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Mediator Med23 deficiency in smooth muscle cells prevents neointima formation after arterial injury

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Dear Editor,

Vascular smooth muscle cells (VSMCs) that line the arteries and veins, are able to increase their proliferation rate greatly following vessel wall injury and in atherogenesis. Changes in VSMC proliferation rate and differentiation state have been proposed as required procedures of injury repair. The regulation of the balance between proliferation and differentiation is of key importance in vascular biology. Our previous works have shown that Mediator MED23 subunit restricts smooth muscle cell lineage development and promotes growth related gene expression¹. Recently, we investigated the roles of Med23 in VSMC proliferation and differentiation as well as injuryinduced neointima formation by generating and examining the smooth muscle specific Med23-knockout $(Med23^{sm-/-})$ mice. Our results showed that Med23 regulates the balance of VSMC growth and differentiation in mouse aorta. The $Med23^{\text{sm}-/-}$ mice showed enlarged lumen and impaired contractility of aorta. RNA profiling of the aorta samples revealed that gene expression programs switching between proliferation and differentiation upon the presence or absence of Med23. More importantly, we found that Med23 deficiency prevented neointima formation after vascular injuries, through repressing VSMC proliferation. Collectively, our data demonstrate that Med23 has positive effects on VSMC plasticity and plays a novel pathological role in vascular injury induced neointima formation through promoting the proliferation and growth of VSMCs, thus providing a novel mechanism understanding the VSMC plasticity and related vascular diseases.

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Previously we have demonstrated that Mediator MED23 plays a repressing role in RhoA signaling pathway and smooth muscle cell fate, but acts as a positive regulator for Ras-targeted genes and the adipocyte fate¹. Nevertheless, the roles of Med23 in regulating smooth muscle development and function have not been examined in vivo in mice. Since homogenous knockout of Med23 is embryonic lethal², we generated smooth muscle specific *Med23*knockout ($Med23^{sm-/-}$) mice to investigate the in vivo function of Med23. We crossed conditional Med23knockout mice (Med23^{fl/fl})³ with the aortic smooth muscle actin (*Acta2*)-Cre expressing line⁴ to generate the $Med23^{sm-/-}$ mutant mice. Western blot analysis of tissue samples from Med23^{sm-/-} mice showed specific and efficient Med23 ablation in stomach and aorta, the walls of which are lined with smooth muscle, but not in kidney, spleen, liver, fat tissue or lung (Fig. 1a). Note that Med23 protein level in a rta from control mice was low, suggesting a repressing function of Med23 during smooth muscle cell (SMC) differentiation, whereas essentially no Med23 protein was detected in the aorta from $Med23^{sm-/-}$ mice (Fig. 1a). Western blot analysis of isolated VSMCs from the aortas confirmed Med23 deletion in the VSMCs from the aortas of Med23^{sm-/-} mice (Supplementary Fig. S1f).

Med23^{sm-/-} mice were born at the expected Mendelian ratio, and they appeared to be normal and indistinguishable from their wild-type littermates. Careful analysis of the aortas revealed the differences between control and *Med23*^{sm-/-} mice: *Med23*^{sm-/-} mice exhibited slightly dilated aortas (Fig. 1b, left and middle panels) and enlarged aorta lumen (Fig. 1c) compared with control mice. In spite of this, blood pressure of *Med23*^{sm-/-} mice was comparable to control mice on either normal or high salt diets (Fig. 1d). We then examined the mesenteric arteries, since the mesenteric arterial network that carries about one fourth of the blood pumped from heart

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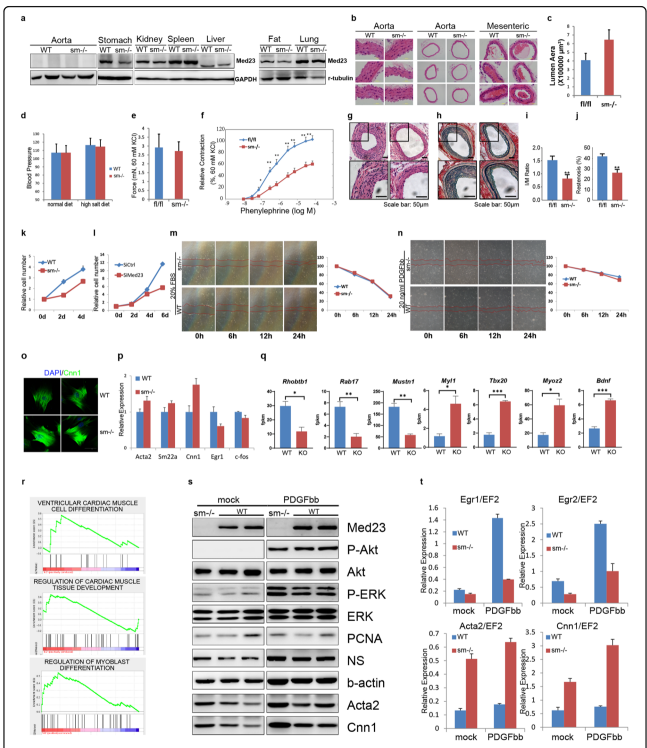


Fig. 1 Med23 deficiency in smooth muscle cells prevents neointima formation after arterial injury. a Immunoblot of Med23 in various tissues isolated from $Med23^{sm-/-}$ and control mice. **b**, **c** Aorta, mesenteric arteries (**b**) and aortic lumen area (**c**) in $Med23^{sm-/-}$ and control mice. **d** Blood pressure of $Med23^{sm-/-}$ and control mice on normal or high salt diets. **e** Isometric contraction of aorta rings prepared from $Med23^{sm-/-}$ and control mice (60 mM KCl). **f** Dose-response curves for PE-induced contraction in $Med23^{sm-/-}$ and control aortas (60 mM KCl). **g** Representative H/E staining of femoral arteries from $Med23^{sm-/-}$ and control mice. **h** Representative elastin staining of femoral arteries from $Med23^{sm-/-}$ and control mice. **i**, **j** Intima-to-media ratio (**i**) and restenosis index (**j**) of femoral arteries harvested from $Med23^{sm-/-}$ and control mice. **k** Cell proliferation of VSMCs isolated from the aorta. **l** Cell proliferation of A7R5 cells after viral-mediated siRNA knockdown of Med23. **m**, **n** Wound healing assay in the presence of either 20% FBS (**m**) or 20 ng/ml PDGFbb (**n**). **o** Immunostaining of $Med23^{sm-/-}$ VSMCs. **p** Expression levels of SMC genes and growth-related genes in the isolated VSMCs. **q**, **r** RNA-seq analysis of the aorta samples from $Med23^{sm-/-}$ mice. **s**, **t** Western blot (**s**) and real-time PCR (**t**) of isolated VSMCs from control and $Med23^{sm-/-}$ mice. NS, nucleostemin. Data are presented as means \pm SEM. *P < 0.05, **P < 0.01, ****P < 0.001, **s control.

contributes significantly to vascular resistance and blood pressure⁵. Consistently, the morphology of mesenteric arteries was similar between $Med23^{sm-/-}$ mice and control mice (Fig. 1b, right panels).

Next, we measured the isometric contraction of aortas. Compared to control mice, the contractility of aorta rings from $Med23^{sm-/-}$ mice was only slightly decreased in high-K⁺ medium (Fig. 1e). However, dose-response curves for the effect of phenylephrine (PE), an α -adrenergic receptor agonist that stimulates blood vessel contraction by activating postsynaptic adrenergic receptors, showed significant reduced levels of PE-induced contraction in Med23-depleted aortas (Fig. 1f). Collectively, $Med23^{sm-/-}$ mice show impaired aortic vasoconstriction, especially in response to PE.

The enlarged lumen and impaired contractility of aortas from Med23^{sm-/-} mice led us to investigate whether these mice possess other changes in vascular function. Neointima formation, which causes unfavorable restenosis after endovascular interventions such as stenting or angioplasty, is a pathological healing response induced by vessel wall injuries. We examined the occurrence of neointima formation in Med23^{sm-/-} mice in a femoral artery wire injury model⁶. Four weeks after the injury, the femoral arteries from control mice showed typical neointima formation, in which smooth muscle cells and their extracellular matrix deposits came out inside the internal elastic lamina and evade the lumen area of the blood vessel (Fig. 1g, h, left panels). Interestingly, we observed greatly reduced level of neointima formation in femoral arteries harvested from Med23^{sm-/-} mice (Fig. 1g, h, right panels). Consistently, the femoral arteries harvested from $Med23^{sm-/-}$ mice showed substantial decrease of both intima-to-media (I/M) ratio (defined as the intimal area divided by the medial area that was calculated as the area encircled by the external elastic lamina minus the intima area, Fig. 1i) and restenosis index (percentage of luminal narrowing, defined as the intimal area divided by the area encircled by the internal elastic lamina, Fig. 1j), in comparison to those from WT mice, which demonstrated ample neointima formation. Therefore, Med23 deficiency represses neointima formation that contributes to restenosis.

Studies indicate that medial SMCs will be activated and start to replicate and migrate in response to vessel wall injury^{7,8}. With the stimulating from local growth factors, such as platelet-derived growth factor (PDGF)⁹, SMCs migrate from tonica media to intima, and proliferate further there. Although the proliferation of VSMC plays a central role in injury repair, excess proliferation of VSMC may lead to arterial injury induced neointima formation. In order to investigate the cellular mechanism behind the reduced neointima formation in *Med23*^{sm-/-} mice after arterial injury, we examined cell proliferation and

migration of VSMCs isolated from the aorta (Fig. 1k-n). Comparing to WT VSMCs, VSMCs from Med23^{sm-/-} mice showed reduced rate of cell proliferation during the 4 days period assayed (Fig. 1k). Consistently, viralmediated siRNA knockdown of Med23 in a rat smooth cell line, A7R5, led to reduced cell proliferation rate (Fig. 11) and lower DNA replicating rate, indicated by EdU incorporation (Supplementary Fig. S1a-c). Whereas in the wound healing assays stimulated with either 20% FBS or 20 ng/ml PDGFbb, VSMCs from Med23^{sm-/-} mice were able to cover the wounded area to the same extend as those from control mice (Fig. 1m, n). siMed23 A7R5 cells also showed similar migration ability as control cells in the wound healing assay (Supplementary Fig. S1d, e). Since cell proliferation, migration and increased cell size all contributed to wound healing, we next performed transwell assay to specifically measure the migration capacity of Med23^{sm-/-} VSMCs. Note that both wound healing assay and transwell assay measure cell migration in vitro, which may not reflect cell migration capacity in vivo. However, transwell assay is closer to migration models. We found that VSMCs from Med23^{sm-/-} mice were able to migrate as fast as those from control mice (Supplementary Fig. S1f, g). These results indicate that the reduced neointima formation observed in Med23^{sm-/-} mice is not caused by the reduced ability of VSMCs to migrate to intima, but rather caused by the reduced proliferation rate of VSMCs in the absence of *Med23*.

Comparing with many other terminal differentiated cell types, one special character of VSMC is its plasticity of switching between differentiation and growth state, which is essential for vessel wall injury repairing, restenosis, atherosclerosis, etc. The expression levels of Med23 in normal aorta samples are already very low (Fig. 1a), suggesting that Med23 level may be repressed in VSMCs to maintain the differentiated contractile phenotype. To look through the growth and differentiation of VSMCs, we examined the morphology and gene expression of isolated VSMCs from $Med23^{sm-/-}$ mice. First, western blot analysis confirmed specific and efficient Med23 deletion in VSMCs isolated from the aorta of Med23^{sm-/-} mice (Supplementary Fig. S1f). SMC marker CNN1 is a microfilament-associated protein that functions in the contraction of smooth muscles¹⁰. Immunostaining revealed increased level of Cnn1, which forms highly organized stress fibers that may alter cell contraction properties, in *Med23*^{sm-/-} VSMCs comparing to control cells (Fig. 10). In addition to Cnn1, the expression levels of SMC genes such as Acta2 and Sm22a were increased in the isolated $Med23^{sm-/-}$ VSMCs, whereas the expression levels of growth-related genes like Egr1 and c-fos were decreased (Fig. 1p), which is consistent with the genes regulated by MED23 in a mesenchymal stem cell line and adipose-derived stem cells¹. To further investigate the changes in gene expression profiling associated with Med23 ablation, we performed RNA-seq analysis of the aorta samples from $Med23^{sm-/-}$ mice (Fig. 1q, r). We found that Rab17 (a member of RAS oncogene family), Rhobtb1 (Rho-related BTB domain containing 1) and musculoskeletal gene Mustn1 were among the down-regulated genes in $Med23^{sm-/-}$ samples, in addition to Egr1. While the expression of muscle-related genes Myl1, Tbx20, Myoz2 and brain derived neurotrophic factor (Bdnf) were up-regulated in $Med23^{sm-/-}$ samples (Fig. 1q). Taken together, our data indicate that Med23 deletion affects the balance between growth and differentiation of VSMCs in mouse aorta, with growth related genes down-regulated and SMC lineage-related genes up-regulated.

PDGF induces SMC migration and further proliferation, and represses smooth muscle differentiation, by specifically suppressing the expression of contractile protein genes, inducing the phosphorylation of Elk-1 and increasing the association of Elk-1 with SRF¹¹. In order to examine whether Med23 promotes cell proliferation via upregulating PDGF signaling, we compared the downstream effectors of PDGFbb in Med23^{sm-/-} VSMCs and control cells. Upon PDGFbb stimulation, VSMC samples from both control and Med23^{sm-/-} mice showed significantly increased levels of phosphorylated ERK (p-ERK) and phosphorylated Akt (p-Akt) (Fig. 1s). Therefore, Med23 deletion does not affect the downstream signaling of PDGFbb. In the absence of PDGFbb, the expression levels of growth-related genes *Egr1* and *Egr2* were slightly lower in $Med23^{sm-/-}$ samples than in WT samples. Egr1 and Egr2 exhibited huge increases in expression levels in WT samples upon PDGFbb stimulation, in contrast to only moderate increase in $Med23^{sm-/-}$ samples (Fig. 1t). On the other hand, both the RNA and protein levels of muscle lineage genes Acta2 and Cnn1 were significantly higher in $Med23^{sm-/-}$ samples than in WT samples in the absence of PDGFbb (Fig. 1s, t). Upon PDGF stimulating, the expression of Acta2 and Cnn1 further increased in $Med23^{sm-/-}$ samples but remained at low levels in control samples (Fig. 1t). These results indicate that the reduced neointima formation in Med23^{sm-/-} mice is not caused by defects in injury-induced signal transductions of SMC, but rather caused by the changes in transcription of SMC growth and differentiation genes, which is specifically controlled by Med23 subunit of the Mediator.

Taken together, our results suggest that *Med23* is important for the function of smooth muscle in vivo, through maintaining the balance of SMC growth and differentiation. Moreover, *Med23* is involved in neointima formation induced by arterial injury, which may improve our understanding on vascular diseases.

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Conflict of interest

The authors declare that they have no conflict of interest.

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