Imatinib Mesylate (STI571)-Induced Cell Edge Translocation of Kinase-Active and Kinase-Defective Abelson Kinase: Requirements of Myristoylation and src Homology 3 Domain

Akiko Fujita, Tomoyuki Shishido, Yunfeng Yuan, Eiji Inamoto, Shuh Narumiya, and Naoki Watanabe

Department of Pharmacology, Kyoto University Faculty of Medicine, Kyoto, Japan (A.F., Y.Y., E.I., S.N., N.W.); and Laboratory of Molecular Oncology, Nara Institute of Science and Technology, Nara, Japan (T.S.)

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ABSTRACT

4-[(4-Methyl-1-piperazinyl)methyl]-N-[4-methyl-3-[[4-(3-pyridinyl)-2-pyrimidinyl][amino]-phenyl]benzamide methanesulfonate (STI571) is the first successful target-based drug with excellent potency against chronic myelogenous leukemia. Studies on this compound have illuminated potentials and problems of kinase inhibitors in the treatment of cancer. As found in crystal structures, STI571-bound Abelson kinase (abl) is believed to form closed conformation with N-terminal regulatory domains. Here we present evidence of distinct STI571-induced modulation of abl functions using high-resolution live-cell imaging approaches. Within lamellipodia of fibroblast cells, STI571 was found to induce rapid translocation of abl to the lamellipodium tip. Quantitative analysis yielded 0.81 and 1.8 μM for EC_{50} values of STI571-induced cell edge translocation of abl-KD-green fluorescent protein (GFP) and wild-type abl-GFP, respectively. It also revealed adverse response of drug-resistant abl-T334I to STI571, suggesting that drug binding to abl-GFP triggers translocation. N-myristoylation and the src homology 3 (SH3) domain were required for this translocation, whereas disruption of intramolecular interactions of these motifs enhanced cell-edge association of abl. An intact C-terminal last exon region in abl, but not its F-actin binding, was required for efficient cell-edge translocation. Moreover, single-molecule observation revealed an STI571-induced rapid increase in slow diffusive species of abl in both the tip and the body region of lamellipodia. These results suggest that although activated abl translocates to the cell edge at its open state, STI571 can also bind and lock abl in the open and membrane-tethered conformation as long as the SH3 domain and the C-terminal region are intact. High-resolution imaging can be a powerful tool for elucidating inhibitor modulation of abl functions under intracellular environment.

Abl is a nonreceptor tyrosine kinase that is implicated in cellular processes such as cell-substrate adhesion, apoptosis, cell cycle regulation, and stress response (Pendergast, 2002; Suzuki and Shishido, 2007). Expression of a chimeric Ber-abl protein, which arises from a reciprocal translocation between chromosomes 9 and 22, causes chronic myelogenous leukemia (CML) (de Klein et al., 1982). STI571, also called imatinib or Gleevec, induces hematological remissions in 90% of patients in the chronic phase of CML (Kantarjian et al., 2002). However, patients receiving this drug may develop drug resistance associated with kinase domain mutations. This led to the development of novel abl kinase inhibitors with a higher potency and broader inhibition spectrums against mutated kinases (Quintas-Cardama et al., 2007). However, several mutations remain resistant to new inhibitors.

In abl, N-terminal CAP (NCAP), SH3, and SH2 domains precede the kinase domain (Woolding et al., 2003). In abl Ib, a major splice variant of c-abl in humans (type IV in mouse), the NCAP region is myristoylated. This myristoylated N-terminal peptide binds the kinase C-lobe (Hantschel et al., 2003; Nagar et al., 2003), conferring autoinhibited

ABBREVIATIONS: CML, chronic myelogenous leukemia; STI571 [imatinib mesylate; Gleevec (US) or Glivec (Europe/Australia)]; 4-[(4-methyl-1-piperazinyl)methyl]-N-[4-methyl-3-[[4-(3-pyridinyl)-2-pyrimidinyl][amino]-phenyl]benzamide methanesulfonate; Abl, Abelson kinase; Bcr, breakpoint cluster region; EGFP, enhanced green fluorescent protein; SH3, src homology 3; abl-1, Abelson kinase interactor 1; Pag, proliferation-associated gene; GFP, green fluorescent protein; NA, numerical aperture; MES, 4-morpholineethanesulfonic acid; NCAP, N-terminal CAP; PD166326, 6-[(2,6-dichlorophenyl)-2-(3-hydroxymethylphenylamino)-8-methyl-8H-pyrido[2,3-d]pyrimidine-7-one.
conformation of abl. The SH3 domain also plays an important role, forming a bridge with the linker between SH2 and kinase domains in autoinhibited abl (Barila and Superti-Furga, 1998). The SH3 domain also interacts with several ligands such as Abi-2, 3BP-1, and Pag/macrophage 23-kDa stress protein (Cicchetti et al., 1995; Dai and Pendergast, 1995; Wen and Van Etten, 1997). An in vitro random mutagenesis analysis of Bcr-abl has identified mutations that give rise to STI571 resistance not only in the kinase domain but also in these N-terminal regulatory domains (Azam et al., 2003). It is noteworthy that several residues conferring STI571 resistance in Bcr-abl are identical with the residues that convert c-abl to an oncogene (Van Etten et al., 1995; Barila and Superti-Furga, 1998; Brasher and Van Etten, 2000), suggesting a common mechanism for drug susceptibility between Bcr-abl and c-abl. Such regulatory domain mutations seem to destabilize the autoinhibited conformation (Azam et al., 2003). In crystals (Schindler et al., 2000; Nagar et al., 2003, 2006), N-terminal regulatory domains in the STI571-bound abl form a closed autoinhibited conformation. It is therefore widely accepted that STI571 stabilizes the closed conformation of the N-terminal half of abl.

With regard to the C terminus downstream of the kinase domain, its F-actin binding down-regulates the kinase activity (Woodring et al., 2001; Hantschel et al., 2005). Through proline-rich motifs between the kinase and the C-terminal F-actin binding domains, abl interacts with ab-1 and ab-2 (Dai and Pendergast, 1995; Shi et al., 1995), which bridge the association with its substrate, mammalian enabled (Tani et al., 2003). Although the structure of full-length abl has not been solved and the precise relationship between the kinase and the F-actin binding domains remains to be elucidated, the association of abl with F-actin and its regulatory proteins may have close relationship with the regulation of its kinase activity.

Cell imaging-based assays involving Ca$^{2+}$ indicators, chemiluminescence, fluorescence resonance energy transfer/bioluminescence resonance energy transfer, and phosphorylation-specific antibodies can be useful tools for drug development and drug target evaluation. They enable rapid assay prepration by eliminating steps of protein purification and development, and drug targets for evaluation. They enable rapid assay preparation by eliminating steps of protein purification and biochemical reconstitution. They may include all required components in assay systems. However, it is often problematic to distinguish whether drugs exert the effect directly or secondarily through distinct targets. For example, Crk-II-based fluorescence resonance energy transfer probes change the emission profile upon phosphorylation by either abl or epidermal growth factor receptor (Kurokawa et al., 2001; Ting et al., 2001). Here we report a more versatile method to monitor drug modulation of abl functions that is direct observation of molecular behavior of GFP-tagged abl using high-resolution fluorescence microscopy. We show that STI571 induces rapid translocation of abl to the tip of lamellipodia in fibroblast cells. We have developed several quantitative analysis methods, including single-molecule observation (Watanabe and Mitchison, 2002; Miyoshi et al., 2006). Our domain analysis within the abl structure reveals STI571-induced rapid cell-edge translocation of abl possibly through promoting complex formation with putative cellular partner(s).

Materials and Methods

Plasmids and Reagents. pcDNA3-abl and pcDNA3-ablKD encoding cDNA of murine type IV c-abl and its kinase-defective K290M mutant, abl-KD, were provided by David Baltimore. mRFP1 (Campbell et al., 2002) and mCherry cDNAs (Shaner et al., 2004) were gifts from Roger Y. Tsien. For preparation of wild-type and deletion mutants of abl, appropriate fragments of abl cDNA were amplified by polymerase chain reaction and introduced into pEGFP vectors (Clontech Laboratories, Mountain View, CA). Point mutations were introduced using QuickChange site-directed mutagenesis kit (Stratagene, La Jolla, CA). For mRFP1 fusion constructs, the coding region of EGFP was replaced with mRFP1 cDNA. The expression vector for mCherry-actin was generated by substituting mCherry cDNA for the coding sequence of EGFP in delCMV-EGFP-actin (Watanabe and Mitchison, 2002). The abl inhibitor STI571 was supplied by Novartis Pharma (Basel, Switzerland).

Live Cell Imaging and Fluorescence Single-Molecule Observation. Live cell imaging and single-molecule observation were carried out as described previously (Watanabe and Mitchison, 2002; Miyoshi et al., 2006). In brief, Xenopus laevis XTC fibroblasts were transfected using SuperFect (Qiagen, Valencia, CA) and maintained after passage into fresh flasks. Before experiments, cells were trypsinized and allowed to spread on a poly-L-lysine (1 mg/ml)-treated coverslip. Cells were then incubated in the medium with or without 1 μM STI571 for 5 min and fixed in Cytoskeleton buffer (10 mM Pipes, pH 6.1, 90 mM KCl, 3 mM MgCl$_2$, 2 mM EGTA, and 0.16 M sucrose) containing 3.7% paraformaldehyde for 20 min at room temperature. The samples were then permeabilized by 0.2% Triton X-100 in Tris-buffered saline (10 mM Tris, pH 7.5, and 150 mM NaCl). A rabbit polyclonal anti-abl antibody (Cell Signaling Technology, Danvers, MA) was used at 1:100 dilution for the primary antibody. Alexa-fluor 594 anti-rabbit antibody and Oregon Green 488 phallolidin (Invitrogen) were used to visualize the primary antibody and filamentous actin, respectively. Images were acquired using Olympus BX52 epifluorescence microscope and plan apochromatic 100×/H11003 objective equipped with a cooled charged-coupled device camera: RTE/CCD-1300-YHS, Yashica CI.512 (both from Roper Scientific, Princeton, NJ), or UIC-QE ( Molecular Devices, Sunnyvale, CA).

Immunofluorescence Analysis. XTC cells transfected with pcDNA3-abl, an expression construct encoding full-length murine type IV c-abl (Tani et al., 2003), were trypsinized and allowed to spread onto poly-L-lysine-coated glass coverslips attached to a handmade flow cell in 70% Leibovitz’s L15 medium (Invitrogen, Carlsbad, CA) without serum for 30 min. The flow cell was placed on the stage of an Olympus (Tokyo, Japan) BX52, BX51, or IX71 microscope equipped with a cooled charge-coupled device camera: RTE/CCD-1300-YHS, Yashica CI.512 (both from Roper Scientific, Princeton, NJ), or UIC-QE (Molecular Devices, Sunnyvale, CA). Immunofluorescence images were acquired at 21 to 23°C using the Metamorph software (Molecular Devices) and Olympus objectives, plan apochromatic 100×/NA, 1.40 or universal apochromatic 150×/NA, 1.45 total internal reflection fluorescence microscope (TIRFM; NA, 1.45), up to 120 min after cells were seeded. For fluorescence single-molecule speckle microscopy, illumination was restricted to the cell peripheral areas. Single-molecule counting was carried out manually.

Materials and Methods

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Results

STI571 Induces Cell-Edge Translocation of abl-GFP Probes. We fused type IV mouse abl with fluorescent proteins (EGFP or mRFP1) at its C terminus and expressed it in X. laevis XTC fibroblasts. These cells adhere tightly to poly-L-lysine-coated glass coverslips and form wide flat lamellipodia (Watanabe and Mitchison, 2002). Placing the cell periphery in a flat geometry enables precise microscopic observations of spatiotemporal regulation of cytoskeletal rearrangement and molecular behavior (Nakagawa et al., 2001;
The merits of our model system also include the ease of cell culture under the microscope because cells grow at room temperature without supply of CO$_2$. We focused on the behavior of abl-EGFP at the cell periphery and found its association with the actin network throughout lamellipodia in XTC cells (Fig. 1A). In lamellipodia, the tip region was the most intensely labeled. These observations are consistent at ASPET Journals on April 14, 2017 molpharm.aspetjournals.org Downloaded from
with the localization of endogenous abl (Woodring et al., 2002; Jin and Wang, 2007). When overexpressed, abl-EGFP induced the formation of finger-shaped actin-rich structures where abl-EGFP was associated. These abl-associated structures dynamically moved and often extended toward the cell periphery against the retrograde actin flow (supplementary Fig. 1 and Movie 1), which fluxes inward at ~1.5 μm/min (Watanabe and Mitchison, 2002). Previous studies have reported analogous effects of abl in the promotion of microspike formation (Woodring et al., 2002) and elongation of the actin comet tail induced by Shigella flexneri (Burton et al., 2005).

Only the negligible effect of STI571 on the localization of abl has been reported previously (Hantschel et al., 2005). By observing abl-EGFP in wide flat lamellipodia, however, we noticed a rapid increase in abl-EGFP signals at the lamellipodium tip as well as at dots in the lamellae region upon perfusion of STI571 (Fig. 1B, Supplemental Movie 2). Simultaneously, a loss in abl-EGFP signals associated with the F-actin network was also observed. These observations suggest a decrease in the affinity between abl and F-actin and the increased affinity of abl to putative partner(s) at the lamellipodium tip induced by STI571.

We also verified that untagged full-length abl translocated to the cell periphery in response to STI571 (Fig. 1C). These results exclude the possibility that the GFP tag attached to the C terminus of abl caused an abnormal response to STI571 by altering the open/closed conformational regulation of abl kinase.

Overexpression of abl-EGFP impaired the spreading of XTC cells. STI571 relieved this inhibition and induced cell-edge protrusion (Fig. 1B, bottom). These observations are consistent with the impaired spreading of NIH3T3 cells by induced dimerization of abl-FKBP (Jin and Wang, 2007). The edge of nontransfected XTC cells protruded less markedly in response to STI571 (data not shown). These results suggest that overexpressed abl negatively regulates cell-edge extension through excessive phosphorylation of its substrates. We therefore wondered whether STI571-induced translocation of abl occurred as a result of dephosphorylation of its substrates. To test this, we generated kinase-dead abl probes, abl-KD-EGFP and abl-KD-mRFP1, in which Lys290 was substituted to methionine. In contrast to wild type, abl-KD-EGFP did not interfere with cell spreading. Only marginal lamellipodium extension was induced by STI571 treatment in cells expressing abl-KD constructs as in control cells, suggesting a low level of phosphorylation of abl substrates in abl-KD-expressing cells (Tani et al., 2003). Upon perfusion of STI571, abl-KD rapidly translocated to the lamellipodium tip and cytoplasmic actin-rich foci to a similar degree as wild-type abl (Fig. 1D). Therefore, dephosphorylation of abl substrates is unlikely to be the mechanism for its STI571-induced translocation.

We next verified that STI571-induced translocation is not caused by a change in F-actin morphology. Translocation of abl-KD occurred without substantial F-actin redistribution (Fig. 1E, supplemental Movie 3). We also confirmed that the accumulation of abl-KD-EGFP was not due to an increase in the thickness of the cell edge (Fig. 1F).

Dose-Response Analysis of STI571-Induced Cell Edge Translocation of abl. Figure 2A shows dose-response analysis of STI571-induced translocation of abl-KD. We measured the ratio of the EGFP fluorescence intensity between the edge and the body region (red). The graph shows the ratio of the average fluorescence intensity of abl-KD-EGFP between the edge area and the body area (n = 6 cells). Error bars show S.D. The dose-response curve (inset) yields 0.81 μM for the EC50 value.

Figure 2A shows dose-response analysis of STI571-induced translocation of abl using a computer-assisted method. A, cells expressing abl-KD-EGFP were treated with increasing doses of STI571. Right, abl-KD-EGFP (green) and the measured area automatically assigned for the lamellipodium edge region (blue) and the body region (red). The graph shows the ratio of the average fluorescence intensity of abl-KD-EGFP between the edge area and the body area (n = 6 cells). Error bars show S.D. The dose-response curve (inset) yields 0.81 μM for the EC50 value. B, dose-response analysis of STI571-induced cell edge translocation of wild-type abl-EGFP measured as in A, except that cells were treated with single-dose STI571 (n = 3–4 cells for each STI571 concentration). Error bars show S.D. The dose-response curve for the fold increase between before and 5 min in edge/body fluorescence ratio (inset) yields 1.8 μM for the EC50 value. C, a tyrosine to phenylalanine mutation at Tyr412 in wild-type abl, the phosphorylation of which impairs STI571 binding (Schindler et al., 2000), increased sensitivity to low-dose STI571 treatment. The graph shows the edge/body fluorescence ratio in a cell treated with 0.5 μM STI571. Only abl-Y412F shows the gradual increase in the ratio. The EC50 value of STI571-induced cell-edge translocation of abl-Y412F-EGFP measured as in B was 1.1 μM. Scale bars, 5 μm.
the body areas in lamellipodia (supplemental Movies 4 and 5) by building a computer-aided quantification method. This method combines image manipulation and measurement commands in the MetaMorph software. This nonbiased analysis yielded 0.81 μM for the EC_{50} value of STI571-induced translocation of abl-KD. This EC_{50} value is comparable with the IC_{50} value of STI571 for the inhibition of abl in cell-based assays, 0.25 to 0.8 μM (Druker et al., 1996; Roumiantsev et al., 2002; Azam et al., 2003; Quintas-Cardama et al., 2007).

We also carried out the same analysis on wild-type abl-EGFP (Fig. 2B). The EC_{50} value of STI571-induced cell-edge translocation of wild-type abl was 1.8 μM, which is slightly higher than that of abl-KD. Especially at lower concentrations, 0.5 to 5 μM, STI571-induced cell-edge translocation of wild-type abl was less marked than abl-KD. Phosphorylation at Tyr412 within the activation loop impairs STI571 binding to abl kinase with an increase in the K_{i} value from 37 nM to 7 μM (Schindler et al., 2000). The importance of this phosphorylation has also been noted by the fact that mutation at Pro131 confers STI571 resistance in cells (Azam et al., 2003) although the P131L mutation renders the kinase more sensitive rather than more resistant to STI-571 in its unphosphorylated state (Roumiantsev et al., 2002). We therefore speculated that autophosphorylation at Tyr412 (Brasher and Van Etten, 2000; Dorey et al., 2001) might reduce the sensitivity of wild-type abl to low-dose STI571. To test this, we introduced Y412F mutation in wild-type abl-EGFP. Abl-Y412F showed better response to low-dose STI571 than wild-type abl (Fig. 2C). The EC_{50} value of STI571-induced cell-edge translocation of abl-Y412F measured as in Fig. 2B was 1.1 μM. These data are in agreement with our interpretation that phosphorylation at Tyr412 could reduce the sensitivity of abl-EGFP to low-dose STI571.

**Binding of STI571 Probes Triggers Cell-Edge Translocation of abl.** The good correspondence between the EC_{50} value of STI571-induced abl translocation and the known IC_{50} value of inhibition of abl by STI571 in cell-based assays supports our interpretation that drug binding drives the translocation of abl probes. To further test whether drug binding to abl probes is the key to STI571-induced cell-edge translocation, we observed the response of the STI571-resistant mutant, abl-T334I, to STI571. T334I mutation, T335I in the abl 1a variant, has been detected frequently among patients with CML who developed drug-resistant disease. Thr334 directly interacts with STI571 in the drug binding pocket of abl (Schindler et al., 2000) and is called the gatekeeper, because T334I shows resistance to most of the kinase inhibitors, including STI571 (Quintas-Cardama et al., 2007). We found that abl-T334I did not translocate to the leading edge in response to STI571 treatment. Moreover, abl-KD-T334I-EGFP associated with the lamellipodium tip seemed to be displaced by coexpressed abl-KD-mRFP1 upon STI571 treatment (Fig. 3, supplemental Movie 6). We observed similar adverse effects of STI571 on the cell-edge localization between wild-type kinase variants abl-T334I and abl (data not shown). In vitro, T334I shows no significant inhibition at STI571 concentrations 200-fold greater than the IC_{50} value of wild-type abl (Corbin et al., 2002). In the following comparison of the translocation efficiency between various abl mutants, we used a high concentration of STI571, ~50 μM, to ensure drug binding to abl mutants, some of which are less sensitive to STI571 than wild-type abl. Among those tested, T334I is the only mutant to be resistant to STI571 at 50 μM. In cells coexpressing constructs with and without the T334I mutation, only the wild-type variant presumably binds STI571 and increases its affinity to the partner(s) at the leading edge. This could account for the displacement of T334I from the cell edge upon STI571 treatment. We conclude that drug binding to abl triggers its cell-edge translocation.

**Release of N-Myristoylated and the SH3 Domain from Autoinhibition Promote Cell-Edge Translocation of abl.** Given the STI571 binding-induced translocation of abl probes, we were motivated to determine which structures in abl are required for translocation. The effects of various deletions and mutations were validated by direct comparison against coexpressed full-length abl because the degree of STI571-induced translocation varied between cells. We tested deletions and mutations introduced into both wild-type abl and abl-KD, which gave identical results (see below). We used a saturating concentration of STI571, 10 to 50 μM, to ensure drug binding to abl mutants, some of which are less sensitive to STI571 than wild type. First, we coexpressed abl-KD-mRFP1 with abl-KD-EGFP and confirmed identical behavior and response to STI571 between two constructs (Fig. 4A). The deletion of the NCAP domain severely impaired STI571-induced cell-edge translocation of abl-KD (Fig. 4B). In abl Ib, a major splice variant of c-abl in human (type IV in mouse), the NCAP region is myristoylated. The myristoylated N-terminal peptide binds the kinase C-lobe (Hantschel et al., 2003; Nagar et al., 2003). We therefore tested the G2A mutant in which glycine at position 2, the target residue for N-myristoylation, had been mutated to alanine. G2A was defective in STI571-induced cell-edge translocation (Fig. 4C, supplemental Movie 7). It is noteworthy that both ΔNCAP and G2A were still able to associate with F-actin structures in lamellipodia.

We next tested the A356N mutation in the binding pocket for the myristoyl group. A356N exhibits high kinase activity as a result of defective autoinhibition (Hantschel et al., 2003).
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A  abl-KD-mRFP1 abl-KD-GFP

B  abl-KD-mRFP1 ΔNCAP-abl-KD-GFP

C  abl-mRFP1 abl-G2A-GFP

D  abl-mRFP1 abl-A356N-GFP

E  abl-KD-mRFP1 abl-KD-P131L-GFP

F  abl-mRFP1 abl-PP-GFP

G  abl-KD-mRFP1 abl-KD-ΔFB-GFP

H  abl-KD-mRFP1 abl-KD-ΔC-GFP

I  abl-WT abl-KD ΔNCAP G2A A356N P131L PP ΔFB ΔC T334I

myc  NCAP SH3 SH2 Kinase  Pro-rich  F-actin binding

1142 aa  STI571-induced translocation  + + +

abl-WT  K290M + + +
abl-KD  + + +
ΔNCAP  b1-1142 WT or K290M + + +
G2A  ΔG2A + + +
A356N  ΔA356N + + +
P131L  WT or K290M ± + +
PP  ΔPP WT or K290M ± + +
ΔFB  1-1110 + - +
ΔC  1-519 ± - -
T334I  WT or K290M T334I ± - +

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We found that A356N is constitutively associated to the cell edge even before STI571 treatment, whereas abl-mRFP1 localized to the cell edge only after STI571 perfusion (Fig. 4D, supplemental Movie 8). These data raise a possibility that the myristoyl group released from the kinase domain contributes to the association of abl to the leading edge.

The deletion and point mutations in the SH3 domain have been studied most extensively as activating mutations (Jackson and Baltimore, 1989; Van Etten et al., 1995; Barila and Superti-Furga, 1998). We tested two point mutations, P131L and P242E/P249E, within the SH3 domain and the SH2-kinase linker region, respectively. Both mutant kinases are highly active as a result of impaired autoinhibitory interactions between the two regions. Ab1-KD-P131L translocated to the cell edge less efficiently upon STI571 treatment than abl-KD (Fig. 4E). In marked contrast, the P242E/P249E mutant (abl-PP) was constitutively associated to the cell edge before and after STI571 treatment (Fig. 4F, supplemental Movie 9). The P131L mutation may disrupt the association to the SH3 domain ligands such as Abi-2, 3BP-1, and Pag/macrophage 23-kDa stress protein (Cicchetti et al., 1995; Dai and Pendergast, 1995; Wen and Van Etten, 1997). These results indicate that release from the intramolecular interaction and subsequent ligand binding of the SH3 domain promote the cell-edge translocation of abl.

**An Intact Last Exon Region but Not Its F-Actin Binding Is Required for Translocation.** We next tested the motifs downstream of the kinase domain in abl. We observed the behavior of abl-ΔFB-KD, in which C-terminal 32 amino acids are deleted. Deletion of this portion eliminates F-actin binding (Hantschel et al., 2005). abl-ΔFB consistently showed diffuse cytoplasmic localization. Upon perfusion of STI571, abl-ΔFB-KD translocated to the cell edge to the same extent as full-length abl-KD (Fig. 4G, supplemental Fig. 2, Movie 10). Thus the cell-edge translocation of abl does not require its ability to bind F-actin. This finding is consistent with the decrease in abl or abl-KD signals associated with the actin network upon STI571 treatment (Fig. 1, A and D). Further shortening of the C terminus, which eliminated the entire C-terminal last exon region, did not abolish but substantially reduced STI571-induced translocation (Fig. 4H, supplemental Movie 11). The deleted region includes multiple proline-rich motifs that bind abl ligands such as v-crk sarcoma virus CT10 oncogene homolog, Nck, and abl-1 (Woodring et al., 2005). Taken together, our data (Fig. 4I) indicate that binding of STI571 to abl induces its cell-edge translocation by promoting the complex formation with putative partner(s) through the myristoyl group, the SH3 domain, and the last exon region.

**A Rapid Induction of Slow Diffusive ab1 by STI571 Revealed by Single-Molecule Observation.** Single-molecule fluorescence microscopy was also useful and elucidated another aspect of STI571-modulated molecular behavior of abl. In lamellipodia, abl-KD-EGFP visualized as single molecules at 20 frames per second exhibited two distinct kinetics of dissociation from cellular structures (Fig. 5A). Both species did not show any directional movement. Because our microscopy would detect directional movement of molecules (Higashida et al., 2004) at up to ~10 μm/s under the setting, active transport of abl is unlikely. The slow diffusive species were more predominant at the lamellipodium tip than in the rest of lamellipodia. Our previous studies (Watanabe and Mitchison, 2002; Miyoshi et al., 2006) have used single-molecule “speckle” microscopy to monitor cytoskeletal association of molecules because immobilization of probes is the key for the formation of discrete speckle signals. We applied the same strategy to distinguish two species. When acquired using the slower exposure time, only a fraction of abl, which mainly consisted of slow diffusive species, were visualized (Fig. 5B, supplemental Movie 12). Images acquired with the 1500-ms exposure revealed that STI571 rapidly increased the slow diffusive abl-KD, not only at the tip but also in the body of lamellipodia (Fig. 5C). The number of single-molecule abl-KD-EGFP (Fig. 5C) and abl-EGFP (Fig. 5D) visualized with the slower exposure times (800–1500 ms) increased significantly after perfusion of STI571. Moreover, single-molecule observation revealed the motion of slow diffusive abl near the cell periphery, which seems to largely represent random diffusive processes (Fig. 5E, supplemental Movie 13). Together with the critical role of N-myristoylation (Fig. 4, C and D), STI571-bound abl most likely accumulates to the lamellipodium tip by codiffusing with putative partners through multiple interactions, including the myristoyl group-membrane interaction, but not simply being targeted to the pre-existing partners at the lamellipodium tip, such as abl-1. Extensive biochemical and optical analysis on such diffusive motions of abl is required to elucidate the underlying mechanisms.

**Discussion**

Using high-resolution live-cell imaging approaches, our present study revealed rapid cell-edge translocation of abl...
kinase induced by its inhibitor, STI571, in lamellipodia of XTC fibroblasts. Several lines of evidence indicate that binding of STI571, but not other indirect mechanisms such as dephosphorylation of abl substrates, induces translocation of abl probes. First, the kinase-dead mutant of abl (K290M) showed similar translocation in response to STI571. Second, the EC$_{50}$ value of STI571-induced cell-edge translocation of abl is comparable with the known IC$_{50}$ value of this drug in other abl-dependent cellular processes. Third, cell-edge association of the drug-resistant abl-T334I mutant was diminished upon STI571 treatment, presumably displaced by STI571-bound wild-type variants. For efficient translocation, multiple abl structures such as N-myristoylation, the SH3 domain, and the last exon region are required. Contrary to the currently prevailing view that STI571 stabilizes the autoinhibited conformation of abl (Nagar et al., 2003; Wang, 2004), our domain analysis (Fig. 4) suggests that STI571 may promote complex formation with putative partner(s) of abl. The presence of two populations of abl with distinct diffusive behaviors before STI571 treatment (Fig. 5, A and B) suggests that the conformation of abl in cells is at dynamic equilibrium (Fig. 6). We designate the open, partner-associated state of abl as the “coassembled” state. Based on our domain analysis, we postulate that upon binding to STI571, this equilibrium shifts to the coassembled state in which the myristoyl group and the SH3 domain are disassembled from autoinhibitory interactions. Released myristoyl group and the SH3 domain together with the last exon region may bridge the interactions with putative cellular partner(s) of abl. Such a complex presumably interacts with the plasma membrane.

Fig. 5. Rapid induction of slow diffusive species of abl by STI571 revealed by single-molecule imaging. A, the graphs show the decay in the number of single-molecule abl-KD-EGFP speckles identified at the first frame of time-lapse images acquired at 50-ms intervals. Molecules associated to the cell contour (lamellipodium tip; magenta) were analyzed separately. Cells treated with 50 μM STI571 for 5 to 15 min were analyzed in the right graphs. Colored lines show the data after normalization for photobleaching. Dashed lines indicate the double exponential fit to the data. The best fit was obtained as follows: before treatment, $t_{1/2}$ = 51 ms (62%) and $t_{1/2}$ = 650 ms (38%) at the lamellipodium tip; $t_{1/2}$ = 58 ms (94%) and $t_{1/2}$ = 1700 ms (5.9%) in the body; after treatment: $t_{1/2}$ = 67 ms (50%) and $t_{1/2}$ = 560 ms (50%) at the lamellipodium tip; $t_{1/2}$ = 64 ms (85%) and $t_{1/2}$ = 660 ms (15%) in the body; n = 5 cells for each condition. B, image of XTC cells expressing a low level of abl-KD-EGFP were acquired by varying exposure times. On the right, illumination was attenuated to 12.5% of the left. Note that only a fraction of abl-KD speckles were visualized as a discrete spot (marked by magenta circles) with the 1500-ms exposure. C and C’, the number of abl-KD-EGFP speckles visualized with the 1500-ms exposure time (marked by magenta circles) increased rapidly upon varying exposure times. On the right, illumination was attenuated to ~12.5% of the left. Note that only a fraction of abl-KD speckles were visualized as a discrete spot (marked by magenta circles) with the 1500-ms exposure. C and C’, the number of abl-KD-EGFP speckles visualized with the 1500-ms exposure time (marked by magenta circles) increased rapidly upon varying exposure times. On the right, illumination was attenuated to ~12.5% of the left. Note that only a fraction of abl-KD speckles were visualized as a discrete spot (marked by magenta circles) with the 1500-ms exposure. D, the number of abl-KD-EGFP speckles visualized with the 800-ms exposure time increased 2 min after 5 μM STI571 perfusion. Each color in the graph indicates data from the same cell (n = 6 cells). **, p = 0.002, two-tailed paired t test. E, images of single-molecule abl-KD-EGFP speckles in the presence of 10 μM STI571 were panelled at 100-ms intervals. Note the slow diffusive motion of an abl-KD speckle (arrowheads) traveling in the cell-edge region (see also supplemental Movie 13). Scale bars, 5 μm.
membrane through the N-myristate attached to abl. STI571-bound abl may then accumulate to the cell periphery by slow diffusive motion. In protein kinase A, binding of the regulatory RI subunit switches the catalytic subunit to expose the N-myristate attached to the catalytic subunit (Gangal et al., 1999). To our knowledge, our finding is the first example of inhibitor-induced conformational regulation of kinases. This STI571-induced inside-out conformation regulation presumably requires abl-interacting molecule(s).

In crystals of STI571-bound abl, the activation loop is stabilized in an inactive conformation (Schindler et al., 2000). This unique conformation is believed to explain the specificity of STI571. A recent report, however, has shown that the kinase domain of abl in complex with an ATP-peptide conjugate adopts a different inactive conformation, which closely resembles that of src kinases (Levinson et al., 2006). These observations infer flexible nature of the catalytic core of the kinase and its modulation by interacting molecules. The binding of the myristoyl group also induces a conformation change in the kinase C lobe, causing the C-terminal helix to undergo a \( \sim 90^\circ \) bend (Nagar et al., 2003). Currently, it is unknown whether STI571 affects the interaction between the kinase and regulatory domains (Nagar et al., 2003). In this regard, it is interesting to note that Nagar et al. (2003 and 2006) described crystals of myristoylated abl\(^1\)–\(^531\) in complex with another inhibitor, PD166326, which consisted of two crystallographically independent molecules. One was in the inactive state with the SH3–SH2 unit docked to the kinase domain, and the other was in an elongated conformation with the myristoyl group no longer associating to the kinase domain. This example shows possible conformation fluctuation of inhibitor-bound abl between closed and coassembled states. The X-ray-scattering analysis (Nagar et al., 2006) has indicated that the majority of inhibitor-bound abl is in the open state. Our findings suggest that in the presence of cellular factors, STI571 promotes the formation of a complex between abl and partner(s).

Our data show translocation of STI571-bound abl to the lamellipodium tip and the pivotal role of the SH3 domain and the last exon region in this translocation. Abi-1 and abi-2 may be good candidates for the partner of STI571-bound abl. Abi-1 interacts with the SH3 domain and the last exon region of abl. Abi-1 is localized at the lamellipodium tip (Nakagawa et al., 2001) as a component of a protein complex consisting of WAVE/Scar1/2/3, specifically Rac-associated protein 1/PIR121, Nck-associated protein 1, and Hspc300 (Eden et al., 2002). This complex mediates signals from GTP-bound Rac1 and induces nucleation of the dendritic F-actin network in the lamellipodium tip. Abi-1/2 also plays an important role in mediating signaling downstream of abl. Abi-1 promotes tyrosine phosphorylation of WAVE2 (Leng et al., 2005) and mammalian enabled (Tani et al., 2003) by bridging the interaction with abl. In hematopoietic cell lines, Bcr-abl induces clustering of an actin-rich structure containing abi-1, WAVE2, and b1-integrin, whereas mutations in Bcr-abl that impairs interaction with abi-1/2 abrogate this cluster formation (Li et al., 2007). The ability for this integrin clustering is correlated with the accumulation of Bcr-abl-transformed myeloid cells in the spleen (Dai et al., 2001). From these properties, Abi-1/2 may be referred to as a coactivator of abl kinase (Suzuki and Shishido, 2007). Because STI571 abolishes such adhesion clusters (Li et al., 2007), Bcr-abl-induced adhesion clusters in hematopoietic cells seem to differ from the putative protein complex of STI571-bound abl in XTC cells. On the other hand, abl interactors such as Pag (Wen and Van Etten, 1997), retinoblastoma protein (Welch and Wang, 1993), and F-actin (Woodring et al., 2003) inhibit the kinase activity of abl. Based on these findings, a coactivation mechanism has been proposed for cell factor regulation of abl kinase activity (Wang, 2004). Our current findings have now opened up another possibility of a two-component inhibition mechanism by which cellular cofactor(s) may play a role in the kinase inhibitor modulation of abl functions. To fully understand this mode of cofactor modulation of kinase inhibition, identification of partner(s) for STI571-bound abl and biochemical reconstitution combining abl, kinase inhibitors, and such putative partner(s) both in cells and in vitro are required.

**Prospects and Implications.** The regulation of abl has been studied extensively both biochemically and structurally. Although such analyses have provided a tremendous amount of insight, they were unable to detect our current findings, because it will require reconstitution using full-length abl proteins together with putative cellular partner(s). Our current findings imply a possible involvement of such cellular components in the kinase inhibition of abl. All of the quantification methods, edge/body fluorescence ratio measurement (Fig. 2), comparison against red fluorescent protein-tagged full-length variants (Fig. 4), and single-molecule counting (Fig. 5, B–D), can be developed into automatic analysis suited to high-content drug screening and target evaluation. Our study highlights that direct observation of molec-
ular behavior of abl itself, and its variants could elucidate the mechanism of action of inhibitors and the influence of mutations readily under live-cell environment.

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References


Address correspondence to: Dr. Naoki Watanabe, Department of Pharmacology, Kyoto University Faculty of Medicine, Yoshida Konoe-cho, Sakyoku, Kyoto, Japan 606-8501. E-mail: naoki-w@four.med.kyoto-u.ac.jp