semiquantification of fluorescent dot intensity on Fluoppi specimens between PLOD2-alone-expressing cells and integrin β1–PLOD2-coexpressing cells (Figures S6A and S6B). We also repeated Fluoppi assay to obtain clear image of fluorescent dot formation only on the integrin β1–PLOD2 cotransfection into the PLOD2-knockout cells (Figure S6C). Tracking the PLOD2-integrin β1 complex by Fluoppi system using integrin β1-Red expressor in combination with PLOD2-Red, the fluorescent foci were first colocalized with ER-GFP in the early phase of transfection, suggesting they were at the ER (Figures S6C and 3D) and then Monti-red-labeled integrin β1 protein transited to filopodia and the plasma membrane in the cotransfected cells (Figure 3D). Immunoprecipitation assay using anti-Flag Ab on the cotransfected HEK293T cell lysates indicated binding of integrin β1 both to full-length PLOD2 and ΔPKHD-PLOD2, which demonstrated direct binding of these two proteins, and integrin β1 bound to the site outside of the PKHD domain of PLOD2 (Figures 3E and 3F).
Figure 4. Hydroxylation of Integrin \( b1 \) in Presence of PLOD2 and Its Significance in Intracellular Localization of Integrin \( b1 \)

(A) LC/MS of integrin \( b1 \) purified from lysate of the 293T cells coexpressing integrin \( b1 \)-Flag and WT-PLOD2 in comparison with that from the cells expressing integrin \( b1 \)-Flag and \( \Delta \)PKHD-PLOD2. The main peaks of integrin \( b1 \) were highlighted with the hatched box.

(B) Hyperview of the highlighted peaks. Arrowhead represented the fragment of integrin \( b1 \) (position of \#651-658 a.a. containing three lysines; AFNKGEKK) from the WT-PLOD2-transfected or form the \( \Delta \)PKHD-PLOD2-transfected lysate, which showed shift of the peak between these two integrin \( b1 \) (spectra; 484.5 to 486 m/z).

(C) Recombinant PLOD2-6xHis protein from HEK293T was purified in 150 mM imidazole using cobalt resin. Purification of PLOD2 was confirmed by Coomassie staining and immunoblotting.

(D) Hydroxylation reaction of PLOD2 was carried out in vitro as described in Methods. Collagen peptides or no peptide substrates were used as controls for reaction. Data are means ± s.d. from three technical replicates for one biological replicate (*p < 0.05, Student’s t-test).

(E) Expression of integrin \( b1 \) mutants replacing Lys\#654 to Ala (K654A), Lys\#657 to Ala (K657A), Lys\#658 to Ala (K658A), or the triple Ala-substituted mutant replacing the lysine (3KA).

(F) Intracellular localization of the integrin \( b1 \) of these three mutants on HSC-2. Arrowhead indicated integrin \( b1 \) at the plasma membrane. Only in the transfected with triple Ala mutant did integrin \( b1 \) lack its membranous localization. Scale bar = 20 \( \mu \)m.

Further immunoprecipitation experiments of endogenous PLOD2 in HSC-2 cells revealed an interaction with the immature integrin \( b1 \) (Figure S6D).

Direct Hydroxylation of Integrin \( b1 \) by PLOD2 for Its Activation

Because the result of the specific binding between PLOD2 and integrin \( b1 \) would indicate that integrin \( b1 \) might be a substrate for PLOD2, hydroxylation of the integrin \( b1 \) protein purified from the lysate of the 293T transfected with wild-type (WT) PLOD2 was analyzed by mass spectrometry (LC/MS) using the integrin \( b1 \) derived from \( \Delta \)PKHD-PLOD2 transfected cells. The main spike, indicated with the hatched box in the over-viewed LC/MS chromatogram of integrin \( b1 \) from the cotransfected cells with integrin \( b1 \) and WT-PLOD2 (Figure 4A), contained the unique spike (indicated by arrow) representing the molecular weight of triple lysine hydroxylation (encoded amino acids between 651 a.a. and 658 a.a., “AFNKGEKK”), which was not observed in the integrin \( b1 \) protein from the integrin \( b1 \)-\( \Delta \)PKHD-PLOD2 cotransfected sample (Figure 4B, hyperview of 4A). The shift of the integrin \( b1 \) peak indicated by the arrow in the WT-PLOD2 cotransfected sample is two-fold higher than that of the integrin \( b1 \) in the \( \Delta \)PKHD-PLOD2 cotransfected sample (Figure 4B). Additionally, we performed an experiment monitoring the activity of PLOD2 employing substrate peptides according to the previous reports (Takaluoma et al., 2007; Guo et al., 2017). The result showed specific increase of succinate derived from \( z \)-ketoglutarate and substrate (integrin \( b1 \) substrate peptide; AFNKGEKK) incubated with purified PLOD2 as well as positive control sample using collagen peptide (IKGIGIKIG) sample (Figures 4C and 4D). The stability of integrin \( b1 \) harboring the single Lys-replacement with Ala (K564A, K657A, and K658A) did not seem significantly impaired inside cells transfected with each integrin \( b1 \) mutant; however, triple Lys-mutated integrin \( b1 \) (K564A + K657A + K658A) seemed markedly unstable (Figure 4E). Moreover, the triple Lys-mutative integrin \( b1 \) was not recruited to the plasma membrane in the transfected HSC-2 cells, in contrast to the cells with the single Lys-mutative integrin \( b1 \), which still retained its membranous localization as well as in those with WT-integrin \( b1 \) (Figure 4F). In this regard, flow cytometry analysis revealed that approximately 20% of the WT-transfected BHK cells showed plasma membrane expression of integrin \( b1 \), whereas no significant number of cells showed membrane expression of integrin \( b1 \) in the triple K-A mutant-transfected cells, which indicated loss of membrane localization without PLOD2 activity (Figures S7A and S7B). Thus, the magnitude of hydroxylation on integrin \( b1 \) by PLOD2 might be critical for its intracellular stability.

PLOD2 Facilitates Metastasis of SCC Cells In Vivo

Based on these findings, we examined whether PLOD2-affected invasion/metastasis of SCC cells in vivo using the intrathoracic metastatic mouse model xenografted with HSC-2 cells. Stably GFP-expressing HSC-2 cells from the PLOD2-knockout clone (PLOD2-KO; KO#31, Figure S8A) were xenografted into the thoracic cavity of athymic nude (nu/nu) mice and evaluated for development of metastatic foci 40 days post-implant in comparison with mice xenografted with parental HSC-2 cells (PLOD2-WT). As to cellular behaviors, PLOD2-KO (KO#31) carrying the heterozygous knockout of PLOD2 did not show significant differences in cellular growth compared with parent HSC-2 (Figure S8B), but endogenous expression of integrin \( b1 \) was seriously attenuated (Figure S8C). In the KO#31 clone, plasma membrane localization of integrin \( b1 \) was also disturbed, which suggested impaired function of integrin \( b1 \) (Figure S8D). Reflecting these results, cellular motility of KO#31 was also seriously decreased in comparison with the parental HSC-2 cells (Figure S8E, image and graphs). Whereas, using PLOD2-KO clone (#31) of HSC-2, we performed rescue experiment by reintroducing wild-type PLOD2. The result showed the revertant (revPLOD2)
A

PLOD2-WT

PLOD2-KO (KO#31)

thoracic cavity

dorsal ventral lung
dorsal ventral lung

Bright field

EGFP

Merge

B

PLOD2-WT

PLOD2-KO (KO#31)

thoracic cavity

ventral

lung

ventral

lung

HE

PLOD2

Integrin β1

x10

x10

x40

x40
restored cellular motility, which was similar to the original HSC-2 by regaining expression of maturated integrin β1 (Figures S9A and S9B). This corroborated requirement of integrin β1 expression for invasive property of HSC-2 cells, and it was regulated by PLOD2. Further we performed the time-dependent chase experiment for integrin β1 under presence of PLOD2 on PLOD2-KO clone with or without WT-PLOD2 and ΔPKHD-PLOD2 transfection. Result showed significant induction of mature integrin β1 was detected in response to time-dependent increase of WT-PLOD2 (Figure S10). Whereas overexpression of ΔPKHD-PLOD2, deficient form of hydroxylation activity (Figure 4B), induced the instability of integrin β1, which suggested the necessity of hydroxylation for integrin β1 maturation. Consequently, in the tumor metastasis mouse model, implant of the PLOD2-KO (KO#31) HSC-2 clone showed drastic elimination of metastatic lesions inside the thoracic cavity in vivo except for local growth of lung tumors, which was clearly different from the PLOD2 WT cell-implanted mice with multiple metastatic foci on pleura (Figures 5A and S11). Immunohistochemistry of the tumors derived from the PLOD2-WT cell-xenograft and KO#31 cell-xenograft was corroborative with tumor cells with or without coexpression of PLOD2 and functional integrin β1 (Figure 5B).

In this study, we found an additional functional aspect of PLOD2 as a direct regulator of cancer invasion/metastasis through its specific interaction with a major integrin family molecule, integrin β1. Namely, PLOD2 is a key regulator for activation of integrin β1 expressed in head and neck SCCs, which stabilizes and activates integrin β1 by hydroxylation via specific binding. This event leads to accelerated cellular motility needed to form metastatic sites for the SCC cells in vivo. Based on results of our analysis, tumor invasion via the PLOD2-mediated activation of integrin β1 seems not to be mediated by the EMT (epithelial-mesenchymal transition) process in cancer cells but by the direct effect of interactions of these two molecules. Marked increase in cellular motility of SCC cells was observed by integrin β1-PLOD2 protein-protein interaction without altering CDH1 or SNAIL expression (Friedl, 2004; Friedl et al., 2004; Canel et al., 2013) as shown in Figure 2C, and the increase appeared strongly affected by catalytic activity of PLOD2 (Figures 2G, 2H, and 4D). Importantly, instability of integrin β1 protein from knockdown of PLOD2 (downregulation of PLOD2) was observed in cancer cells of diverse origins such as esophageal SCC cells (KYSE30), lung adenocarcinoma cells (A549), uterine cervical SCC cells (HeLa), hepatocellular carcinoma cells (KYN-2), and breast...
invasive ductal adenocarcinoma cells (MCF7) as well (Figure S13A). In addition, we examined change of integrin β6 and integrin α5 expression by treating with PLOD2-siRNA in HSC-2, HSC-3, and Ca9-22 cells, because integrin β6 and α5 were reported to be expressed, in addition to β1 on head/neck SCCs (Koivistro et al., 2000). The result was shown as Figure S13B, which demonstrated significant attenuation of integrin β6 and α5 by PLOD2-siRNA treatment in these cells. As an exception, in HSC-3 cells, unlike other two SCC lines, decrease of α5 was not clearly observed, whereas integrin β6 was decreased as well as β1. This might be because of the HSC-3’s different cellular property from HSC-2 and Ca9-22 (both were cloned from primary SCC lesion), which was cloned from the lymph nodal metastasis and with highly expressed Vimentin (Figure 2C). These observations imply that PLOD2 might play a pivotal role for regulation of integrin β1 involved in invasion/metastasis of tumors with diverse lineages. Additionally, integrin β1 knockdown had less of an effect on migration in Figure S4B, than siPLOD2 throughout. PLOD2 may in fact provide an approach to target integrin β1 along with some other invasion-promoting targets. A large screening for PLOD2 targets may be required in the future.

We demonstrated that PLOD2 hydroxylase modified integrin β1 protein as a direct substrate. In the intracellular molecular binding assay (Fluoppi assay), PLOD2 and integrin β1 formed complex at the ER (ER) as shown in Figures 3B, 3C, and S6, then the integrin β1 moved to filopodia and the plasma membrane (Figure 3D). ER localization of PLOD2 has been reported in the previous studies (Chen et al., 2017; Gjaltema et al., 2016), which is consistent with our present result, indicating that PLOD2 first hydroxylizes integrin β1 at the ER, following which the integrin β1 is recruited to the cell surface functional site after the modification.

In previous studies on modification of collagen by PLOD2, hydroxylation of lysine within the X-K-G motif was reported to be essential for polymerization of collagen fibers (Kovnikko and Phlajaniemi, 1998; Mlynyharju and Kovnikko, 2004; Takaluoma et al., 2007; van der Slot et al., 2003). Based on earlier findings, the putative lysine-rich motif for hydroxylation was found at position of #651-658 a.a., “AFNKGEKK,” in integrin β1 (Figure S13C), which may point to a target motif for PLOD2. In fact, the mass spectrometry result revealed “AFNKGEKK” as the supershifted peak among integrin β1 fragments, which appeared only in the active PLOD2-treated sample (Figures 4A and 4B), and coincidently, the triple Lys substitution with Ala (K654A/K657A/K658A) abolished membrane localization of integrin β1 (Figures 4F and S7). Taking these together, the hydroxylation of these lysine within AFNKGEKK sequence by PLOD2 is critically implicated in gain of integrin β1 function in vivo.

We generated the PLOD2-knockout HSC-2 clone, PLOD2-KO (KO#31; heterozygous genomic knockout of PLOD2, PLOD2−/−), and tracked its dynamics in vivo in comparison with those of the parental HSC-2 (wild type; PLOD2-WT) using the mouse xenografted model. Intrathoracic implant of PLOD2-KO cells showed loss of pleural dissemination and pulmonary metastasis; it only showed local growth of tumor, whereas PLOD2-WT cells developed multiple metastatic foci covering the entire thoracic cavity. These data suggested that cancer cell motility was critically regulated by the presence of PLOD2. Immunohistochemistry demonstrated coexpression of PLOD2 and integrin β1 at invasive cancer nests in the metastatic tumor tissue derived from the mice implanted with PLOD2-WT cells, whereas integrin β1 expression in cancer nests was markedly attenuated by defect in PLOD2. This histological finding corroborated the significant relationship between integrin β1 and PLOD2 in vivo as was proven, further, by in vitro molecular assay. Finally, we confirmed that the expression pattern of integrin β1 and PLOD2 in SCC tissues surgically obtained from patients was similar to that of tumors in the mouse model.

Therefore, we conclude that integrin β1, a critical motor molecule for SCC invasion/metastasis, requires PLOD2 for its stabilization and functional cellular localization via their specific interaction through hydroxylation (Figure 7). Our results indicate that specific suppression of PLOD2 activity may provide one of the chief clues for inhibiting cancer invasion/metastasis in future cancer therapeutics.

Limitations of the Study

This study provides a crosstalk between PLOD2 and integrin β1 in cellular motility, and the hydroxylation on integrin β1 by PLOD2 is critical for its intracellular stability. However, we could not optimize the in vitro

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**Figure 6. Expression of PLOD2 and Integrin β1 in Head and Neck SCCs from the Patients**

Immunohistochemistry using anti-PLOD2 Ab and anti-integrin β1 Ab were performed on the SCC tissues derived from oral cavity, pharynx, and larynx, respectively. Box at the HE images (left column) indicated the field shown as hyperview in the middle and right column. Scale bar = 50 μm.
reaction of integrin β1 hydroxylation, because of an insoluble integrin β1 mutant substrate, and we were unable to directly detect the function of PLOD2 and integrin β1 in patient-derived primary SCC samples. Further enzymatically and clinically studies remain to be determined.

METHODS
All methods can be found in the accompanying Transparent Methods supplemental file.

SUPPLEMENTAL INFORMATION
Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.100850.

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AUTHOR CONTRIBUTIONS
Conceptualization, K.S., E.K.; Methodology, Y.U., K.S.; Investigation, Y.U., K.S., H.I., I.S., Y.K., M.S.; Writing—Original Draft, K.S., E.K.; Writing—Review & Editing, E.K.; Funding Acquisition, Y.U., K.S., E.K.; Resources, Y.U., K.S., A.H.; Supervision, E.K.

DECLARATION OF INTERESTS
The authors declare no competing interests.
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Supplemental Information

PLOD2 Is Essential to Functional Activation of Integrin β1 for Invasion/Metastasis in Head and Neck Squamous Cell Carcinomas

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Figure S1. Cell growth and migration by the suppression of PLODs, Related to Figure 1.
(A) mRNA expression of the protein hydroxylases was semi-quantitatively examined in oral SCC lines (HSC-2, HSC-3 and Ca9-22) and non-neoplastic keratinocytes (HaCaT) by RT-PCR. (B) Effects of two different oligos for PLOD2 knockdown were confirmed by immunoblot and wound healing assay. #1: custom siRNA synthesis based on previously reports (Sada et al., 2016). #2: on-target plus siRNA (set of 4 oligos). (C) Oral SCC cell growth was measured by MTT assay 48 hours after transfection of siRNA. Data are means±s.d. from 3 biological replicates. (D) Distribution of integrin β1 (red) by the knockdown of PLOD1 and PLOD3 gene. Cytoskeleton (green) and nuclei (blue) were stained with phalloidin and Hoechst, respectively. Scale bar = 20 μm
(A) Representative images of plasma membrane localization of CDH1 (red). Plasma membrane was marked with GFP (green) using a CellLight membrane-GFP. Membrane localization of CDH1 merged with GFP were quantified by fluorescence intensity (right panel). Quantitative results are mean ± s.d. from 3 biological replicates with more 10 cells per replicates (n.s. = not significant, Student’s t-test). Non-targeting siRNA (siCtrl) and siPLOD2 are shown by gray and white box, respectively. Scale bar = 20 µm

(B) PLOD2-knockdown cells were stained for integrin β1, and filopodial protrusions (magenta) were detected by FiloQuant (Jacquement et al., 2017) as a plugin for the ImageJ. Protrusion number (right panel) was quantified by FiloQuant analysis. Results are mean ± s.d. from 3 biological replicates with 20 - 50 cells per replicates (*p < 0.05, Student’s t-test). Scale bar = 20 µm

(C) Representative images of plasma membrane localization of integrin β1 (red). Plasma membrane was marked with GFP (green) using a CellLight membrane-GFP. Fluorescence intensity of integrin β1 localized membrane was calculated and the results are mean ± s.d. from 3 biological replicates with more 10 cells per replicates (*p < 0.05, Student’s t-test). Scale bar = 20 µm

Figure S2. Membrane localization of CDH1 and integrin β1 in the PLOD2-knockdown oral cancer cells, Related to Figure 2.
Figure S3. Intracellular localization of integrin β1, Related to Figure 2.
Magnification of image in Figure 2B. Scale bar = 10 µm.
**Figure S4. Expression of integrin β1 in the PLOD2-knockdown oral cancer cells, Related to Figure 2.**

(A) The intensity of immunoblot band (Fig. 2C) was quantified by densitometric analysis. Band intensity was normalized to beta-Actin band intensity. Data are means±s.d. from 3 biological replicates (*p < 0.05, Student’s t-test).

(B) Cell migration was detected by wound healing assay after transfection of integrin β1 siRNA (siβ1). Images were taken 0 and 24 h after wound formation (scale bar = 400 μm), and the wound width was estimated using fluorescence microscopic images. Data are means±s.d. from 3 technical replicates for 1 biological replicate (*p < 0.05, Student’s t-test as compared to siCtrl).

(C) Expression of integrin β1 in the siβ1-transfected cells by immunoblotting using anti-integrin β1. Band intensity was normalized to beta-Actin band intensity. Data are means±s.d. from 3 technical replicates for 1 biological replicate (*p < 0.05, Student’s t-test).
Figure S5. Distribution of integrin β1 by PLOD2 expression, Related to Figure 2.

(A) The intensity of immunoblot band (Fig. 2F) was quantified by densitometric analysis. Band intensity was normalized to beta-Actin band intensity. Data are means±s.d. from 3 biological replicates (*p < 0.05, Student’s t-test).

(B) Representative images of plasma membrane localization of integrin β1. Distribution shift of integrin β1 were quantified by fluorescence intensity of intracellular and plasma membrane-localized β1. Quantitative results are mean±s.d. from 3 biological replicates with more 10 cells per replicates (*p < 0.05, Student’s t-test). Scale bar = 20 μm

(C) The percentages of lysosome localization of integrin β1 were calculated by immunofluorescence image of β1 overlap with lyso-GFP vs total β1. Pearson’s correlation coefficient was used to measure correlation between integrin β1 and lysosome. Quantitative results are mean±s.d. from 3 biological replicates with more 10 cells per replicates (n.s. = not significant, Student’s t-test).

(D) HSC-2 cells were transfected with the expression vector of ΔPKHD-PLOD2 cloning to pEF1a-IRES-ZsGreen. After 24h, the cells were treated with MG132. The expression levels and subcellular localization of integrin β1 were observed by immunoblot and immunocytochemical analysis. Scale bar = 20 μm.
Figure S6. The interaction of integrin β1 with PLOD2 in endoplasmic reticulum, Related to Figure 3.
(A) The protein-protein interaction between PLOD2 and integrin β1 was detected using the fluorescent-based Fluoppi system. Representative images of intracellular foci were observed at high-resolution by the fluorescence microscopy. Scale bar = 10 μm (B) The fluorescence intensity per unit area of foci was measured. Data are obtained as the averages of nine spots from three independent biological replicates. Error bar represent s.d. (*p < 0.05, Student’s t-test). (C) Marked aggregation of fluorescent dots was detected only in the PLOD2-knockout HSC-2 cells cotransfected with integrin β1-Red and PLOD2-Ash. Nuclei (blue) were stained with Hoechst. Results represent the three images from three independent biological replicates. Scale bar = 20 μm (D) Lysate (Input) from HSC-2 cells was immunoprecipitated (IP) with control IgG or anti-PLOD2 antibody. Endogenous integrin β1 was coprecipitated with endogenous PLOD2 in HSC-2 cells.
Figure S7. PLOD2 activity and plasma membrane translocation of integrin β1, Related to Figure 4.
(A) BHK (baby hamster kidney) cells were transfected with empty control vector, WT-integrin β1 (WT) and mutant integrin β1 (triple KA). Expression of integrin β1 was confirmed by immunofluorescence staining without permeabilization. Scale bar = 20 μm. (B) Flow-cytometry analysis of integrin β1 expression in transfected BHK cells without permeabilization. Cells were gated on the basis of their SSC and FSC (upper panels). In dot plot (middle panels), the boxed area shows the positive populations of cell surface. Histogram and mean fluorescence intensity (MFI) shown in lower panels.
Figure S8. Analysis of PLOD2 KO cells, Related to Figure 5.

(A) Sequence of Exon 7 in PLOD2 gene targeting by CRISPR-Cas9 system. PLOD2 knockout cells (KO #2, #31) were isolated as heterozygous deletion and frameshift mutation. gRNA (guide RNA) for using CRISPR-Cas9 system are shown. (B) Cell growth of PLOD2-KO HSC-2 cell was measured by MTT assay. Results are mean ± s.d. from 3 biological replicates. (C) Expression of PLOD2 and integrin β1 in the PLOD2-knockout cells were examined by immunoblotting. (D) Distribution of integrin β1 (red) and E-cadherin (CDH1: green) by the knockout of PLOD2 gene. Scale bar = 20 μm (E) Cell migration of PLOD2-knockout cell lines was detected by wound healing assay using 24-well plate coated with collagen. Images were taken 0, 24 and 48 h after wound formation (scale bar = 400 μm), and the wound width was estimated using microscopic images. Results are mean ± s.d. from 3 technical replicates for 1 biological replicate (*p < 0.05, Student’s t-test as compared to WT).
Figure S9. Integrin β1 maturation by re-expression of PLOD2 in PLOD2-knockout cells, Related to Figure 5.

(A) After stable transfection of empty control vector or WT-PLOD2 to PLOD2-knockout cells (KO #31), the cells were selected with G418 for 4 weeks. Stable cell lines expressing PLOD2 and ZsGreen were generated. These revertant cell lines (revCtrl clone #1, #2 and revPLOD2 clone #3, #5) were seeded and cultured in 24-well plates for 48 hours. Re-expression of PLOD2 and integrin β1 were detected by immunoblotting. The intensity of immunoblot band (right panel) was quantified by densitometric analysis. Band intensity was normalized to beta-Actin band intensity. Data are means±s.d. from 3 biological replicates (*p < 0.05, Student’s t-test). (B) Migration rate of revertant cells was measured by wound healing assay. Images were taken 0, 24 and 48 hours after wound formation (scale bar = 400 μm), and the wound width was estimated using fluorescence microscopic images. Data are means±s.d. from 3 technical replicates for 1 biological replicate (*p < 0.05, Student’s t-test between revCtrl and revPLOD2).
**Figure S10.** Integrin β1 maturation in a PLOD2-dependent manner, Related to Figure 5.

(A) PLOD2-knockout cells (KO #31) were transiently transfected with empty control vector (upper panel), WT-PLOD2 (middle panel) and ΔPKHD-PLOD2 (lower panel). Cell lysate was collected at each time points after transfection, and the expression of integrin β1 and PLOD2 were detected by immunoblotting.

(B) The intensity of immunoblot band was quantified by densitometric analysis. Band intensity was normalized to beta-Actin band intensity. Each band intensity of empty vector, WT-PLOD2 and ΔPKHD-PLOD2 overexpression are shown by blue, red and green box, respectively. Data are means±s.d. from 3 biological replicates (*p < 0.05, Student’s t-test as compared to empty vector).
Figure S11. Fluorescence imaging of PLOD2 KO cells in vivo, Related to Figure 5.
Fluorescence imaging of the invasive and metastatic tumors in the thoracic cavity of mice injecting PLOD2-WT and PLOD2-KO HSC-2 cells are shown. Scale bar = 50 μm
Figure S12. Colocalization of PLOD2 and integrin β1 in the SCC tissues, Related to Figure 6

(A) Immunohistochemistry using anti-PLOD2 Ab and anti-integrin β1 Ab were performed on the SCC tissues derived from larynx and oral cavity. Box at the HE images (left panel) indicated the field shown as hyperview in the middle and lower panel. Scale bar = 1 mm  (B) Subcellular colocalization (lower panels, arrowhead) of PLOD2 (green) and integrin β1 (red) in each case of oral SCC was assessed by fluorescent immunostaining using anti-PLOD2 and anti-integrin β1 antibodies. Nuclear counterstain with Hoechst 33258 was applied. Results represent the images from five independent area for the three cases. Scale bar = 20 μm
Figure S13. Expression of integrin β1 in PLOD2-knockdowned cancer cells, Related to Figure 4.

(A) Expression levels of integrin β1 in PLOD2 siRNA-transfected cancer cell lines were detected by immunoblotting. KYSE30 (esophageal ca.), A549 (lung ca.), HeLa (cervical ca.), KYN2 (liver ca.) and MCF7 (breast ca.) were used. (B) Expression of integrin β1, β6 and α5 in the PLOD2-knockdowned cells were examined by immunoblotting. (C) Amino acid sequence of integrin β1. Hydroxylation site (box), transmembrane domain (dashed line) and cytoplasmic domain (under line) are shown.
Table S1. Oligonucleotides, Related to Figures 1 – 4.

| Gene name | Forward primer (5’ to 3’) | Reverse primer (5’ to 3’) |
|-----------|---------------------------|---------------------------|
| ALKBH3    | TGGCTTTGTTTGACGTGAAAG     | GTGCCAGTGAGGATTTGGTT      |
| ASPH      | GGAGATTTTATTTCACCTGGG     | CTTTGCGTTTATCCATCAGTC     |
| EGLN1     | GCCCCATTTGCGACATTGAAC     | CCCCACAACCTTTCTACCTGGTTAG|
| EGLN2     | AGCCCCAATGCAAGCTTCTC      | AGTGTTAGAGGTGGCTGGT       |
| EGLN3     | ATCAGACAGGCTGTCCTTCCTA    | CTTGGCACTCCAATCCTGGT      |
| FIH1      | GTGGGCCAGGAAGATTGCA      | GGGTGTAAGACGAGTATG        |
| FTO       | GAGAGCGCGAAGCTAAGAAA     | AATGAGGATGCGAGATACC       |
| ITGB1     | AATGAAAGGGGCTGTTGGTAG    | CGTTGCTGGTCACCAAGTA       |
| JMJD2A    | CCTCAGTGCCTGTCTGATTAGCT  | CCAGTGAAGAGGACATTCG       |
| JMJD6     | GCCACAAACTACTAGAGAGC     | TTGGAACCCTGTAGCTCCTCC     |
| MINA53    | TACTTGGCTCCTTGGTGG       | GGAGATGGCTGCTGAGAT        |
| NO66      | TGTCTTACGGGTATTCTCAATTCC| TCCCCACCTCTCAACACCTC      |
| OGFOD1    | TTTTGAAGATTTCCGGTCTGG    | GACCATGAAACGACCACTT       |
| PLOD1     | GCAGCAGGATGGTCTGTCTGA    | GGGCTTTGCTGTAGCTGGT       |
| PLOD2     | GCGTTCTCTTCCGGTCCTC      | GTGTTAGCTGGTCAGGATG       |
| PLOD3     | GGTACGAGGACCAGTGCTCTTA  | GAAGGTTGGAAGGTGTTG        |
| TET1      | GGGCAGCAGTTTCACTACAGAGGC| GGGGTTCGGTTTCACTTTTT      |
| beta-Actin| CCTCAGTGAAGATTTGCTCAGAGG| TGCCAATGTTGATGACTGG       |

Primer sets of Cloning and plasmid construction

| Primer set | Oligonucleotides | Primer set | Oligonucleotides |
|------------|------------------|------------|------------------|
| PLOD2-WT   | TTACTAGTGCCATG   | Same forward primer with PLOD2-WT | TTGCGGCCGCTCAGAGGTCTTCTTCTGAGAT |
| for pEF1-IRES-ZsGreen | GGGATGGGCTGCTAGAGTCTTCTTCTGAGAT | for PLOD2-WT | TTGCGGCCGCTCAGAGGTCTTCTTCTGAGAT |
| PLOD2-ΔPKHD for pEF1-IRES-ZsGreen | GGGATGGGCTGCTAGAGTCTTCTTCTGAGAT | Same forward primer with PLOD2-WT | TTGCGGCCGCTCAGAGGTCTTCTTCTGAGAT |
| PLOD2-ΔPKHD for pcdnA3.1(+) | GGGATGGGCTGCTAGAGTCTTCTTCTGAGAT | Same forward primer with PLOD2-WT | TTGCGGCCGCTCAGAGGTCTTCTTCTGAGAT |
| PLOD2-Red for pMonti-Red-MNL | GGGGAAGCTTACCATTG | GGGGGAAGCTTACCATTG | TGTGAAG |
| Integrin β1-WT for pcdnA3.1(+) | GGGGAAGCTTACCATTG | TGTGAAG | GGGGGAAGCTTACCATTG |
| Integrin β1-WT for pcdnA3.1(+) | GGGGAAGCTTACCATTG | TGTGAAG | GGGGGAAGCTTACCATTG |
| Integrin β1-WT for pcdnA3.1(+) | GGGGAAGCTTACCATTG | TGTGAAG | GGGGGAAGCTTACCATTG |
| EGFP for pcdnA3.1(+) | TTTCTCAGGAGGTGAGCA | TTTCTCAGGAGGTGAGCA | TTTCTCAGGAGGTGAGCA |

Annealing oligonucleotides

| Primer set | Oligonucleotides | Primer set | Oligonucleotides |
|------------|------------------|------------|------------------|
| Myc-tag for PLOD2-WT & ΔPKHD | GGGGGAATCCGGC | GATCATCAGAGGGTTCTTCTTCTGAGATG | GATCATCAGAGGGTTCTTCTTCTGAGATG |
| FLAG-tag for Integrin β1-WT & mutants | GGGGGAATCCGGC | GATCATCAGAGGGTTCTTCTTCTGAGATG | GATCATCAGAGGGTTCTTCTTCTGAGATG |
| Myc-tag for EGFP | CAGGCTCCTCTTCTTCTGAGATG | TTTCTCAGGAGGTGAGCA | TTTCTCAGGAGGTGAGCA |

Table S1. Ueki et al
TRANSPARENT METHODS

Cell lines

Human cancer cell lines used in the present study were as follows: HSC-2, HSC-3, derived from the lymph nodes of different patients with metastatic poorly differentiated SCC, and Ca9-22 derived from the patients with gingival SCC (JCRB Cell Bank: Japanese Collection of Research Bioresources Cell Bank) (Yamamoto et al., 1986; Momose et al., 1989; Matsui et al., 1998). Cells were maintained in RPMI 1640 containing 10% FBS with 100 unit/mL penicillin and 100 mg/mL streptomycin (Life Technologies) at 37°C with 5% CO₂. These cell lines were passaged for less than 6 months and then replaced with those of early passages.

PLOD2 KO cell lines

PLOD2 knockout cell clones were generated by CRISPR/Cas-9 based genome engineering technology. The plasmids hCas9 and gRNA_Cloning Vector were a gift from George Church (Addgene plasmid # 41815 and #41824) (Mali et al., 2013). gRNA vector including PLOD2 targeting sequence (5’- GCTGATCATCATCATTATCC -3’) was prepared by following Church lab protocol (https://media.addgene.org/41824/). hCas9 and PLOD2 gRNA vector were co-transfected into cells using Lipofectamine
3000 Transfection Reagent (Thermo Fisher Scientific) by following manufacturer’s protocol. Single cell cloning was achieved by serial dilution, and genotype of each clones was analyzed by genomic PCR followed by Sanger DNA sequencing. Following primers were used to amplify genomic sequence including PLOD2 targeting site, forward primer: 5’-TCCCTTAGGAGCCTTGGAAAACA and reverse primer: 5’-ATCCAGCCAGGTGACATAAACCA. Sanger DNA sequencing was carried out using the forward primer and BigDye terminator v3.1 cycle sequencing kit (Applied Biosystems).

For revertant cell lines re-expressing PLOD2 in PLOD2-knockout cells, the PLOD2-IRES-ZsGreen plasmid was transfected into PLOD2-knockout cells. After selection with 0.1 mg/ml of Geneticin (G418, Gibco) for 4 weeks, PLOD2 re-expressing stable lines were newly cloned.

**Animal models**

A total of 1.0×10^6 cells of human oral squamous cell carcinoma cell lines stably expressing Green fluorescent protein (GFP-HSC-2) were transplanted into the pleural cavity of a 7-week-old female nude mice (BALB/c nu/nu, CLEA Japan Inc.) by direct injection. 40 days after initial administration of the GFP-HSC-2 cells, all mice were
autopsied and their organs examined by fluorescence stereomicroscopy (Leica M165 FC).

The animal studies in the present work were approved by the Animal Research Committee of the Niigata University School of Medicine. All mouse procedures and euthanasia, including cell transplantations, were performed painlessly or under anesthesia, and within the strict guidelines of the Experimental Animal Center of Niigata University School of Medicine.

**Plasmid construction and transfection**

cDNA were prepared from HSC-2 cells using an RNeasy kit (Qiagen) and a SuperScript III kit (Invitrogen). Human *PLOD2* CDS (NM_000935.2) and *Integrin β1* CDS (NM_002211.4) were amplified from HSC-2 cDNA and NHDF cDNA, respectively. They were cloned into pBluescript SK\(^+\) vector (Stratagene). To generate the PLOD2 wild type and mutant plasmid, *PLOD2* genes were subcloned into a pEF1-IRES-ZsGreen vector at the *Nhe I* / *Spe I* compatible cohesive sites and *Sma I* site. The synthetic myc-tag oligonucleotides were annealed and inserted into a pEF1-IRES-ZsGreen (Clontech) vector at the *Sma I* and *BamH I* site. Furthermore, *PLOD2-myc* gene were subcloned into a pcDNA3.1(+) at the *Nhe I* / *Spe I* compatible
cohesive sites and Not I site. For integrin β1-FLAG plasmid, integrin βI gene were subcloned into a pcDNA 3.1(+) vector (Thermo Fisher Scientific) at the BamH I - Xho I. The synthetic flag-tag oligonucleotides were annealed and inserted into a pcDNA 3.1(+) vector at the Xho I and Apa I site. For Myc-EGFP plasmid, EGFP gene were amplified from the pEGFP-N1 vector (Clontech) and subcloned into a pcDNA 3.1(+) at the Xho I - Apa I site. The synthetic myc-tag oligonucleotides were annealed and inserted into the Nhe I-Xho I sites.

PLOD2-Red plasmid was constructed by cloning the cDNA encoding PLOD2 into the Hind III - Not I of the vector pMonti-Red-MNL (MBL). Integrin β1-Ash plasmid was constructed by cloning the cDNA encoding integrin βI into the BamH I - Not I of the vector pAsh-NML Ver.2 (MBL).

The site-directed mutagenesis for integrin β1 was performed by PCR using a single mutant primer and PfuUltra High-Fidelity DNA polymerase (Agilent Technologies). After Dpn I digestion the PCR products were transformed into the competent E.coli cells. The nucleotide sequences of all constructs obtained by PCR were confirmed by DNA sequencing. The primer sets used in this study are shown in Table S1.

For the plasmid transfection, Cells were transiently transfected with PLOD2 and integrin βI plasmids using Lipofectamine 3000 (Thermo Fisher Scientific) according to
the manufacturer’s instructions. The stable clone expressing EGFP was selected in 0.6 mg/ml of Geneticin (Gibco).

**Cell proliferation assay**

Cells (4×10⁴) were seeded on a 48-well plate. After 48 hours of siRNA transfection, cell proliferation was evaluated by MTT assay using Cell Count Reagent SF (Nacalai Tesque) according to the manufacturer’s protocol.

**Wound healing assay**

The myc-EGFP plasmid was transfected into HSC-2, HSC-3 and Ca9-22 cells. After selection with 600 μg/ml of Geneticin for 4 weeks, GFP-expressing stable lines were newly cloned and used in wound healing assay. Cells were seeded in collagen type-1 coated 24-well plates. 48 hours after siRNA treatment was made a scratch using a cell scratcher was done (Iwaki, Asahi Technoglass). The wounded monolayer was washed with PBS, exchanged medium, and photographed using a microscope every time. All data were obtained from at least three independent experiments.

**Conventional and quantitative PCR**
Total RNA was extracted from prepared cells using RNeasy plus mini kit (QIAGEN) according to the manufacturer’s protocol. Complementary DNA was synthesized from 1 µg of total RNA using Revertra Ace (Toyobo) for performing mRNA analysis. Each cDNA was amplified by PCR with z-Taq polymerase (TaKaRa). Quantitative real-time PCR was performed using SYBR Green RT-PCR Kit (Applied Biosystems). Data analysis was followed by \( \Delta \Delta^{{CT}} \) methods using the step one plus instruction (Applied Biosystems). The primer sets used in this study are described in Supplementary Table S1.

**RNA interference**

Each siRNA (25 nM) was transfected using Lipofectamine RNAi MAX reagent (Thermo Fisher Scientific) according to the manufacturer’s protocol. ON-TARGET plus SMART pool siRNA for human PLOD2 (#5352), PLOD1 (#5351) and PLOD3 (#8985) were purchased from Dharmaco. The custom siRNA of PLOD2 (Sada et al., 2016) and integrin β1 (Kato et al., 2014) was synthesized by Sigma-Aldrich. Negative control scrambled siRNA was purchased from Sigma-Aldrich.

**Immunoblotting and immunoprecipitation**
The cell pellets were lysed with the RIPA buffer (50 mM Tris/HCl pH 7.5, 150 mM NaCl, 0.5% sodium deoxycholate, 1% NP-40, 0.1% SDS) containing the complete mini protease inhibitor (Roche). Samples were separated on 10% SDS-PAGE gels followed by transfer to PVDF membrane. Nonspecific binding was blocked by incubation in 5% skim milk in TBS-T. Blots were probed with specific primary antibodies followed by appropriate horseradish peroxidase-conjugated secondary antibody. Positive signals were detected by a chemiluminescence system (Supersignal West Pico or Femto substrate; Thermo Fisher Scientific) and quantification of band intensity was calculated using a ChemiDoc Touch imager (Bio-Rad). Anti-PLOD1 (Sigma-Aldrich), anti-PLOD2 (Proteintech), anti-PLOD3 (Sigma-Aldrich), anti-Integrin β1 (CST) anti-E-cadherin (BD Biosciences), anti-SNAIL (Abcam), anti-ZO-1 (Abcam) anti-Vimentin (Santa Cruz Biotechnology) and anti-Actin (Merck) antibodies were used for immunoblotting.

For immunoprecipitation, HEK 293T cells were transiently co-transfected with the expressor plasmid of integrin β1-flag and PLOD2-myc. Cells were lysed in IP buffer (50 mM Tris-HCl pH7.5, 150 mM NaCl, 1% Brij 20, 0.5% n-octyl-β-D-glucoside, 0.5% Triton-X, 1 mM PMSF) supplemented with protease inhibitors cocktail (Roche). Cell lysates were incubated overnight at 4°C with anti-FLAG M2 affinity gel
Beads were washed six times with wash buffer (50 mM Tris-HCl pH7.5, 150 mM NaCl, 0.2% NP-40) and were boiled in 2× sample buffer (125 mM Tris-HCl pH 6.8, 4% SDS, 20% Glycerol, 0.01% Bromophenol blue, 10% β-Mercaptoethanol). Equal amounts of proteins were separated on 10% SDS-PAGE gels and immunoblotted with the anti-myc monoclonal antibody (9E10, Sigma-Aldrich) and horseradish peroxidase-conjugated secondary antibody (Mouse TrueBlot ULTRA: anti-mouse Ig HRP, Rockland).

**Fluoppi assay**

The protein-protein interaction between PLOD2 and integrin β1 was detected using the fluorescent-based technology detecting protein-protein interaction (Fluoppi) system (MBL life science), in which PPIs were visualized as foci (Koyano et al., 2014; Yamano et al., 2015). PLOD2 was subcloned into a pMonti-Red-MNL plasmid to fuse with a Monti-Red (homo tetramer red fluorescent protein from Montipora sp.) tag. Integrin β1 was subcloned into a pAsh-MNL ver.2 plasmid to fuse with an Ash (homo oligomerized protein assembly helper) tag. These plasmids of PLOD2-Red and integrin β1-Ash were co-transfected into HSC-2 or HeLa cells using Lipofectamine 3000 (Thermo Fisher Scientific) according to the manufacturer’s instructions, and intracellular foci were
observed by the Olympus IX71 fluorescence microscope and analyzed using cellSens standard software (Olympus).

**Mass spectrometric analysis of integrin β1**

Extracts of $5 \times 10^6$ cells expressing the integrin β1-flag and PLOD2-myc were prepared on ice for 30 minutes in IP buffer. Integrin β1-flag proteins were immunoprecipitated by using anti-FLAG M2 affinity gel and eluted by $3\times$ FLAG peptide. Immunoprecipitated integrin β1-Flag were separated SDS-PAGE, the gel was then stained with Coomassie Blue, and protein bands corresponding to the proteins of interest were excised and digested with LysC protease (Roche) according to standard protocol. The extracted peptides were desalted and concentrated using RP-C18 StageTip columns and the eluted peptides used for mass spectrometric analysis. All LC/MS-MS experiments were performed by using standard protocol (ACROSABLE Inc. Sendai, Japan). Peptides were analyzed using an amaZon ETD ion trap mass spectrometer (Bruker Daltonics). Spectral data were compared to the Swiss-Prot Homo sapiens database using Mascot (version 2.3.01; Matrix Science) to identify the protein. The search criteria were: trypsin enzyme, allowance of up to two missed cleavage peptides, mass tolerance ±0.5 Da, MS/MS tolerance ±0.5 Da, fixed modifications of cysteine
carbamido-methylation, variable modifications of lysine hydroxylation.

**Flow-cytometry**

Human integrin β1 WT and triple KA mutant plasmid was transfected into BHK (baby hamster kidney) cells. Cells were harvested by trypsinization and washed twice with PBS containing 2% FBS. The cells incubated with anti-integrin β1 antibody (Abcam) for 20 min at 4°C, and then washed twice with PBS containing 2% FBS. For detection of β1 integrin, cells were incubated with anti mouse-Alexa fluor 488 (Thermo Fisher Scientific) for 20 min at 4°C, and further washed twice PBS containing 2% FBS. Cells were detected on a BD FACS Calibur cell analyzer (BD Biosciences) and data were analyzed using FlowJo VX software.

**Purification and activity of PLOD2**

*PLOD2* cDNA were subcloned into the pCMVViR-TSC vector (Sakaguchi et al., 2014) and the PLOD2-myc-6×His expressor was transfected into HEK293T cells. Extracts of $3 \times 10^6$ cells expressing the PLOD2-myc-6×His were prepared on ice for 30 minutes in IP buffer. Cell lysate was load into a column of TALON metal resin (Takara) and washed twice with wash buffer (50 mM Tris-HCl pH7.5, 150 mM NaCl, 10 mM
imidazole), and further eluted with 150 mM imidazole. The eluted PLOD2 was dialysed with 50 mM HEPES pH7.4 and 150 mM NaCl, and then concentrated with the Amicon ultra-0.5 centrifugal filters (Millipore). Protein concentration was estimated with the BCA protein assay (Pierce), Nanodorp Spectrophotometer at 280 nm (Thermo Fisher Scientific) and CBB staining of SDS-PAGE.

Hydroxylation reaction of PLOD2 were carried out in 50 μL volume of reaction buffer based on previously reported method (Guo et al., 2017). Briefly, the purified PLOD2 (approximately 1.0 μg) was reacted with the reaction buffer (50 mM HEPES pH7.4, 50 μM FeSO4, 100 μM α-KG, 500 μM ascorbate, 0.125 mM peptide substrate) for 120 min at 37 °C. The collagen peptide substrate (IKGIKGIKG) and integrin β1 peptide substrate (AFNKGEKK) were synthesized at 95% purity by Scrum Inc (Tokyo Japan). 1 μL of the reaction solution was diluted in 25 μL of 50 mM HEPES pH7.4 buffer and measured succinate production in the reaction solution using Succinate-Glo assay following the manufacturer’s procedure (Promega). The luminescence was detected using a Centro LB 960 microplate luminometer (Berthold).

**Immunofluorescence**

Cultured cells were fixed with PBS containing 4% paraformaldehyde and washed
with 0.1% Triton X-100 in phosphate-buffered saline (-). PLOD2 was stained with rabbit anti-PLOD2 antibody (Proteintech), Integrin β1 was stained with mouse anti-integrin β1 antibody (Abcam) and E-cadherin was stained with mouse anti-E-cadherin antibody (BD Biosciences) as a primary antibody, and further treated with Cy3-labeled goat anti-rabbit (H+L) antibody (Jackson), as a secondary antibody. Cell nuclei were counterstained with 4’,6-diamidino-2-phenylindole (Hoechst 33258 dye; Molecular probes). CellLight ER-GFP BacMam 2.0 (Thermo Fisher Scientific) was used for ER marker. For the staining of actin filaments (F-actin), Alexa 488-conjugated phalloidin (Molecular Probes) was used. The fluorescent signals were detected by the Olympus IX71 fluorescence microscope and analyzed using cellSens standard software (Olympus). For quantification, the fluorescence intensity of plasma membrane-localized integrin β1 vs total integrin β1 was calculated using Metamorph software (Molecular devices). Colocalization analysis and calculation of filopodial protrusions were performed using a Coloc 2 and FiloQuant (Jacquement et al., 2017) as a plugin for the ImageJ software. CellLight Membrane-GFP BacMam 2.0 (Thermo Fisher Scientific) was used for plasma membrane marker.

**Immunohistochemistry and patient-derived tissues**
Human oral, pharyngeal and laryngeal carcinoma tissues were fixed in 10% buffered formalin solution. Paraffin-embedded sections were deparaffinized, and antigen retrieval was performed by autoclaving (Pascal S2800, DakoCytomation) in sodium citrate buffer. Then endogenous peroxide activity was quenched by incubation in 0.3% hydrogen peroxide (Kanto Chemical Co.) for 20 minutes at room temperature. Slides were incubated with a 1:250 dilution of anti-PLOD2 antibody (Proteintech) and anti-integrin β1 antibody (CST) in CanGet signal immunostain solution B (Toyobo) overnight at 4°C, and further incubated with anti-rabbit HRP antibody (CST). After the slides were washed with phosphate buffered saline (-), they were subjected to the DAB detection system (Nichirei).

The study using patient-derived tissues was approved by the ethics committees of Niigata University and Niigata Cancer Center Hospital. Informed consent for use of materials was also obtained from each patient.

Quantification and Statistical Analysis

Data were represented as mean ±standard deviation (s.d.). Comparisons between two groups were carried out using Student’s t test (MS Excel). A value of P < 0.05 was considered significant.