Activation of the Non-receptor Tyrosine Kinase cSrc in Macrophage-rich Atherosclerotic Plaques of Human Carotid Arteries

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To determine the involvement of the non-receptor tyrosine kinase cSrc in plaque destabilization in carotid atherosclerosis (CAS), which is responsible for cerebral infarction, we performed quantitative and morphological detection of phosphorylated active cSrc (p-cSrc) and histopathological examination in CAS lesions. We examined carotid endarterectomy specimens obtained from 32 CAS patients. Each specimen was used for immunoblot and immunohistochemical analyses of p-cSrc, histopathological analysis, and image analysis of macrophage content. There was a strong positive correlation between cSrc activation on blots and macrophage content on sections. When we defined the macrophage-rich plaque (MRP) and the macrophage-poor plaque (MPP) as having macrophage content more and less than 5%, respectively, the p-cSrc density and the occurrence of plaque hemorrhage and thrombus formation were significantly increased in the MRP group (n=18) compared to the MPP group (n=14). p-cSrc immunoreactivity was localized in lesional endothelial cells, macrophages, and smooth muscle cells, which contained proinflammatory substances: the upstream oxidized low density lipoprotein, tissue factor and osteopontin, and the downstream active forms of extracellular signal-activated kinase and p38 and nuclear factor-κB. Our results suggest that cSrc activation in lesional cells contributes to plaque destabilization in CAS via persistent inflammation.

Key words: atherosclerosis, carotid artery, Src, inflammation, plaque instability

I. Introduction

Several investigations have indicated that carotid atherosclerosis (CAS) is a major cause of cerebral infarction [1]. Acute-onset ischemic brain attack occurring in CAS patients results from acute occlusion of the lumen by atherothrombosis or hemorrhage in the plaques or from thromboembolism originating in atherothrombotic lesions [36, 37]. These events associated with CAS appear to be based on plaque instability, as described in studies on coronary atherosclerosis [11]. Emerging evidence suggests implications for persistent inflammation in plaque destabilization of carotid arteries [32]. Several studies on atherosclerosis of carotid and coronary arteries have shown that plaques containing a lot of macrophages are unstable, while those containing only a few macrophages are stable [11, 24, 32, 36, 37].

It has been shown that vascular endothelial cells (VECs), T-lymphocytes, and vascular smooth muscle cells (VSMCs) as well as macrophages participate in atherogenic inflammation [24, 32, 37]. VECs exposed to pathological conditions including risk factors for atherosclerosis release chemokines into blood and express cell adhesion molecules (CAMs) on the apical cell surface. Chemokines stimulate circulating monocytes to express integrin αMβ2 as the receptor for CAMs, and the binding of CAMs to integrin αMβ2 induces monocyte transmigration into the subendothelial vascular wall tissue. Migrating monocytes are transformed to macrophages in the presence of monocyte colony-
stimulating factor released from T-lymphocytes. In atherosclerotic plaques, migrating VSMCs secrete collagen to form neointima, while macrophages secrete matrix metalloproteinases to degrade the surrounding extracellular collagen matrix. The latter leads to plaque destabilization, as evidenced by the vulnerability of the fibrous cap, followed by ulceration, hemorrhage, rupture, and thrombus formation, or both. Macrophages also secrete vascular endothelial growth factor, resulting in neovascularization in the plaques. The newly formed capillary vessels are mechanically vulnerable and readily rupture to form hematoma in the plaques, so-called plaque hemorrhage. All plaque-composing cells also secrete several cytokines and growth factors. These observations would indicate the crosstalk between the lesional cells in atherosclerotic plaques, and suggest that macrophage content is a marker of inflammation and destabilization of atherosclerotic plaques.

In atherogenesis, angiotensin II, oxidized low density lipoprotein (OxLDL), activated factor VII (FVIIa), and osteopontin have been shown to trigger inflammatory response, by binding to their receptors: angiotensin II type 1 receptor (AT-R), scavenger receptor, tissue factor, and integrin αvβ3, CD44 or fibronectin, respectively [3, 8, 21, 24, 26, 27, 35, 38]. Actually, recent in vivo studies have shown increased levels of OxLDL, tissue factor, and osteopontin in macrophage-rich or symptomatic plaques [10, 14, 23]. Moreover, atherothrombosis is enhanced in animals overexpressing tissue factor [12] and osteopontin [6], while it is attenuated in animals deficient in tissue factor [34] and osteopontin [33]. These observations suggest critical roles for tissue factor and osteopontin in atherogenesis. Several in vitro studies have demonstrated that extracellular stimuli induced by angiotensin II, OxLDL, tissue factor, and osteopontin activate the non-receptor tyrosine kinase cSrc, collectively. cSrc is a proto-oncogene product that was identified as an intracellular homologue of the Rous sarcoma virus oncogene product vSrc [35]. cSrc is a 60 kDa protein, expressed ubiquitously in several types of cells including inflammatory cells, and localized in the cytosol under physiological conditions. The binding of extracellular ligands, such as growth factors, cytokines, factor VIIa, and osteopontin, to their respective membrane-bound receptors results in recruitment of the cytosolic cSrc to the intracellular domains of the receptors followed by phosphorylation-mediated activation of cSrc. The activated cSrc starts inflammatory and proliferative cell signaling via activation of mitogen-activated protein kinases (MAPKs) such as extracellular signal-regulated kinase (ERK), c-Jun N-terminal kinase (JNK), and p38 as well as transcription factors including nuclear factor-κB (NF-κB) [3–5, 7, 15, 21, 35]. However, it remains to be determined whether cSrc activation may occur in association with atherogenic inflammation and plaque destabilization in vivo. To address this issue, we investigated morphological and quantitative analyses of active form of cSrc phosphorylated at amino acid residue Y416 (p-cSrc) and verified the correlation between macrophage content and cSrc activation in CAS lesions. This is the first report concerning both immunohistochemical detection and quantitative measurement of p-cSrc in atherosclerotic plaques of human carotid arteries.

II. Materials and Methods

Subjects and tissue preparation

This study was approved by the Ethics Committee of Tokyo Women’s Medical University and was carried out on 32 CAS patients [ages: 50–82 (69.7±8.2) y] who underwent carotid endarterectomy (CEA). The clinical features of the examined cases are summarized in Table 1. Each CEA specimen was divided into equal two parts and processed for making freshly frozen materials and 30% sucrose-immersed, optical cutting temperature (OCT) compound-immbedded frozen materials. Both of them were stored at −80°C until use.

Immunoblot analysis

The primary antibodies employed in immunoblotting were mouse monoclonal IgG against p-cSrc (Clone 9A6, Upstate, Lake Placid, NY, USA; 1:5,000) and mouse monoclonal IgG against β-actin (Clone AC-15, Sigma, St. Louis, MO, USA; 1:10,000). Freshly frozen materials were homogenized with 10-fold volumes of radioimmunoprecipitation assay (RIPA) buffer consisting of 50 mM Tris-buffered saline, pH 7.5 (TBS), 1% sodium dodecyl sulfate (SDS), 1% glycerol, 1% deoxycholic acid, 0.1% Tween-20, and 0.3% 1,4-dithiothreitol (DTT) as well as the protease inhibitor cocktail Complete MIni™ (Roche Diagnostics, Mannheim, Germany) according to the manufacturer’s instructions. Ho-

| Characteristics | n | % |
|-----------------|---|---|
| Total number of subjects | 32 | 100 |
| Sex | | |
| Male | 28 | 87.5 |
| Female | 4 | 12.5 |
| Phenotype | | |
| Symptomatic case | 12 | 37.5 |
| Asymptomatic case | 20 | 62.5 |
| Risk factor | | |
| Hypertension | 24 | 75 |
| Hyperlipidemia | 14 | 43.8 |
| Diabetes mellitus | 10 | 31.3 |
| Smoking | 10 | 31.3 |
| Ischemic heart disease | 13 | 40.6 |
| Therapeutic drugs | | |
| Aspirin | 28 | 87.5 |
| Statin | 9 | 28.1 |
| Fibrate | 0 | 0 |
| TZD | 0 | 0 |
| ACEI | 4 | 12.5 |
| ARB | 13 | 40.6 |

ACEI, angiotensin II-converting enzyme inhibitor; ARB, angiotensin II receptor blocker; TZD, thiazolidinedione.
mogenates were centrifuged for 15 min at 4°C with 15,000 g to obtain supernatant samples. Protein concentration of each sample was measured by the Bradford method [29], and samples were subsequently mixed with equal volume of Laemmli’s buffer and boiled for 5 min. Aliquots of total protein extracts (30 µg/lane) were loaded on a 10% SDS-polyacrylamide gel, electrophoresed, and transferred onto a polyvinylidene difluoride (PVDF) membrane (Millipore, Tokyo, Japan). Blots were pretreated overnight at room temperature with 5% bovine serum albumin (BSA) in TBS containing 0.1% Tween-20 (TBS-T) solution, and then incubated with the anti-p-cSrc antibody in Can Get Signal solution (Toyobo, Tokyo, Japan) for 1 hr at room temperature followed by horseradish peroxidase (HRP)-labeled antiserum IgG antibody (DakoCytomation, Kyoto, Japan; 1:10,000) in Can Get Signal solution for 1 hr at room temperature. Immunoreaction for p-cSrc was visualized by the chemiluminescence method using the ECL-plus kit (GE Healthcare, Buckinghamshire, UK). Blots were then treated for 30 min at 50°C with a stripping buffer composed of 50 mM TBS, 2% SDS, and 1% β-mercaptoethanol, rinsed in TBS-T, treated overnight at room temperature with 5% skim milk/TBS-T solution, and incubated with the anti-β-actin antibody in 5% skim milk/TBS-T solution followed by the HRP-labeled anti-mouse IgG antibody in 5% skim milk/TBS-T solution. Immunoreaction for β-actin was detected with the ECL kit (GE). Lanes processed with omission of the primary antibodies or by incubating with non-immune mouse IgG served as negative reaction controls. Immunoreactive signals for both p-cSrc and β-actin were detected by the HRP-labeled anti-mouse IgG antibody, and β-actin was detected with the HRP-labeled anti-β-actin antibody. Immunoreaction was visualized by the ECL method. Briefly, sections were incubated with the anti-p-cSrc antibody in Can Get Signal solution for 1 hr at room temperature. Immunoreaction for p-cSrc was visualized by the chemiluminescence method using the ECL-plus kit (GE Healthcare, Buckinghamshire, UK). Blots were then treated for 30 min at 50°C with a stripping buffer composed of 50 mM TBS, 2% SDS, and 1% β-mercaptoethanol, rinsed in TBS-T, treated overnight at room temperature with 5% skim milk/TBS-T solution, and incubated with the anti-β-actin antibody in 5% skim milk/TBS-T solution followed by the HRP-labeled anti-mouse IgG antibody in 5% skim milk/TBS-T solution. Immunoreaction for β-actin was detected with the ECL kit (GE). Lanes processed with omission of the primary antibodies or by incubating with non-immune mouse IgG served as negative reaction controls. Immunoreactive signals for both p-cSrc and β-actin were detected by the HRP-labeled anti-mouse IgG antibody, and β-actin was detected with the HRP-labeled anti-β-actin antibody. Immunoreaction was visualized by the ECL method.
the photogenic data were digitally imported into a personal computer using Photoshop image software. Both the CD68-immunoreactive area (mm$^2$) and the total area (mm$^2$) in a plaque were measured using NIH image software.

**Statistics**

The relationship between macrophage content and cSrc activation in the examined cases was statistically verified by nonparametric correlation analysis. Macrophage content in each plaque was expressed as a percentage of the CD68-immunoreactive area in the total area on a section. cSrc activation in each plaque was expressed as a relative value of the densitometric p-cSrc/β-actin ratio on a lane. Linear equation ($y=ax+b$) and correlation coefficient ($r$) were obtained, and $r$ value more than 0.6 was considered to be strongly correlated. The examined lesions were then divided into macrophage-rich plaques (MRPs) and macrophage-poor plaques (MPPs), as described previously [23]; MRPs and MPPs were defined as having macrophage content more than 5% and less than 5%, respectively. The densitometric p-cSrc/β-actin data in the MRP and MPP groups were expressed as mean ± standard deviation, and compared by unpaired Student’s t-test between the two groups. The histopathological findings were compared by using chi square test between the two groups. $P$ value less than 0.05 was considered statistically significant.

**III. Results**

**Relationship between macrophage content and cSrc activation**

Immunoreactive signals for p-cSrc and β-actin appeared on immunoblots at 66 kDa and 42 kDa, respectively (Fig. 1A). No immunoreaction product was detectable on negative reaction control lanes. There was a strong positive correlation between the CD68-immunoreactive areas on sections and the optical p-cSrc densities on blots (Fig. 1B). The 32 examined cases were divided into 14 MPP cases and 18 MRP cases. Unpaired Student’s t-test revealed that the optical p-cSrc densities on blots were significantly increased in the MRP group compared to the MPP group (Fig. 1C).

**Comparison of histopathological findings between the MPP and MRP groups**

The histopathological findings were compared by chi square test between the MRP and MPP groups (Table 2). Based on the hallmarks of plaque instability, the occurrence of plaque hemorrhage and thrombus formation significantly increased in the MRP group than in the MPP group. By contrast, there was no significant difference in the occurrence of ulceration and calcification of the CAS plaques between the two groups.

**Immunohistochemical observations of p-cSrc and related substances**

No immunoreaction product deposit was visible on negative reaction control sections (data not shown). At a low

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**Table 2. Comparison of histopathological features between the two groups**

| Characteristics         | MPP group (cases) | MRP group (cases) | $P$ value |
|-------------------------|-------------------|-------------------|-----------|
| Number of subjects      | 14                | 18                |           |
| Plaque morphology       |                   |                   |           |
| Ulceration              | 2                 | 3                 | NS        |
| Hemorrhage              | 3                 | 13                | 0.011     |
| Thrombus formation      | 0                 | 9                 | 0.0018    |
| Calcification           | 12                | 16                | NS        |

MPP, macrophage-poor plaque; MRP, macrophage-rich plaque; NS, not significant.
magnification, p-cSrc immunoreactivity was distributed in the CAS plaques (Fig. 2), and seemed to increase in the MRP group compared with the MPP group. The distribution of p-cSrc appeared to overlap with that of CD68 and SMA to some extent; however, the cSrc immunoreactivity was less intense in large foamy macrophages neighboring the lipid core. At a higher magnification, p-cSrc immunoreactivity was localized in the cytoplasm of VECs lining the inner surface of carotid arteries and newly formed capillary vessels in atherosclerotic lesions as well as macrophages and migrating VSMCs (Fig. 3). p-cSrc immunoreactivity in the cytoplasm of these lesional cells was colocalized with OxPC, tissue factor, osteopontin, p-ERK, p-p38, and NF-κB (Fig. 4). NF-κB immunoreactivity was also localized in the nucleus of the lesional cells.

**IV. Discussion**

In the present study, it is noteworthy that p-cSrc was detected in human CAS lesions. There was a strong positive correlation between the CD68-immunoreactive areas on sections and the p-cSrc-immunoreactive densities on blots. This suggests that cSrc activation is enhanced in parallel with an increase in macrophage content in atherosclerotic lesions. The fact that both plaque hemorrhage and thrombus formation as the histopathological hallmarks of plaque instability were more common in the MRP group than in the MPP group shows the close relevance between macrophage content and plaque destabilization, as previously discussed [16]. The lack of difference in the occurrence of calcification between the two groups suggests dual roles for osteopontin in atherogenesis, including plaque inflammation and extracellular calcification. Furthermore, p-cSrc immunoreactivity was localized in VECs, macrophages, and VSMCs in CAS plaques. Several experimental studies have indicated that macrophage content in atherosclerotic lesions reflects inflammatory activity and plaque instability [11, 24, 32, 36, 37]. Given these observations, cSrc activation in lesional cells may be involved in inflammation and instability of atherosclerotic lesions.

Some in vitro studies have demonstrated that the binding of OxLDL, FVIIa, and osteopontin to their specific receptors gives rise to cSrc activation, resulting in the activation of inflammatory cell signaling [17, 24, 35]. In relation to this, we demonstrated the colocalization of p-cSrc with OxLDL, tissue factor, and osteopontin in lesional cells such as VECs, macrophages, and VSMCs. After uptake into the lesional cells via scavenger receptors, OxLDL in the cytoplasm activates cSrc [9, 17, 20]. It is known that inflamed plaques have greater levels of the hepatocyte-produced, plasma-mediated FVIIa, as a result of increased endothelial permeability in the lesions. This leads to increased chances of FVIIa binding to the receptor tissue factor, resulting in cSrc activation [8, 26–28, 35]. Similarly, osteopontin binding to the receptor integrin α₅β₃, CD44 or fibronectin induces cSrc activation [24]. In addition, the binding of angiotensin II, one of the most important factors involved in atherogenesis, to AT₁R also brings about cSrc activation [3, 21, 30]. These studies attributed cSrc activation to increased oxidative stress and cytosolic calcium levels.

It has been shown that cSrc activation exerts critical effects on the induction of proinflammatory gene products via activation of the MAPK pathway and the NF-κB pathway [3–5, 7, 8, 15, 21, 26, 27, 35, 38]. In relation to this, we demonstrated the colocalization of p-cSrc with p-ERK, p-p38, and NF-κB in CAS lesions, as previously described [19, 21, 22, 25, 31]. Several experimental studies have shown that activated cSrc catalyzes the phosphorylation of phosphatidylinositol 3-kinase (PI3K) and protein kinase C (PKC) [4, 5, 7, 8, 15, 26, 27, 35] to activate them. PI3K activates ERK and p38 via activation of Ras and Raf-1, respectively, while PKC activates ERK and NF-κB via activation of Raf-1 and inhibitor of κB kinase, respectively. Alternatively, both the...
MAPK and NF-κB pathways induce tissue factor [2, 18, 22] and osteopontin [38], which have been shown to be overexpressed in symptomatic CAS plaques [10, 14]. Moreover, both the binding of FVIIa and osteopontin to their specific receptors, as mentioned above, consequently induce cSrc activation again. Thus, it is likely that cSrc activation may contribute to persistent inflammation in CAS lesions.

Collectively, we demonstrated the following four findings using CEA specimens: (1) the strong positive correlation between macrophage content and cSrc activation, (2) the increased vulnerability of MRPs compared to MPPs, (3) the localization of p-cSrc in VECs, macrophages, and VSMCs, and (4) the colocalization of p-cSrc with the upstream OxLDL, tissue factor, and osteopontin and the downstream activated ERK, p38, and NF-κB. Our results suggest that cSrc activation in lesional cells mediates persistent inflammation in CAS plaques and contributes to plaque destabilization, and point to the possibility that the inhibition of cSrc may ameliorate these pathological conditions. Recent investigations have documented implications for cSrc activation in transactivation and nuclear translocation of the transcription factor signal transducer and activator of transcription (STAT) in atherosclerosis models [21, 39]. Little is known, however, about the relationship between cSrc and STAT in atherogenesis. Answers to this and other questions concerning the modulation of cSrc activity need to be addressed to further the understanding of the pathomechanism of plaque destabilization and the prevention of ischemic events in the brain.

**Fig. 3.** Photomicrographs of MRP sections immunostained for p-cSrc (A–C, E), CD68 (D) and SMA (F). Sections (C, E) are consecutive to sections (D, F), respectively. Immunoreaction on all sections is visualized by the ABC method with DAB as the chromogen (brown). Arrowheads, blank arrows, and solid arrows indicate VECs, macrophages, and VSMCs, respectively. Bar=50 µm (A–F). ABC, avidin-biotin-immunoperoxidase complex; DAB, 3,3’-diaminobenzidine tetrahydrochloride; MRP, macrophage-rich plaque; p-cSrc, active form of cSrc phosphorylated at Y416; SMA, α-smooth muscle actin; VECs, vascular endothelial cells; VSMCs, vascular smooth muscle cells.
Fig. 4. Photomicrographs of MRP sections immunostained for p-cSrc (D–F, J–L), OxPC (A), tissue factor (B), osteopontin (C), p-ERK (G), p-p38 (H), and NF-κB (I). Panels (A–C, G) indicate identical regions to panels (D–F, H), respectively, on consecutive sections. Panels (H, I) indicate identical regions to panels (K, L), respectively, on the same sections. Immunoreaction on all sections is visualized by the ABC method with DAB as the chromogen (brown) (A–L), and in addition, panels (K, L) show double immunostaining for p-cSrc on panels (H, I), respectively, by the PIC method with DAB/ NiCl$_2$ as the chromogen (blue & brown). Arrowheads, blank arrows, and solid arrows indicate VECs, macrophages, and VSMCs, respectively. Bar=50 µm (A–L). ABC, avidin-biotin-immunoperoxidase complex; DAB, 3,3'-diaminobenzidine tetrahydrochloride; MRP, macrophage-rich plaque; NF-κB, nuclear factor-κB; OxPC, oxidized phosphatidylcholine; p-cSrc, active form of cSrc phosphorylated at Y$^{416}$; p-ERK, active form of extracellular signal-regulated kinase phosphorylated at Y$^{204}$; PIC, polymer-immunocomplex; p-p38, active form of p38 mitogen-activated protein kinase phosphorylated at Y$^{182}$. 
VI. References

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