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Title

PUCHI regulates Very Long Chain Fatty Acid biosynthesis during lateral root and callus

formation

Short title: PUCHI regulates VLCFA biosynthesis

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Abstract

Lateral root organogenesis plays an essential role in elaborating plant root system architecture. In *Arabidopsis*, the AP2-family transcription factor PUCHI controls cell proliferation in lateral root primordia. To identify potential targets of PUCHI, we analyzed a time course transcriptomic dataset of lateral root formation. We report that multiple genes coding for very long chain fatty acid (VLCFA) biosynthesis enzymes are induced during lateral root development in a PUCHI-dependent manner. Significantly, several mutants perturbed in VLCFA biosynthesis show similar lateral root developmental defects as *puchi-1*. Moreover, *puchi-1* roots display the same disorganized callus formation phenotype as VLCFA biosynthesis deficient mutants when grown on auxin-rich callus inducing medium. Lipidomic profiling of *puchi-1* roots revealed reduced VLCFA content compared to WT. We conclude that PUCHI-regulated VLCFA biosynthesis is part of a pathway controlling cell proliferation during lateral root and callus formation.

139 words

lateral root formation | morphogenesis | VCLFA | callus formation | cell division

Significance

Lateral root organogenesis enables plant roots to branch and improve foraging for resources. Using a systems biology approach, we discovered expression of very long chain fatty acid (VLCFA) biosynthesis pathway genes are induced downstream of PUCHI, a transcription factor controlling cell division and morphogenesis during lateral root formation. Furthermore, we report that regulation of the VLCFA biosynthesis pathway by PUCHI is conserved for both lateral root development and callus formation, the first step in *in vitro* plant regeneration.

Author contributions

M.J.B., L.L., and S.G. designed research; T.D.C., J.L., T.G., Y.B., Q.D., V.V., F.T., U.V., S.D. performed research; T.D.C., J.L., T.G., Y.B., M.L., U.V., P.G., J.D.F., S.D., H.F., M.J.B., L.L., and S.G. analyzed data; and T.D.C., M.J.B., L.L. and S.G. wrote the paper.

Introduction

Plant root system architecture (RSA) is the three-dimensional configuration of a whole root system in its living environment (1). It is a major determinant of plant viability and crop yield, and a target for breeding to improve crop performance under various stresses (2, 3). Root branching is of particular importance because it largely determines the overall surface area of the root system and its spatial organization in the soil. The molecular mechanisms of root branching have been extensively studied in the model plant *Arabidopsis thaliana*. Lateral roots (LRs) originate from a small group of xylem-pole pericycle cells of the primary root that are primed by auxin to acquire founder cell identity (4). These founder cells undergo a succession of anticlinal and periclinal cell divisions that eventually result in the formation of a dome-shaped lateral root primordium (LRP; 5–8). The LRP emerges through overlaying root tissues to become a LR (9).

Lateral root development is an excellent experimental system to study *de novo* meristem formation (10). Moreover, recent studies have shown that lateral root formation shares common mechanisms with organ regeneration in tissue culture, especially the first step of callus formation (11–14). While many genes involved in lateral root development have been identified, little is known about the mechanisms that progressively organize the LRP into a new root meristem (15). LRP organization is not dependent on a stereotypical cell division pattern and therefore on cell lineage (6, 7). It is a dynamic process dependent on complex gene regulatory networks and on cell-cell interactions including hormonal and biomechanical signals (6, 9, 16). Interestingly, inference of the gene regulatory network involved in LR formation suggested an early patterning mechanism defining the central region and flanks of the LRP and identified genes involved in this process (17). One such gene encodes the AP2/EREBP-family transcription factor PUCHI which was previously shown to control cell proliferation during LRP formation (18). The *puchi-1* mutant LRP exhibits additional anticlinal and periclinal cell divisions from early stages and produces abnormally enlarged flank cells (18). However, little is known about the pathways that are regulated by PUCHI.

Very long chain fatty acids (VLCFAs) are fatty acids with 20 or more carbons synthesized in the endoplasmic reticulum from long chain-fatty acyl-CoA (16 or 18 carbons) by the fatty acid elongase complex (SI Appendix, Fig. S1). This complex catalyzes rounds of 2-carbon elongation in a 4-step mechanism, involving a 3-ketoacyl-CoA synthase (KCS), a 3-ketoacyl-CoA reductase (KCR), a 3-hydroxyacyl-CoA dehydratase (HACD), and a trans-2,3-enoyl-CoA reductase (ECR). Multiple KCS enzymes with various expression patterns have been described and their substrate affinity is thought to be responsible for the final VLCFA chain length (19-21). In contrast, only a limited number of genes that encode functional enzymes catalyzing each of the subsequent steps of the elongation cycle has been identified in Arabidopsis. KETOACYL REDUCTASE 1 (KCR1) encodes an Arabidopsis KCR enzyme (22) while PASTICCINO 2 (PAS2; 23) and PROTEIN TYROSINE PHOSPHATASE-like (PTPLA) encode two HACD enzymes (24), whilst the product of the ENOYL CO-A REDUCTASE/ECERIFERUM 10 (ECR/CER10) gene has ECR activity (25). These enzymes are physically linked together by PASTICCINO 1 (PAS1; 27). VLCFAs are components of various classes of membrane, storage and extracellular lipids (27). VLCFA-containing sphingolipids and glycerolipids provide unique properties to membranes promoting inter-leaflet interactions and lipid rafts formation. As such, VLCFAs influence membrane dynamics during cell division (28, 29), the distribution of membrane proteins such as auxin transporters (26, 30), and are involved in non cell-autonomous processes of growth regulation (31-33). In addition, VLCFA-containing lipids, or their derivatives, are thought to act as second messengers in response to various stimuli (34, 35). Finally, VLCFAs are components of hydrophobic extracellular material such as skin barriers in mammal (36) or suberin and cuticular waxes in plants (27).

In this study, we identified genes acting downstream of PUCHI using a time course LR transcriptomic dataset (37) combined with a gene regulatory network (GRN) inference algorithm (17). We found PUCHI-dependent expression of genes involved in the biosynthesis of VLCFAs during LR development. Consistently, VLCFA mutants display defects in LR development similar to *puchi-1*. VLCFAs were previously described to regulate the competence of pericycle cells to generate calli on an auxin-rich callus-inducing medium (CIM) (33). We show that PUCHI is also expressed in CIM-treated roots and regulates the expression of genes involved in VLCFA biosynthesis during CIM-induced callus formation. Accordingly, CIM-treated roots of the *puchi-1* mutant show reduced level of VLCFAs as compared to wild type and exhibit comparable phenotype to VLCFA biosynthesis mutants. Our results indicate that PUCHI-

regulated VLCFA biosynthesis is part of a pathway controlling cell proliferation during LR and callus formation.

Results

PUCHI regulates VLCFA biosynthesis genes during lateral root development

To gain insight into the function(s) of PUCHI during LR development, we took advantage of the time-course transcriptomic dataset profiling every stage of LRP organogenesis (37) to identify candidate genes whose expression is regulated by this transcription factor. We employed the TDCor algorithm (17) to search the LR dataset for genes exhibiting an expression profile highly similar to that of *PUCHI* (Pearson's correlation coefficient > 0.80) when shifted back in time by 3 hours. Our *in silico* analysis retrieved 217 potential target genes whose expression profiles are correlated with that of *PUCHI* (SI Appendix, Table S1). A Gene Ontology (GO) enrichment analysis using BiNGO (38) revealed that 71 GO biological processes were significantly overrepresented in this group of putative downstream genes (SI Appendix, Table S2). Among them, the "VLCFA biosynthesis" category stood out as one of the most strongly overrepresented biological processes (p-value = 0.006). In the LR dataset, PUCHI transcript abundance rapidly rises after LR induction and peaks at around 12 hours after gravistimulation, which corresponds to the time when the first round of anticlinal cell division is observed (Fig. 1A, black line). PUCHI transcript levels then gradually decrease over time. Expression profiles of genes encoding key enzymes for each step of the VLCFA elongation cycle displayed comparable dynamics, albeit shifted in time. These included KCS1, KCS2, KCS9, KCS17 and KCS20 genes, all encoding members of the KCS enzyme family catalyzing the first step of VLCFA elongation, as well as KCR1, PAS2 and ECR/CER10, which encode enzymes catalyzing the second, third, and fourth steps of VLCFA elongation, respectively, and PAS1 (Fig. 1A and SI Appendix, Fig. S2).

We validated this inference approach with a classical transcriptomic analysis of the *puchi-1* mutant complemented with a dexamethasone (DEX) inducible PUCHI-GLUCOCORTICOID RECEPTOR (GR) protein fusion expressed under the control of the *PUCHI* promoter. Treatment of *pPUCHI::PUCHI:GR/puchi-1* seedlings with DEX targets the recombinant transcription factor to the nucleus to regulate gene expression (39, 40) and restore the wild type (WT) LR phenotype (*SI Appendix*, Fig. S3). We used the auxin naphthaleneacetic acid (NAA) to induce

LR formation and therefore increase the number of cells in which the PUCHI promoter is activated. Forty-six genes were significantly regulated (fold change > 1.5 and *p*-value < 0.05) after 4h NAA + DEX treatment compared to NAA alone. Only 4 of these genes were common with the genes recovered from our *in silico* analysis including 1 gene involved in VLCFA biosynthesis (*KCS8*) thus supporting activation of VLCFA biosynthesis genes downstream of PUCHI. However, this activation was lost when cycloheximide was added (*SI Appendix*, Table S3). Hence, whilst expression of a set of genes encoding for the entire VLCFA elongation pathway was stimulated in a PUCHI-dependent manner during LR formation, these genes were not primary targets of PUCHI.

To confirm the PUCHI-dependent activation of the VLCFA biosynthesis pathway during LR development, we compared expression levels of VLCFA biosynthesis genes in WT and *puchi-1* roots during LR formation by RT-qPCR. An auxin-dependent LR induction system (LRIS, modified from 41); *SI Appendix*, Fig. S4) was used to synchronously induce lateral root formation along the whole primary root. Transcript levels of *KCS1*, *KCS2*, *KCS20*, *KCR1*, *PAS2* and *ECR/CER10* were elevated after LR induction by NAA treatment in WT, but this response was disrupted in the *puchi-1* loss-of-function mutant background (Fig. 1B). Hence, genes encoding key components of the fatty acid elongase complex responsible for VLCFA biosynthesis are induced during LRP development, and this is dependent on the PUCHI transcription factor.

PUCHI controls the spatial expression of VLCFA biosynthesis genes during LR development

We next analyzed the spatial expression pattern of VLCFA biosynthesis genes during LR development using promoter-reporter gene fusions in WT and *puchi-1* backgrounds. A functional *pPUCHI::GFP:PUCHI* reporter is expressed throughout LRP during early stages of development, then the expression is excluded from the tip and limited to the flanks (Fig. 2A; 18). The functional *pKCS1::KCS1:GFP* reporter was expressed in the outermost layer of cells in LRP (Fig. 2B). *KCS6*, *KCR1*, *PAS2*, and *PAS1* GUS reporter transgenes were strongly expressed in developing and emerging LRP (Fig. 2C,E-G), whereas *pKCS20::GUS* was expressed in a few cells of LRP (Fig. 2D). Interestingly, these transgenes displayed distinct expression patterns in emerging LRP, with *pKCS6::GFP:GUS*, *pKCS20::GUS* and *pPAS1::GUS* showing stronger

activity at the base of emerging LRs (Fig. 2C,D,G) whereas expression of *pKCS1::KCS1:GFP*, *pKCR1::GFP:GUS* and *pPAS2::GUS* was stronger at their tips (Fig. 2B,E,F).

When introgressed into the *puchi-1* background, we observed a clear reduction in *pKCS1::KCS1:GFP* and *pKCS6::GFP:GUS* expression in developing LRP (Fig. 2H,I). In contrast, *pKCS6::GFP:GUS* and *pKCS20::GUS* expression was clearly enhanced in *puchi-1* emerging LRs, especially at their tips (Fig. 2J). Loss of function of PUCHI also caused a clear and consistent decrease in expression of *pKCR1::GFP:GUS*, *pPAS2::GUS* and *pPAS1::GUS* in developing LRP and emerging LRs (Fig. 2K-M). Interestingly, expression of these reporters became confined to several basal and flank cells in a majority of emerging LRs. These contrasted changes in reporter genes expression might not completely reflect the endogenous expression of the corresponding genes and suggest that feedbacks might occur. Nevertheless, our analyses confirmed that key VLCFA biosynthesis genes are expressed during LRP development and that their expression patterns in LRP are dependent on PUCHI.

PUCHI inhibits LR initiation but is necessary for later organ development

To better understand the role of PUCHI during LRP development, we carefully characterized the impact of the *puchi-1* mutation on LR formation. Emerged LR and non-emerged LRP were counted along the primary root of 9 day-old seedlings (Fig. 3A). LRP located in the LR formation zone, *i.e.* rootwards of the most recently emerged LR, were scored as "developing LRP" (Fig. 3A, 42). Although the density of emerged LR was not significantly different between *puchi-1* and WT, the density of developing LRP in the LR formation zone was almost three times higher in the mutant compared to WT (Fig. 3B). Consistent with this observation, the distances between two consecutive LRP were much shorter intervals in *puchi-1* than in WT with clusters of developing LRP either along a longitudinal (*i.e.* along a protoxylem pole) or radial (*i.e.* along opposite protoxylem poles) axis (Fig. 3C-D, 43). Thus, our data reveals that PUCHI represses LRP initiation in the pericycle and controls the spacing between LRP. This is consistent with recent results linking PUCHI to lateral inhibition of LR development by a peptide hormone-receptor signaling pathway (44).

In the root branching zone, *i.e.* shootward of the most recently emerged LRs (Fig. 3A, 42), a strikingly high number of non-emerged LRP were present in the mutant, consistent with the observation of higher LRP initiation rate in the LRP formation zone (Fig. 3B). To test if they

correspond to arrested or delayed LRP, we marked and counted the number of emerged LRP in the root branching zone first 9 days after germination and then again after 4 extra days of growth. In both WT and *puchi-1*, we observed that none or very few new LRs had emerged during these additional days, consistent with the hypothesis that these non-emerged LRP have stopped developing (*SI Appendix*, Table S4). Thus, our data suggest that in addition to LRP initiation density, loss of *PUCHI* function also impacts LRP development. To confirm this, we used a gravistimulation-based LR induction system (45, 46) to analyze the kinetics of LRP development in *puchi-1* compared to WT and scored the developmental stages reached by LRP (5) 18 and 48 hours after induction (Fig. 3E). Gravistimulation induced the initiation of LRP development in almost 100% of the *puchi-1* and WT seedlings. However, a delay in *puchi-1* LRP development was already observed 18 hours post gravistimulation compared to WT. At 48 hours, a majority of WT LRP had emerged while most *puchi-1* LRP only reached developmental stages IV, V or VI (Fig. 3E). 6 days after gravistimulation, 100% (53/53) of induced LRP had emerged in the WT background whereas only 70% (35/50) did so in the *puchi-1* background. Hence, PUCHI is required for normal developmental progression of LRP and LR emergence.

VLCFA mutants display similar defects in lateral root development as puchi-1

We performed the same lateral root phenotyping assay for VLCFA mutants. We focused on mutants in KCS genes expressed downstream of PUCHI during lateral root development (kcs1-5, kcs9, kcs2 kcs20), and ECR/CER10 genes because (i) mutants in KCR1, PAS1 and PAS2 display severe and pleiotropic developmental phenotypes (22, 23), (ii) loss-of-function phenotype for those KCS, and ECR genes have been described (21, 25, 26, 33, 47), and (iii) functional redundancy and substrate specificity of KCS enzymes have been studied (21, 48). We did not observe any significant differences in LRP formation and development between kcs9 single mutants nor kcs2 kcs20 double mutant and WT seedlings. However, kcs1-5 mutant, a null allele for the KCS1 gene (33), displayed a root branching phenotype similar to puchi-1, albeit milder (Fig. 3B-E). An ECR mutant (cer10-2) produced similar but weaker phenotype (SI Appendix, Fig. S5). Moreover, double kcs1-5 kcs20 and kcs1-5 cer10-2 mutants resulted in more pronounced LRP development defects compared to the corresponding single mutants (SI Appendix, Fig. S6). Hence, VLCFA biosynthesis loss-of-function mutants exhibited similar defects in LRP development as puchi-1. These defects were weaker than those observed in

puchi-1, possibly due to the fact that PUCHI may simultaneously regulate multiple VLCFA genes and possibly other pathways, and that VLCFA enzymes may act redundantly as indicated by our double mutant analysis.

PUCHI and VLCFAs control pericycle cell proliferation on callus inducing medium

VLCFAs were recently shown to control the ability of pericycle cells to form calli in Arabidopsis roots (33). This prompted us to investigate the role of PUCHI during callus formation. When seven-day old pPUCHI::GFP:PUCHI seedlings were incubated on CIM, reporter gene expression was observed in developing calli (Fig. 4A). Before being transferred to CIM, WT and puchi-1 roots displayed comparable anatomy (Fig. 4B, upper panels). After 4 days on CIM, both WT and *puchi-1* roots responded to the hormonal treatment with pericycle cell proliferation (Fig. 4B, lower panels). However, whereas WT roots produced dome-shaped calli, the *puchi-1* mutant generated a continuous layer of dividing cells along its entire primary root. This phenotype was similar and stronger than the fused-calli phenotype displayed by the VLCFA biosynthesis deficient mutant kcs1-5 (Fig. 4B, SI Appendix, Fig. S7; 33). DEX treatment restored a WT callus formation phenotype in pPUCHI::PUCHI:GR/puchi-1 plants on CIM medium (Fig. 4C) thus confirming that the callus formation phenotype of puchi-1 roots on CIM is caused by loss of function of *PUCHI*. We next analyzed if *PUCHI* is required for the expression of *VLCFA* genes in the context of CIM-induced callus formation. pKCS1::KCS1:GFP, pKCS6::GFP:GUS and pKCR1::GFP:GUS expression were studied after CIM treatment in WT and puchi-1 backgrounds. All three reporter genes were expressed in calli in WT plants and this expression was lost in puchi-1 (SI Appendix, Fig. S8). Hence, PUCHI and VLCFAs restrict pericycle cell proliferation during callus formation and PUCHI is necessary for expression of VLCFA biosynthesis genes in response to CIM treatment.

puchi-1 displays altered VLCFA content

To further demonstrate that the *puchi-1* phenotypes were linked to changes in VLCFA content, we analyzed the fatty acid composition in WT, *kcs1-5* and *puchi-1* CIM-treated roots. CIM treatment was chosen because of the strong associated phenotypes and as it generated a large amount of material for lipidomics. The total amount of fatty acids was similar in all three lines but the amount and proportion of VLCFAs were significantly lower in *kcs1-5* and *puchi-1* compared to WT (Fig. 4D; *SI Appendix*, Fig. S9). Interestingly, a detailed analysis showed that

C22 fatty alcohols (22:0-OH) and C24 sphingolipids (h24:0) were significantly reduced in *puchi-1* while precursors (C16, C18) were more abundant (*SI Appendix*, Fig. S10). VLCFAs are precursors of suberin and cutin biosynthesis. A suberin and cutin-like layer was recently described overlying developing LRP (49, 50). Suberin and root cutin are composed from dicarboxylic acid (DCA), ω-hydroxy acids (ωOH) and fatty alcohols (OH) (50, 51). The level of all C18-containing fatty acids typical of suberin/cutin (ωOH-18, 18-DCA and 18-OH) was similar between *puchi* and WT (*SI Appendix*, Fig. S10). However, we cannot exclude qualitative changes in composition of the suberin/cutin layer in front of LRP that would be regulated by PUCHI.

Discussion

We report that spatio-temporal expression of genes encoding VLCFA biosynthetic enzymes is coordinately regulated by PUCHI during LR formation. Interestingly, while expression of non-redundant *KCR1*, *PAS2* and *PAS1* genes (52, 53) is induced by PUCHI throughout developing LRP in a similar manner, partially redundant *KCS* genes involved in the biosynthesis of VLCFAs of various chain lengths (21, 48, 53) displayed more diverse expression patterns and regulation. Our data suggest that VLCFAs of specific chain lengths may be synthesized and required in different cell types and/or at different stages of LRP development where they might contribute to LRP patterning.

The *puchi-1* loss-of-function mutant has been reported to exhibit LRP with abnormal cell division patterns and with higher cell proliferation rates compared to WT at the organ flanks (18). We report a delay in LRP development and emergence, as well as higher initiation density along the primary root, with frequent clustering of primordia under both normal and tissue culture (CIM) conditions. This is consistent with previous studies indicating that VLCFAs are important to restrict cell proliferation (31). It was recently shown that VLCFA biosynthesis was also necessary to organize cell proliferation into distinct calli in Arabidopsis roots grown on CIM (33). Importantly, callus formation shares common steps with the LRP formation pathway (12–14). We showed here that the *puchi-1* mutation causes roots to form a fused and continuous calli when treated with CIM, a phenotype reminiscent and even stronger than the one reported for *kcs1-5* and other mutants (33). Interestingly, roots of *puchi-1* and VLCFA mutants produced more LRP than WT roots and these LRP exhibit ectopic cell divisions, suggesting that PUCHI

regulates a pathway restricting pericycle cell proliferation that is conserved between LR development and *in vitro* callus formation. Given that defects in VLCFA biosynthesis lead to cell over-proliferation in different contexts such as shoot vasculature (31, 54), CIM-induced calli (33; this study), and in LRP (26), the regulation of VLCFA biosynthesis genes by PUCHI may explain part of the *puchi-1* LRP and CIM phenotype. PUCHI-regulated VLCFA biosynthesis might also be critical for cell patterning in lateral root primordia as evidenced by *puchi-1* phenotype and specific expression profiles of individual VLCFA biosynthesis genes. Further experiments will be needed to test this interesting hypothesis.

VLCFAs could regulate cell proliferation in several ways. VLCFAs can be incorporated into cell membranes and influence their structural and functional dynamics, especially endocytosis, vesicular trafficking, and exocytosis, impacting cell cytokinesis or the targeting to the membrane of important factors, such as auxin transporters (26, 28, 30). More generally, VLCFA content and composition may modulate hormonal signaling dynamics, especially cytokinins and auxins (31, 55). Last, VLCFA distribution could impact the mechanical properties of the tissues by modifying cell wall composition in the endodermis (22) or tissues that overlay LRP (49).

In conclusion, using a systems biology approach we found that the expression of multiple important enzymes catalyzing each of the four steps of the VLCFA elongation cycle are upregulated during LRP development and this is dependent on the AP2/EREPB transcription factor PUCHI. In addition, the *puchi-1* loss-of-function mutant shares similar LRP and callus phenotypes with mutants impaired in VLCFA biosynthesis. Hence, during root branching and root-derived callus formation the PUCHI transcription factor stimulates the expression of key VLCFA biosynthesis genes to regulate cell proliferation, organogenesis and organ spacing.

Materials and Methods

Plant materials, transgenic plant generation, plant growth conditions, and methods for root phenotyping, microscopy and cytological analyses, transcriptome and gene expression level analyses, callus formation assay, and lipidomics profiling by GC-MS are described in *SI Appendix*, *SI Materials and Methods*. Primers used are listed in *SI Appendix*, Table S5. PUCHI::GR expression data were deposited in the Gene Expression Omnibus (GEO) of the National Center for Biotechnology Information (NCBI) under accession number GSE128721.

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Figure Legends

Figure 1. VLCFA biosynthesis genes are induced during LRP development in a PUCHI-dependent manner. (A) Transcripts accumulation levels are expressed in arbitrary units (A.U.). The gray dotted line indicates the time point when *PUCHI* expression reaches a maximum after LR induction. (B) VLCFA biosynthetic gene expression by RT-qPCR in WT (blue) and *puchi-1* roots (orange). Lateral root formation is inhibited in control plants treated with NPA (darker shade), while on NAA (brighter shade) lateral root initiation is induced synchronously along the primary root in both WT and *puchi-1*. Roots were harvested after 24h treatment on 5 μ M NPA or 10 μ M NAA. Normalization was achieved with the *CYCLIN-DEPENDENT KINASE A;1* (*CDKA;1*) gene. The calibrator cDNA is WT under NPA treatment. Data are represented as Mean \pm SEM (standard error of the means) of three biological replicates. Significance was determined by Student's t test (* P < 0.05, ** P < 0.01).

Fig. 2: VLCFA genes are expressed in developing LRP and their expression patterns are dependent on PUCHI. (A) Expression of *pPUCHI::GFP:PUCHI* in *puchi-1* background is observed in developing LRP and is gradually confined to their base and flanks. (B) to (M) Expression patterns of various translational and transcriptional reporter constructs of VLCFA biosynthesis genes in typical WT (B to G) and *puchi-1* (H to M) LRP and newly emerged LR. Scale bars = 50μm. Numbers indicate the percentage of LRP or LR displaying the corresponding expression pattern. n = 30-40 seedlings for each GUS assay.

Figure 3. LR initiation is enhanced but LRP development is delayed in *puchi-1* mutant. (A) Root branching events were scored in three distinct developmental zones of Arabidopsis primary root regarding lateral root formation as suggested in (42). (B) Density of developing LRP, emerged LR, delayed LRP, and total LR initiations (LRP + LR) in 9-day old WT, puchi-1 and kcs1-5 seedlings. Developing LRP are LRP scored in the LR formation zone. Delayed LRP are defined as those located in the branching zone of the primary root but have not crossed the epidermis. Data are represented as Mean \pm SEM of three biological replicates; number of seedlings \geq 20 in each repeat. Significance was determined by Student's t test. (C) Frequency distribution of distances between two consecutive LRP in WT, kcs1-5 and puchi-1 roots. Each bin of the histogram represents a range of 300 μ m. Number of LRP = 222, 208 and 228 for WT, kcs1-5 and puchi-1, respectively. The orange bar in each histogram indicates the mean LRP distance in each genotype. The stars in the histograms for kcs1-5 and puchi-1 indicate the significant difference between these mean distances compared to that of WT. (D) Examples of longitudinal and radial clusters of LRP in *puchi-1* roots. Arrowheads indicate LRP. Scale bars = 50µm. (E) Distribution of developmental stages as described by Malamy and Benfey (5) achieved by gravistimulationinduced LRP formation in WT, kcs1-5 and puchi-1 roots at 18 and 48 hours after the gravistimulation. Data are represented as Mean ± SEM of three biological replicates, with

number of seedlings \geq 20 in each repeat. For all occasions, * P < 0.05, ** P < 0.01, *** P < 0.001.

Figure 4: Callus formation is enhanced in *puchi-1* roots. (A) *pPUCHI::GFP:PUCHI* expression in calli induced by CIM, scale bars = 50 μm. (B) 7-day-old *puchi-1*, kcs1-5 and WT seedlings grown on ½ MS medium (top), and after 4 more days of growth on CIM (bottom). (C) pPUCHI::PUCHI:GR/puchi-1 plus dexamethasone (DEX) restores callus formation in *puchi-1* roots on CIM. n = 40 for WT, kcs1-5 and puchi-1. n = 20 for pPUCHI::PUCHI:GR/puchi-1, scale bars = 0.1mm. (D) Global fatty acids of WT, puchi-1 and kcs1-5 roots after 4 days on CIM. The results are the sum of carboxylic acids, α,ω-dicarboxylic acids, ω-hydroxy acids, fatty alcohols and 2-hydroxy fatty acids. Data are represented as Mean ± SEM of six, six and five biological replicates for WT, puchi-1 and kcs1-5, respectively. * P < 0.05, ** P < 0.01, *** P < 0.001.