Structural basis of inactivation of Ras and Rap1 small GTPases by Ras/Rap1-specific endopeptidase from the sepsis-causing pathogen Vibrio vulnificus

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Multifunctional autoprocessing repeats-in-toxin (MARTX) toxins are secreted by Gram-negative bacteria and function as primary virulence-promoting macromolecules that deliver multiple cytopathic and cytotoxic effector domains into the host cytoplasm. Among these effectors, Ras/Rap1-specific endopeptidase (RRSP) catalyzes the sequence-specific cleavage of the Switch I region of the cellular substrates Ras and Rap1 that are crucial for host innate immune defenses during infection. To dissect the molecular basis underpinning RRSP-mediated substrate inactivation, we determined the crystal structure of an RRSP from the sepsis-causing bacterial pathogen Vibrio vulnificus (VvRRSP). Structural and biochemical analyses revealed that VvRRSP is a metal-independent TIKI family endopeptidase composed of an N-terminal membrane-localization and substrate-recruitment domain (N lobe) connected via an inter-lobe linker to the C-terminal active site–coordinating core–substrate-recruitment domain (N lobe) via a functional conformation of the C lobe to facilitate recruitment of cellular substrates. Collectively, these results indicate that RRSP is a critical virulence factor that robustly inactivates Ras and Rap1 and augments the pathogenicity of invading bacteria via the combined effects of its N and C lobes.

Numerous Gram-negative bacterial pathogens have the ability to deliver bacterial “effector” proteins into the host cytoplasm to modulate a multitude of host cell functions (1). Direct effector delivery systems, known as type III (2, 3), IV (4, 5), and VI (6, 7) secretion systems, play a central role in pathogenic interactions between bacteria and host cells. In addition to the typical effector transfer systems, many Gram-negative pathogenic bacteria, including the sepsis-causing pathogen Vibrio vulnificus, secrete multifunctional autoprocessing repeats-in-toxin (MARTX) toxins via the atypical type 1 secretion system (8, 9). Once secreted, these toxins translocate their effector domains modularly into host cells and undergo autoproteolysis, resulting in the release of functionally discrete effectors in the cytosol (10, 11). MARTX toxins are believed to form pores in the host cell plasma membrane via repeat-containing regions located at the N and C termini that facilitate effector module translocation, although how the repeat regions form pores remains unknown (10–12). Autoproteolysis essential for MARTX toxin function is mediated by a core element of the toxins, namely the cysteine protease domain that is allosterically activated by inositol hexakisphosphate exclusively present in the host cytosol (13).

Within host cells, different effectors display distinct cytopathicity or cytotoxicity, and the overall toxicity depends on the combined virulence of individual MARTX toxin effectors (12, 13).
Because pathogenic bacteria confer variation on the effector domain content of MARTX toxins by homologous recombination events, MARTX toxin-mediated translocation of novel effector domains emerges spontaneously (15, 16). Thus, characterizing the cellular functions of individual effector domains can provide a comprehensive understanding of how MARTX toxins contribute to bacterial pathogenesis.

To date, more than 10 MARTX effector domains have been annotated, including several domains of unknown function in Vibrio species (17), among which the fifth domain of a MARTX toxin from Vibrio vulnificus CMCP6 is a Ras/Rap1-specific endopeptidase (RRSP) (18). Notably, bacterial strains that produce a MARTX toxin harboring an RRSP are more virulent than those that do not (15). RRSP catalyzes sequence-specific cleavage of the Switch I region of the cellular substrates Ras and Rap1 (3672–3855) harboring residues Asp-3721 and Arg-3841 of four helices (26). The cytotoxicity of the RRSP from V. vulnificus CMCP6 depends on the C-terminal subdomain (residues 3672–3855) harboring residues Asp-3721 and Arg-3841 that are critical for maintaining the structural stability of the RRSP (24). Although these results revealed that RRSP hydrolyzes the critical cellular substrates Ras and Rap1, the molecular mechanisms by which it recognizes and processes cellular substrates remain unknown.

In this study, we determined the crystal structure of an RRSP from the sepsis-causing pathogen V. vulnificus CMCP6 (VvRRSP), and we demonstrated that it is a metal-independent TIKI family endopeptidase. We further dissected the structural features, identified catalytic and substrate recognition residues, and unveiled the roles of the subdomains in the functions of the enzyme.

Results

Crystal structure of VvRRSP

A previous study reported that the MARTX toxin effector RRSP is an enzyme possessing site-specific processing activity against the small GTPases Ras and Rap1 that act as cellular switches during signal transduction, including innate immune responses (18). Computational protein sequence analysis revealed that RRSP belongs to the TIKI superfamily (22). To elucidate the structural basis of Ras and Rap1 processing by RRSP, we attempted crystallization of the full-length domain, corresponding to amino acid residues 3580–4089 of the MARTX toxin from V. vulnificus strain CMCP6 (Fig. 1A). However, the resulting crystals failed to diffract to high resolution. Deletion of N- and C-terminal flexible regions resulted in a core construct (residues 3596–4072) that produced crystals diffracting X-rays to 2.7 Å (Table 1), with four VvRRSP molecules in the asymmetric unit (Fig. 1B; see under “Experimental procedures” for detailed information on structure determination and refinement). The structure of each of the four protein chains is essentially identical, except for the N-terminal α1 helix spanning residues 3596–3609 (Fig. 1C). Size-exclusion chromatography revealed that VvRRSP exists as a monomer in solution (Fig. 1D); hence, the structure of chain A was considered representative of the subunit structure and was used in further analysis. The enzyme activity of VvRRSP (residues 3596–4072) was confirmed by in vitro KRas cleavage assay, demonstrating that the construct was functional and processed Ras and Rap1 (Fig. 1E).

Molecular details of VvRRSP

The VvRRSP structure can be divided into an N-terminal lobe (N lobe, residues 3596–3722) and a C-terminal lobe (C lobe, residues 3740–4072; Fig. 2A), and the N lobe can be further subdivided into the membrane localization N1 domain (MLD), consisting of four helices (α1–α4), as characterized previously (23), and the N2 domain comprising three helices (α5–α7). The MLD resembles a pointing finger in which the first α-helix extends from the remaining three-helix bundle (Fig. 2A and Fig. S1A). This bundle (α2, α3, and α4) forms a hydrophobic core, in which Tyr-3613, Ile-3616, Leu-3620, Val-3623, Ile-3632, Leu-3638, Ile-3642, Tyr-3645, Leu-3658, Leu-3661, Val-3665, and Leu-3669 are arranged to maintain structural integrity (Fig. 2B). The structural arrangement of the MLD in VvRRSP is markedly different from that of the two structures of isolated MLDs determined previously by nuclear magnetic resonance (NMR; PDB code 2N9W; Fig. S1B) (26) and crystallography (PDB code 4ERR; Fig. S1C), suggesting that this domain may be highly flexible. The VvRRSP MLD is similar to the membrane localization C1 domain in the Pasteurella multocida toxin (PMT) (23), in which the first α-helix of the MLD differs but the remaining three helices superimpose well in the two protein structures (Fig. 2C). Residues Phe-3636 on the α3 helix and Phe-3670 on the α4 helix make critical contributions to the structural stabilization of the MLD by engaging in extensive hydrophobic contacts with the groove lined by Met-3674, Gln-3678, Asp-3682, Ala-3685, Pro-3689, Ala-3693, and Phe-3696 on the α5 and α6 helices of the N2 domain (Fig. 2D).

The C lobe can also be separated into two subdomains (Fig. 2A), the C1 domain (α8–α12 and β1–β3), and the C2 domain (α13–α22 and β4–β11) that adopts a typical α + β TIKI fold (Fig. S2A). The C lobe is firmly integrated with the N lobe, mainly via hydrophobic interactions between Leu-3710, Met-3713, and Trp-3717 on α7 in the N2 domain; Ile-3845, Ile-3848, and Tyr-3849 on α12 in the C1 domain; His-3996, Met-3999, Tyr-4000, and Gly-4003 on α20; and Gly-4039 and Val-4041 on β9 in the C2 domain (Fig. 2E). A previous study reported that Asp-3721 and Arg-3841 are critical for structural stability of RRSP (24). The structure of RRSP reveals that Asp-3721 on α7 forms a salt bridge with Arg-3841 on α12 that helps to maintain the structure between N2 and C1 domains (Fig. 2E).

A long loop (residues 3723–3739, referred to as the interlobe linker) connects the two lobes (Fig. 2F). In particular, residues Tyr-3723 and Leu-3724 play a critical role in maintaining structural integrity between the N2, C1, and C2 domains (Fig. 2F).
The inter-lobe linker appears to function as a hinge to bestow flexibility between the two lobes. To evaluate the importance of the loop, residues Tyr-3723 and Leu-3724 were mutated to alanine, and in vitro KRas cleavage assays revealed that RRSP enzyme activity was completely abolished (Fig. 2G). Thus, these two residues might be essential for controlling flexibility.

Structural comparison using PDBeFold identified structural similarity between V. vulnificus RRSP and TIKI superfamily proteins (22), including PMT (PDB code 2EBF; RMSD = 2.79 Å for 404 Cα atoms), HopBA1 (PDB code 5T09; RMSD = 2.96 Å for 146 Cα atoms), Bcr135 (PDB code 3B55; RMSD = 2.47 Å for 140 Cα atoms), and ChaN (PDB code 2G5G; RMSD = 2.95 Å for 145 Cα atoms), all of which contain a parallel β-sheet core surrounded by α-helices (Fig. S2).

C2 catalytic domain

TIKI superfamily proteins share a common core domain consisting of a central β-sheet surrounded by α-helices (22). Likewise, the C2 domain in VvRRSP has a six-stranded β-sheet, flanked by helices on both sides (Fig. S2A). Five of the six β-strands of the C2 domain (β4, β5, β6, β7, and β10) form the characteristic parallel β-sheet core observed in all TIKI proteins (Fig. S2, B–F).

It has been proposed that enzymatically active TIKI family proteins share a catalytic mechanism with the erythromycin esterase (Ere) family, in which a conserved His–Glu pair is essential for substrate hydrolysis (27). Structural comparison of V. vulnificus RRSP with the Ere family protein Bcr135 revealed that functional residues are highly conserved (Fig. 3, A–E). The predicted functional residues His-3902, His-4030, Glu-3900, and Glu-3930 (forming the 2His/2Glu motif) in VvRRSP are located near β4, β5, and β7 strands within the parallel β-sheet core of the C2 domain (Fig. 3, B and C, and Fig. S3). We subsequently investigated whether these 2His/2Glu residues are essential for catalytic activity (Fig. 3F). Mutant proteins did not undergo any significant conformational changes compared with WT RRSP according to the results of circular dichroism (CD) spectrometry analysis (Fig. S4).

We further evaluated the importance of the 2His/2Glu residues for the hydrolysis of endogenous Ras (Fig. 4A) or Rap1 (Fig. S5) in HEK293T cells transfected with each VvRRSP.
mutant. The results clearly showed that mutation of Glu-3930 or His-4030 completely abolished enzymatic activity, consistent with the results of in vitro KRas cleavage assays. Interestingly, mutation of Glu-3900 or His-3902 inhibited enzymatic activity significantly, but not completely, in cells (Fig. 4A). Consequently, ectopic expression of RRSP\textsubscript{E3930L/H4030L} for 16 h did not cause the cell-rounding that was observed with wildtype (WT) RRSP (Fig. 4B), as typically observed in RRSP-expressing cells (24). Mild cell-rounding was observed following RRSP\textsubscript{E3900L} or RRSP\textsubscript{E3902L} expression (Fig. 4B).

Green fluorescent protein (GFP)-fused MLD of RRSP localizes to the plasma membrane in HeLa cells (23), and we expected similar localization for V\textsubscript{v}RRSP in this study. We therefore tested whether the V\textsubscript{v}RRSP mutations affected cellular localization, and no difference was found between WT and mutant proteins at 16 h post-transfection; all proteins were mainly localized to the HEK293T cell membrane, where their substrates reside (Fig. S6). We further evaluated the ability of the mutants to cleave endogenous Ras at 16 h post-transfection by confocal microscopy analysis, and WT RRSP efficiently processed endogenous Ras, whereas mutants lacking essential catalytic residues were inactive (Fig. 4C). Thus, these results collectively demonstrate that V\textsubscript{v}RRSP is a Ras/Rap1-specific endopeptidase that requires the essential 2His/2Glu catalytic residues in the C2 domain.

RRSP cleaves between residues Tyr-32 and Asp-33 in the Switch I region (\textsuperscript{30}DEYDPTIEDSY\textsuperscript{40}) in K/H/N-Ras, and \textsuperscript{30}EKYDPTIEDSY\textsuperscript{40} in Rap1A/B) of the cellular substrates Ras and Rap1 following sequence-specific substrate recognition (19). Remarkably, the sequences in the Switch I region are dominated by several negatively charged residues (Fig. S7) that might be involved in forming contacts with the RRSP active site. In addition to the catalytic His/Glu residues, a highly conserved basic residue is found near the active site residues in all TIKI proteases (22). The corresponding residue in V\textsubscript{v}RRSP is Arg-4001 (Fig. 3, B and C). We therefore presumed that Arg-4001 may be a substrate recognition residue. Proteolytic processing assays with RRSP mutated at Arg-4001 revealed complete loss of catalytic activity, indicating that this residue may indeed play a role in substrate recognition (Fig. 4D).

During refinement of the RRSP structure, we noticed that the electron density map around residues Arg-3988–Gln-3992 located close to the active site in the C2 domain is disordered (Fig. 4E), indicating flexibility and a potential involvement in substrate recognition. Given that the catalytic residues of V\textsubscript{v}RRSP were successfully assigned, we hypothesized that substitution of catalytic residues with nonfunctional residues might stabilize the active site and the adjacent flexible region. We subsequently determined the structure of the RRSP\textsubscript{E3930L/H44030L} mutant at a resolution of 2.3 Å (Table 1). The structure of the mutant RRSP is essentially identical to that of WT RRSP (RMSD of Ca atoms is less than 0.5 Å), with minimal differences in the conformation of the active sites containing the essential residues (Fig. 4E). Like WT RRSP, the active site of the V\textsubscript{v}RRSP\textsubscript{E3930L/H44030L} mutant is arranged similarly to that of BcR135 (Fig. S8). Luckily, the flexible region spanning residues Arg-3988–Gln-3992 in molecule A of the mutant is ordered and adopts a loop structure close to the active site (Fig. 4E). To evaluate the involvement of this region in the function of RRSP, we selected Arg-3988 based on its interactions with
Figure 2. Molecular details of VvRRSP. A, structure of VvRRSP divided into subdomains. The N1, N2, C1, and C2 domains of VvRRSP are colored blue, gray, green, and magenta, respectively. The linker region between the N and C lobes (inter-lobe linker) is colored orange. B, three-helix bundle within the N1 domain. Residues forming the hydrophobic core of the bundle are indicated. C, superimposition of the VvRRSP N1 domain (blue) onto the PMT (PDB code 2EBF) C1 domain (orange). D, structural stabilization of the N1 domain. The critical residues Phe-3636 of α3 and Phe-3670 of α4 that are involved in structural stabilization are indicated. Residues involved in forming the groove in the N2 domain are shown in light blue surface representation and are indicated. E, structural integration between the C lobe and the N2 domain. Helix α20 in the C2 domain (magenta), α12 in the C1 domain (green), and α7 in the N2 domain (light blue) are displayed in cartoon representation. Residues involved in hydrophobic and charge interactions between the domains are indicated. F, inter-lobe linker connecting the two lobes shown in stick representation. The N2 domain in the N lobe and the C1 and C2 domains in the C lobe are colored light blue, green, and magenta, respectively. The enlarged box shows residues Tyr-3723 and Leu-3724 that may be critical for controlling flexibility between the N and C lobes. G, in vitro KRas cleavage assays showing the importance of residues Tyr-3723 and Leu-3724. KRas* indicates cleaved KRas.
the Switch I region of Ras, dominated by negatively charged residues (Fig. S7). This residue was mutated to glutamate, and the overall conformation was confirmed by CD spectroscopy (Fig. S4). In vitro KRas cleavage assays showed that the mutation completely inhibited enzymatic activity (Fig. 4D), suggesting that the flexible loop may be involved in substrate recognition.

Because we could not observe an interaction between WT RRSP and Ras, we used the inactive RRSP mutant to examine interactions with KRas. However, in vitro pulldown assays revealed that the nonfunctional RRSP was similarly unable to maintain interaction with KRas (Fig. 4F and Fig. S9). No differences in the active sites were observed between WT and mutant RRSP structures. Therefore, these results suggest that residues Glu-3900, Glu-3930, His-3902, and His-4030 in the active site may spontaneously adopt the catalytically active conformation in the presence of substrate.

Comparison of VvRRSP and homologs revealed the highest similarity with the C1/C2 domains of PMT (PDB code 2EBF; RMSD = 2.96 Å for 404 Cα atoms; Fig. S2), even though VvRRSP shares only 26% amino acid sequence identity with PMT. PMT also belongs to the TIKI superfamily, but it lacks the catalytic 2His/2Glu motif in the common β-sheet core (Fig. S10, A and B). Accordingly, endopeptidase activity against KRas was not observed for the PMT C1/C2 domain (Fig. S10C). No metal bound to VvRRSP during refinement of its structure, we therefore performed isothermal titration calorimetry, and no direct interaction was detected between VvRRSP and benzamidine (data not shown). Together, these results indicate that VvRRSP is a metal-independent TIKI family endopeptidase.

**Structure of Ras/Rap1-specific endopeptidase**

TIKI proteases have a wide variety of domains appended to the conserved TIKI fold structure that might act to promote enzymatic function (30). VvRRSP has an N-terminally extended N lobe as part of its MLD. Thus, to further assess the role of the N lobe, we constructed various deletion mutants (Fig. 6A) and evaluated them by in vitro KRas cleavage assay (Fig. 6B). Deletion of the N lobe and inter-lobe linker (Δ3580–3736), leaving...
only the C lobe, resulted in complete loss of enzymatic activity toward KRas (Fig. 6B). Similarly, the C2 catalytic domain alone (residues 3856–4072) was unable to hydrolyze KRas (Fig. 6B). Interestingly, deletion of the MLD N1 domain (Δ3580–3671) significantly reduced the efficiency of KRas cleavage activity compared with WT RRSP (Fig. 6B). Thus, these results suggest that the N lobe and inter-lobe linker are crucial for substrate recruitment or recognition via integration with the C lobe to induce the functionally competent conformation of VvRRSP. It is worth mentioning that the inter-lobe linker may act as a flexible hinge to properly arrange the two lobes into a functional conformation that is enzymatically active. In particular, Tyr-3723 and Leu-3724 on the linker may be key residues regulating flexibility (Fig. 2, F and G). Only deletion of α1 (Δ3580–3609) within the MLD failed to affect the enzymatic activity (Fig. 6B).

Consistent with the in vitro assay results, the cleavage of endogenous Ras was entirely blocked in HEK293T cells transfected with plasmid expressing the N lobe deletion mutant (residues 3737–4089), and the activity of the MLD-deleted RRSP (residues 3672–4089) was significantly impaired (Fig. 6C). We further analyzed the cellular localization and enzymatic activity of VvRRSP by confocal microscopy. As reported previously (23), the MLD alone (residues 3580–3671) is mainly localized to the membrane in HEK293T cells that do not exhibit Ras cleavage activity when ectopically expressing this truncated construct (Fig. 6D). However, the MLD-deletion mutant (residues 3672–4089) or the C lobe mutant (residues 3737–4089) was dominantly expressed in the cytoplasm, again without enzymatic activity (Fig. 6D). Remarkably, the α helix appears to be critical for membrane localization of RRSP. Without the α helix (Δ3580–3609), membrane localization does not occur, resulting in reduced enzymatic activity, indicating that proper membrane localization of RRSP is required for enzymatic function (Fig. 6D). It is worth noting that in the structure the α helix points away from the MLD helix bundle (Fig. 2A). Taken together, these results demonstrate that the N lobe of RRSP performs dual roles: localization to the membrane via the α helix of the MLD and promoting enzymatic function.

**Discussion**

Although mainly studied in the context of broad medical research in tumor initiation, progression, and metastasis, Ras proteins also play fundamental roles in immunity and inflammation (20). The Ras superfamily of small GTPases can be
divided into at least six subfamilies (Ras, Rho, Ran, Rab, Rheb, and ARF). Among these, Ras subfamily proteins K/H/N-Ras and Rap1A/B have an identical amino acid sequence in their switch loops (18). This feature may confer the specificity of RRSP toward these two cellular substrates. The activation of Ras-dependent extracellular signal-regulated kinase-1 (ERK) signaling plays crucial roles in T-cell responses and Fcγ receptor-mediated phagocytosis (31–33). ERK signaling is also related to Toll-like receptor-mediated chemokine production in dendritic cells (34). GTP-bound Ras activates the phosphoinositide 3-kinase pathway that is crucial for the activation and functioning of all immune cells (32, 35, 36). Rap1 is also implicated in the activation of the innate immune system in various immune cells (37–40). Thus, it can be anticipated that robust inactivation of Ras and Rap1 by RRSP would lead to the failure of innate immune defenses and consequent enhancement of the pathogenicity of invading bacteria.

Bacterial effectors use diverse mechanisms to modulate the activity of host small GTPases and interfere with corresponding host-signaling pathways. Many effectors delivered by type III and IV secretion systems modulate the GTPase cycle of proteins belonging to the Rho, Rab, and ARF families by mimicking the functions of host regulators (41). Effectors also modulate the activity of small GTPases via modifications such as adenylation, glucosylation, N-acetylglucosamination, and proteolytic cleavage (42–46). Thus, bacterial effectors preferentially target molecular switch GTPases to efficiently subvert diverse cellular responses in the host–pathogen interactions.

Figure 5. VvRRSP is a metal-independent Ras-specific endopeptidase. A, in vitro KRas cleavage assays showing metal-independent RRSP activity. KRas* indicates cleaved KRas. B, effects of protease inhibitor cocktails including phenylmethanesulfonyl fluoride, pepstatin, leupeptin, benzamidine, and bestatin on the KRas cleavage activity of RRSP. Trypsin was used as a control. C, in vitro KRas cleavage assay showing that benzamidine inhibits the activity of RRSP. Ethanol was used as a control. D, effect of benzamidine (4 mM) on the KRas cleavage activity of RRSP. Data are representative of two independent experiments with similar results. E, superimposition of the structure of KRas complexed with GTP (PDB 5W22, white) onto that of KRas complexed with GTP and benzamidine (PDB 4DSO, magenta). Switch I and II regions in the structure of KRas complexed with GTP and benzamidine are colored yellow and green, respectively. Benzamidine and GTP are shown in stick representation.
The RRSP effector from the highly virulent *V. vulnificus* strain CMCP6 clinical isolate was initially identified as a MARTX toxin effector that is cytotoxic in HeLa cells (24), and it was later confirmed as a site-specific endopeptidase that effectively processes Ras and Rap1 (18). However, a lack of structural information has impeded our understanding of the mechanisms underlying how it recognizes and processes its cellular substrates. Our current structural analysis reveals that *Vv*RRSP shares an active site architecture with TIKI superfamily proteins (22), including Ere family enzymes known to catalyze the hydrolysis of esters and proteins (27). It has been proposed that enzymatically active TIKI domain proteins share a catalytic mechanism with Ere family enzymes, wherein the His–Glu pair conserved among TIKI domain proteins activates a water molecule for nucleophilic attack on the carbonyl group of ester and peptide bonds (22, 27). His-85 and Glu-113 form the His–Glu pair in Bcr135, corresponding to His-3902 and Glu-3930 in *Vv*RRSP (Fig. 3A). It remains uncertain whether *Vv*RRSP shares the same catalytic mechanism with TIKI proteins because the His-4030–Glu-3930 pair was assigned as the essential catalytic residues in *Vv*RRSP, rather than the His-3902–Glu-3930 pair (Fig. 4, A and B). However, we cannot rule out the possibility that *Vv*RRSP shares a catalytic mechanism with TIKI proteins because a water molecule was observed close to Glu-3900 and Glu-3930 in WT *Vv*RRSP (Fig. S11A) and in the center of the active site of the catalytically nonfunctional mutant (Fig. S11B).

The structure of *Vv*RRSP revealed that the functional conformation of the N1 domain is promoted by the N2 domain (Fig. 2D). This structural feature may render the 14-amino acid residue α1 helix suitable for membrane anchoring (Fig. 2A). One side of α1 is composed of hydrophobic residues, and the other side has positively charged residues that may enhance membrane interaction (Fig. S12A). Superimposition of *Vv*RRSP molecules in the asymmetric unit reveals different conformations for the MLD fingers (Fig. 1C), indicating flexibility. Indeed, part of the α1 helix is involved in crystal packing contacts, its flexibility is independent of the rest of the helices of the N1 domain in the RRSP structure, and it may be stabilized by these contacts (Fig. S12B). Thus, we cannot rule out the possibility that the α1 helix conformation in the structure of *Vv*RRSP may be due to the crystal packing contacts. The membrane-interacting conformation of the N lobe likely stabilizes the functional conformation of RRSP in association with the active site-containing C lobe, thereby promoting recruitment of substrates from the membrane.

Pathogenic bacteria have evolved mechanisms to effectively combat host defenses. Expanding our understanding of how bacteria manipulate immune systems in humans will provide
promising strategies for controlling infection. Targeting the Ras and Rap1 signaling switches could offer an efficient way to subvert host signaling systems and augment the virulence of pathogenic bacteria. Because RRSPs are conserved among pathogens infecting mammals, fish, and plants, our detailed structural and functional analysis of VvRRSP might be broadly applicable for the development of strategies to combat powerful virulence mechanisms.

**Experimental procedures**

**Bacterial strains, plasmids, and cell culture**

Bacterial strains and plasmids used in this study are listed in Table S1. *Escherichia coli* strains were grown in Luria-Bertani (LB) medium at 37 °C with appropriate antibiotics. HEK293T cells were purchased from the American Type Culture Collection (ATCC) and cultured at 37 °C with 5% CO2 in HyClone Dulbecco’s modified Eagle’s medium (GE Healthcare) supplemented with 1% antibiotic/antimycotic (Gibco-BRL) and 10% fetal bovine serum (Gibco-BRL).

**DNA cloning and mutagenesis**

Genomic DNA from *V. vulnificus* strain CMCP6 was kindly provided by Dr. Joon Haeng Rhee from Chonnam National University Medical School and utilized as a template for amplifying the DNA fragment encoding the effector domain (residues 3580–4072) of the MARTX toxin, named VvRRSP. The amplified DNA fragment was subcloned into the Ncol and Xhol restriction enzyme sites of the pHis-Parallel1 expression vector, a protein expression vector that attaches an N-terminal His6-tag cleavable using recombinant tobacco etch virus (rTEV) protease. To investigate the critical residues responsible for endopeptidase activity, constructs for expression of (rTEV) protease. To investigate the critical residues responsible for endopeptidase activity, constructs for expression of proteins were constructed using site-directed mutagenesis. DNA fragments corresponding to an N-terminal deletion mutant (residues 3596–4072) was incorporated into the same expression vector using the same method. In addition, to investigate the domains related to substrate recognition and binding, DNA fragments expressing truncated forms of VvRRSP, namely residues 3610–4072, 3737–4072, 3580–3855, and 3856–4072, were subcloned into the pHis-Parallel1 expression vector. For analysis of localization and cytotoxicity, a DNA fragment encoding residues 3580–4089 was amplified and inserted into the Nhel and Apal sites of the pAcGFP vector for expression of C-terminal GFP fusion proteins. Several plasmids expressing mutated or truncated forms of VvRRSP fused with a C-terminal GFP, including RRSPE3900L, RRSPE3930L, RRSPH3902L, RRSPH4030L, RRSPY3723A, RRSPR3988E, and RRSPR4001F were generated using PCR products were subcloned into the NcoI and XhoI sites of the pHis-Parallel1 expression vector. The DNA fragment corresponding to an N-terminal deletion mutant (residues 3596–4072) of the MARTX toxin, named VvRRSP was overexpressed in *E. coli* BL21-CodonPlus (DE3)-RIPL cells by induction with 0.5 mM isopropyl β-D-thiogalactopyranoside (IPTG) at 21 °C for 18 h. Cultured cells were harvested by centrifugation at 5000 × g for 10 min at 4 °C, resuspended in ice-cold buffer A (50 mM Tris-HCl (pH 7.5), 300 mM NaCl), and lysed with a high-pressure homogenizer (Nano DeBEE, B.E.E. International). Crude cell extracts were centrifuged at 25,000 × g for 4 °C for 1 h. The supernatant containing hexaHis-tagged RRSP (residues 3596–4072) was loaded onto a Ni-NTA–agarose column (Qiagen Venlo, The Netherlands) pre-equilibrated with buffer A. Nonspecific resin-bound proteins were washed out with buffer A supplemented with 30 mM imidazole, and hexaHis-tagged RRSP was eluted with buffer A containing 250 mM imidazole. The hexaHis tag was cleaved by incubating the eluted protein with rTEV protease (Gibco) at 4 °C overnight. After protein concentration using a centrifugal concentrator with a 3-kDa cutoff (Amicon Ultra-15; Merck Millipore Ltd., Burlington, MA), size-exclusion chromatography was performed using a Superdex 200 10/300 column (GE Healthcare). Fractions containing RRSP (residues 3596–4072) were pooled and loaded onto a second Ni-NTA–agarose column for further purification, and the flow-through containing the tag-free RRSP (residues 3596–4072) was concentrated to 15 mg/ml and stored at −80 °C after clarification by centrifugation at 16,000 × g for 30 min. Selenomethionine (SeMet)-substituted RRSP (residues 3596–4072) was expressed in *E. coli* B834 (DE3) cells (Merck Millipore Ltd.), a methionine auxotrophic strain, in minimal medium supplemented with 50 mg/ml SeMet and purified as described above for native RRSP (residues 3596–4072) in the presence of 5 mM methionine throughout all purification steps. RRSP3930L/H4030L mutants were also expressed and purified as described for RRSP (residues 3596–4072). Purified proteins include a four-residue cloning artifact (Gly–Ala–Met–Ala) at the N terminus. For biochemical assays, expression of variants, including mutated and truncated forms, was carried out as described for RRSP (residues 3596–4072), but purification involved only one-step affinity column chromatography. Purified variants were stored at −80 °C after clarification by centrifugation at 16,000 × g for 30 min.
Structure of Ras/Rap1-specific endopeptidase

30 min. Expression of KRas was induced in *E. coli* BL21-Codon-Plus (DE3)-RIPL cells by treatment with 0.5 mM IPTG at 21 °C for 18 h. Harvested cells were resuspended in ice-cold buffer A and lysed using a high-pressure homogenizer (Nanoc DeBEE, B.E.E.). The recombinant N-terminally His₆-tagged KRas in the supernatant was purified by Ni-NTA affinity chromatography, and the buffer was exchanged to 50 mM Tris- HCl (pH 7.5) and 300 mM NaCl for *in vitro* Ras cleavage assays. KRas fused with GSH S-transferase (GST) was also expressed and purified for *in vitro* pulldown analysis. The homogeneity of proteins was assessed by 12% SDS-PAGE and visualized using Coomassie Blue staining. Raf1 (residues 54–131) was made similar to that of KRas for pulldown analysis.

Size-exclusion chromatography

A Superdex 200 10/300 GL gel-filtration column (GE Healthcare) was installed on a fast protein LC (FPLC) system (Bio-Rad) equipped with a 28% PEG 4000, and 0.1 M Tris-HCl (pH 8.0), and X-ray diffraction data were performed using AutoBuild in the PHENIX software package. Initial crystals were produced under the same conditions, however, high-quality crystals were obtained for X-ray diffraction. The supernatant was purified by Ni-NTA affinity chromatography, and bovine serum albumin (BSA) was used as a molecular weight standard.

Crystallization, X-ray diffraction, and structure determination

Initial crystallization screening was performed using the sitting-drop vapor diffusion method against commercially available sparse-matrix screening kits (Hampton Research), with drops consisting of 0.4 μl of protein (15 mg/ml) in buffer A and 0.4 μl of reservoir solution in 96-well MRC plates. Initial crystals were produced in several different conditions. However, high-quality crystals for X-ray diffraction were only obtained after optimizing conditions, and 1.4 mM ammonium sulfate, 0.1 mM Tris-HCl (pH 8.0), and 12% glycerol proved optimal. Crystals of RRSP (residues 3596–4072) were frozen in cryoprotectant solution consisting of 1.6 mM ammonium sulfate, 0.1 mM Tris-HCl (pH 8.0), and 12% glycerol and exposed to X-rays in a nitrogen gas stream at -173 °C. Crystals diffracted X-rays to a resolution of 2.7 Å at beamline 5C of the Pohang Accelerator Laboratory (PAL) in Korea. Crystals of SeMet-substituted RRSP (residues 3596–4072) were grown and optimized under the same conditions, and single-wavelength anomalous diffraction data were collected at a resolution of 2.9 Å using the same beamline. Crystals of RRSP, SeMet-substituted RRSP (residues 3596–4072) were grown in 0.2 M LiSO₄, 28% PEG 4000, and 0.1 M Tris-HCl (pH 8.0), and X-ray diffraction data were also collected at beamline 5C of PAL to a resolution of 2.3 Å. All X-ray data were processed and scaled with the HKL2000 software package (47). The structure of SeMet-substituted RRSP (residues 3596–4072) was initially determined with the AutoSol phasing module utilizing the anomalous signals from selenium atoms, and automatic model building was performed using AutoBuild in the PHENIX software package. Initial model building of SeMet-substituted RRSP (residues 3596–4072) was manually performed with COOT (49). The structure of native RRSP (residues 3596–4072) was solved by molecular replacement in MOLREP using the partial model of SeMet-substituted RRSP (residues 3596–4072) as a template (50). COOT was used for model building, and refinement, including the translation-liberation-screw procedure, was carried out with REFMAC5 and PHENIX (48, 51). The native structure was re-refined using the higher resolution mutant structure for further improvement of structural quality. Residues 3988–3992 and 4070–4072 in molecule A, 3989–3992 and 4068–4067 in molecule B, 3989–3992 and 4067–4072 in molecule C, and 3988–3992 and 4067–4072 in molecule D are disordered and thus not included in the final model. The structure of RRSP, was solved using the first model of native RRSP as a template and refined in the same way. Residues 4069–4072 in molecule A, 3989–3991 and 4067–4072 in molecule B, 3989–3993 and 4066–4072 in molecule C, and 3988–3993, and 4066–4072 in molecule D are disordered and not included in the final model of RRSP. Crystallographic data are summarized in Table 1.

In vitro Ras cleavage and pulldown assays

Purified KRas (0.5 μM) was incubated with RRSP or PMT at a ratio of 1:1 or 2:1 in 50 mM Tris-HCl (pH 7.5) and 300 mM NaCl. After incubating the KRas-RRSP or KRas-PMT mixture at 37 °C for 20 min, the reaction was stopped by addition of SDS-PAGE sample buffer. Proteins were separated on a 15% gel and visualized using Coomassie Blue staining. For analysis of inhibitors of RRSP activity, purified RRSP was preincubated with the indicated protease inhibitors at 37 °C for 30 min. After adding KRas, the mixture was further incubated at the same temperature for the indicated time. Samples were separated by SDS-PAGE and visualized by Coomassie Blue staining.

For pulldown assays, plasmids expressing KRas fused with GST were used as bait proteins, mixed with Raf1 (residues 54–131) or variants of RRSP, including mutated and truncated forms, and incubated with GSH-conjugated Sepharose beads (GE Healthcare) in 50 mM Tris-HCl (pH 7.5), 300 mM NaCl, 1 mM GTP, and 5 mM MgCl₂ for 3 h at 4 °C. After intensive washing with incubation buffer, KRas fused with GST was eluted with the same buffer supplemented with 20 mM reduced GSH. RRSP variants eluted from the resin were analyzed by SDS-PAGE and visualized using Coomassie Blue staining.

Transient transfection

HEK293T cells were grown to ~70–80% confluency, and transfection was implemented using X-tremeGENE HP reagent (Roche Applied Sciences, Penzberg, Upper Bavaria, Germany) according to the manufacturer’s instructions.

Endogenous Ras cleavage assay

HEK293T cells, transfected with plasmids expressing RRSP or its variants, were harvested at 16 h after transfection and lysed in lysis buffer (0.5% Nonidet P-40, 150 mM NaCl, 10% glycerol, and 20 mM HEPES (pH 7.2)) supplemented with complete protease inhibitor mixture (Roche Applied Science). After 30 min of incubation on ice, cell lysates were centrifuged at 16,000 × g for 30 min. Supernatants were mixed with LDS-PAGE loading buffer (Life Technologies, Inc.); proteins were separated by SDS-PAGE and transferred onto a polyvinylidene fluoride membrane by the semi-dry blot method; and RRSP, Ras, and Rap1 proteins were detected by Western blotting using a chemiluminescence detection system (ThermoFisher Scien
tific, Waltham, MA). To detect Strep-tagged RRSP and its variants, a horseradish peroxidase (HRP)-conjugated mouse anti-Strep antibody (IBA Lifesciences, Göttingen, Germany) was used according to the manufacturer’s protocol. To detect endogenous Ras, mouse monoclonal anti-Ras antibody (1:2000; Millipore) and HRP-conjugated anti-mouse IgG antibody (1:5000; Cell Signaling Technology, Danvers, MA) were used. To detect endogenous Rap1, mouse monoclonal anti-Rap1 antibody (1:1000; Santa Cruz Biotechnology, Dallas, TX) and HRP-conjugated anti-mouse IgG antibody were used. Actin was detected using an HRP-conjugated rabbit monoclonal anti-β-actin antibody (1:5000; Cell Signaling Technology). All experiments were repeated at least three times, and data are expressed as the mean ± S.D. using Prism (Version 6.0, GraphPad Software).

Cell morphological analysis

HEK293T cells were transfected with plasmids expressing the indicated RRSP C-terminal GFP constructs. After 16 h, cell morphological changes were examined using a FLoid Cell Imaging Station (ThermoFisher Scientific).

Confocal microscopy

HEK293T cells were seeded into μ-slide 8-well plates (Ibidi, Martinsried, Germany) and transfected with plasmids expressing RRSP (residues 3580–4089) or its variants tagged with GFP at the C terminus. After 16 h, cells were fixed in 4% paraformaldehyde in phosphate-buffered saline (PBS) at 25 °C for 10 min. Fixed cells were washed with PBS and permeabilized with 100% methanol for 10 min at −20 °C. After further washing with PBS, cells were blocked with 2% BSA in PBS at 25 °C for 1 h and then incubated with Pan-Ras antibody (1:100; Millipore) overnight at 4 °C. After washing twice with PBS containing 0.1% Triton X-100 (PBS-T) and once with PBS, cells were incubated with anti-mouse IgG secondary antibody Alexa Fluor 568 (1:150; Invitrogen, Waltham, MA) at 25 °C for 1 h. For further washing three times, nuclei were stained with Hoechst 33342 (ThermoFisher Scientific) at 25 °C for 10 min and further washed with PBS-T. To visualize actin filaments, RRSP-transfected cells were stained with rhodamine-phalloidin (1:1000; ThermoFisher Scientific) at 25 °C for 20 min. Mounting medium was applied to stained cells, and images were acquired using a Nikon laser-scanning confocal microscope (C2plus) and analyzed using NIS-Elements software.

Circular dichroism (CD) spectroscopy

CD spectroscopy was used to compare the structural conformation of RRSP WT and its mutants. Pure protein (0.2 mg ml⁻¹) in 50 mM Tris-HCl (pH 7.5) and 300 mM NaCl was subjected to far-UV CD measurements at 20 °C using a 1-mm path length quartz cuvette in a Jasco J-815 CD spectrometer (Jasco, Tokyo, Japan). CD spectra were acquired over the wavelength range of 200–260 nm and converted into mean residue ellipticity (degree cm² dmol⁻¹). Blank spectra obtained using buffer without protein were subtracted.

PDB codes

The atomic coordinates and structure factor amplitudes of WT RRSP and the RRSPF3930L/H4030L mutant have been deposited in the PDB (52) under accession codes 6A8J and 6A7H, respectively.

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