Substrate Recognition of MARTX Ras/Rap1-Specific Endopeptidase

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Substrate recognition of MARTX Ras/Rap1 specific endopeptidase

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Abstract

Ras/Rap1-specific endopeptidase (RRSP) is a cytotoxic effector domain of Multifunctional-autoprocessing repeats-in-toxins (MARTX) toxin of highly virulent strains of Vibrio vulnificus. RRSP blocks RAS-MAPK kinase signaling by cleaving Ras and Rap1 within the Switch I region between Y32 and D33. Although the RRSP processing site is highly conserved among small GTPases, only Ras and Rap1 have been identified as proteolytic substrates. Here we report that Y32-D33 residues at the scissile bond play an important role for RRSP substrate recognition, while nucleotide state of Ras has only a minimal effect. In addition, substrate specificity is generated by residues across the entire Switch I region. Indeed, swapping the Ras Switch I region into either RalA or RhoA, GTPases that are not recognized by RRSP, generated chimeras that are substrates of RRSP. However, a difference in processing efficiency of the Ras Switch I in the context of Ras, RalA, or RhoA indicates that protein regions outside the Ras Switch I also contribute to efficient RRSP substrate recognition. Moreover, we show that synthetic peptides corresponding to the Ras and Rap1, but not RalA, Switch I regions are cleaved by RRSP demonstrating sequence specific substrate recognition. In conclusion, this work demonstrates that the GTPase recognition of RRSP is nucleotide independent, and is mainly driven by the Ras and Rap1 Switch I loop and also influenced by additional protein-protein interactions, increasing the substrate specificity of RRSP.

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Author Contributions
M.B. conceived and conducted most of the experiments, analyzed the data and coordinated the study. Z.L. A.R.L. A.E.R., and B.L.P. produced the synthetic peptides, conducted in vitro cleavage studies, and analyzed data. M.B. and K.J.F.S. wrote the manuscript. All authors reviewed the results and approved the final version of the manuscript.

Notes
M.B. and K.J.F.S. hold a patent application covering RRSP application. The remaining authors declare no competing financial interest.

ASSOCIATED CONTENT
Supporting Information.
Figure S1 (.pdf)
Table S1 (.pdf)
List S1 (.pdf)
Keywords  
bacterial toxin; Ras protein; Ras-related protein 1; GTPase KRas; bacterial pathogenesis; RRSP; endopeptidase; substrate specificity

TEXT 
*Vibrio vulnificus* is a motile Gram-negative foodborne pathogen that can cause life-threatening infections. Ingestion of contaminated seafood or the exposure of open wounds to water harboring *V. vulnificus* can result in fatal septicemia, gastroenteritis and necrotizing fasciitis. The mortality rate of human subjects with liver damage or compromised immune systems can be greater than 50%. The severity of *V. vulnificus* infection is due to the secretion of the cytolytic pore-forming *V. vulnificus* cytolysin (VVC) and the multifunctional autoprocessing repeats-in-toxin (MARTX) toxin, which together diminish the innate immune response and induce tissue necrosis at the site of infection, helping the bacterial to invade and spread systemically. In particular, MARTX has been recognized as the most significant virulence factor for *V. vulnificus* pathogenesis.

MARTX toxins are large protein toxins (3500–5300 aa) secreted by Gram-negative bacteria. MARTX toxins may carry up to five different enzymatically active effector domains that are delivered into the eukaryotic cytosol across the plasma membrane. Once in the cytosol, the MARTX cysteine protease domain (CPD) binds inositol hexakisphosphate, initiating autoprocessing of the toxin at leucine residues located in unstructured regions that link the effector domains. When released in target cells, each effector domain has a unique enzymatic function, resulting in cytopathicity or cytotoxicity.

The toxin effector designated Domain of Unknown Function in position 5 (DUF5) is carried by some, but not all, *V. vulnificus* MARTX toxins. Notably, a *V. vulnificus* strain producing a MARTX toxin with DUF5 shows about 50-fold higher LD₅₀ in mice than strains producing a toxin without DUF5. In addition, delivery of DUF5 alone into cells results in cytotoxicity characterized by cell rounding and lack of cell proliferation. In order to understand how DUF5 dramatically increases the virulence potential of *V. vulnificus*, we recently solved its molecular mechanism within cells. DUF5 directly inhibits the mitogen-activated protein kinases pathway (MAPK) by proteolytically processing the small guanosine triphosphatase (GTPase) Ras, thereby preventing the phosphorylation of extracellular signal–regulated kinases (ERK) and blocking cell proliferation.

Ras is the representative protein of the small GTPase superfamily that includes more than 150 proteins separated into five families: Ras, Rho, Rab, Arf, and Ran. These small GTPases control various cell processes by alternating conformational states finely regulated by guanine-nucleotide-exchange factors (GEFs), which promote formation of the GTP-bound active state, and by GTPase-activating proteins (GAPs), which accelerate the intrinsic GTPase activity to promote formation of the GDP-bound inactive state. In particular, the conformation of the Switch I region (residues 30–38 in Ras) in the GTP-bound state is essential for binding of effectors and activation of downstream signaling cascades. DUF5 was found to cleave Ras between Y32 and D33 within the Switch I region.
There are three human RAS genes (NRas, HRas and KRas) that encode four Ras proteins: NRas, HRas, and two isoforms of KRas, of which KRas4B is the predominant splice variant expressed in many tissues. The primary sequence of the four Ras proteins is highly conserved (85–87% amino acid sequence identity) and they share many structural and biochemical properties. The N-terminal G domain (residues 5–166) is involved in the nucleotide binding and GTP hydrolysis including the highly conserved Switch I region (Fig. 2). The hypervariable regions (HVRs) at the C-termini are subject to different post-translational modifications that direct Ras isoforms to different subcellular locations. In our previous study, we found that DUF5 cleaves full-length KRas, NRas and HRas equally well, demonstrating that the different C-terminal HVRs among Ras isoforms are not involved in substrate recognition by the endopeptidase.

The Ras subfamily, in addition to KRas, HRas, and NRas, includes other similar GTPases like Ras-related proteins 1 and 2 (Rap1 and Rap2), as well as Ral, Rheb, and Rit proteins. These other subfamily members regulate many cellular functions including cell proliferation, differentiation and survival, cell adhesion, mTOR signaling, and neuronal differentiation. Among several small GTPases tested belonging to this Ras subfamily, DUF5 was found to efficiently cleave only Rap1 in addition to Ras, and did not cleave RalA, RheB2, or Rit2. The endopeptidase also did not cleave GTPases from other small GTPase subfamilies, including RhoA, which regulates cytoskeleton assembly. Since DUF5 exclusively cleaves the small GTPases Ras and Rap1, this effector domain was renamed Ras/Rap1-specific endopeptidase (RRSP).

Ras itself is the central node in the cell that coordinates extracellular signals to ultimately regulate expression of genes for cell proliferation, differentiation, and survival through its control of the MAPK signaling cascade. Rap1 mainly controls establishment of cell polarity, activation of integrin-mediated cell adhesion and the regulation of cell–cell contacts. Interestingly, both Ras and Rap1 control signal transduction in response to bacterial infection, likely accounting for why they are targeted by this bacterial toxin, although the biochemical basis for the substrate specificity has not been previously investigated.

In this study, we demonstrate the Switch I sequence confers specificity of RRSP for Ras and Rap1. RRSP is able to cleave Ras and Rap1 Switch I peptides corresponding to residues comprised between 28 and 38, but not a peptide from the closely related RalA. In addition, swapping the Ras Switch I sequence into RalA and RhoA converts these non-substrate GTPases into substrates. Differences in the efficiency of processing of these chimeras reveal a contribution of other regions of the substrates to the recognition. The contribution is unlikely to be related to nucleotide binding since RRSP cleaves both GTP- and GDP-bound Ras. Overall, we show that the substrate specificity of RRSP for Ras and Rap1 is mainly driven by sequence specific recognition of the Switch I sequence of Ras and Rap1 with regions outside of the Switch I region contributing to processing efficiency.
EXPERIMENTAL PROCEDURES

Materials

*E. coli* TOP10 cells were obtained from Life Technologies and *E. coli* BL21(DE3)Magic and pMCSG7 bacterial expression vector were provided by Andrzej Joachimiak (Argonne National Laboratory, IL). Proteins were expressed in the kanamycin-resistant BL21(DE3) (pMagic) *E. coli* strain. The pMagic plasmid provides the rare codon tRNA gene argU (40 and A. Joachimiak, personal communication). Recombinant KRas-Tagless (2–185) proteins loaded with GDP and GppNHp were provided by Bindu Lakshman and Andrew Stephen (Frederick National Laboratories for Cancer Research, MD). The amount of nucleotide bound to KRas was detected by high-performance liquid chromatography and the elution times were compared to a standard curve of either GDP/GppNHp. Common reagents were obtained from Sigma-Aldrich, Fisher or VWR. SspI restriction enzyme and Gibson Assembly® Master Mix were obtained from New England Biolabs. Custom DNA oligonucleotide primers and GBlocks™ as listed in Table S1 were purchased from Integrated DNA Technologies (Coralville, IA). Plasmids were purified by using Epoch spin columns or Qiagen Midi Prep kit according to the manufacturer’s recommended protocol. Plasmids were introduced into *E. coli* BL21(DE3)(pMagic) by electroporation. Recombinant proteins were purified using ÄKTA Xpress Purifier system (GE Healthcare Bio-Sciences, Pittsburgh, PA). Prepacked columns with Nickel Sepharose resin (Ni-NTA) HisTrap column for affinity chromatography or Superdex 200 (26/60) for size exclusion chromatography (SEC) were purchased from GE Healthcare Bio-Sciences.

Expression and purification of RRSP

The N-terminal 6xHis tagged RRSP was purified over a Ni-NTA HisTrap column followed by SEC using the ÄKTA protein purification system as previously described.

Cloning, expression and purification of small GTPase

DNA sequences corresponding to RalA (NP_005393.2) and RhoA (NP_001300870.1) were amplified from templates previously described, using primers designed for Gibson Assembly® (New England Biolabs). PCR products were cloned into the pMCSG7 expression vector previously digested with SspI. The Y32A, D33A and D33E mutations were introduced by site-directed mutagenesis using the pMCSG7-KRas vector as a template. Custom gBlocks™ designed to introduce the modified KRas, RalA and RhoA Switch I sequences were assembled into the SspI-digested pMCSG7 vector using Gibson Assembly®, following the manufacturer’s recommended protocol. Plasmids were confirmed to be accurate by DNA sequencing and then transformed into BL21(DE3)Magic cells. All small GTPases were purified as described above for rRRSP, except all buffers were adjusted to pH 7.5 and contained 10 mM MgCl₂.

In vitro RRSP cleavage assay

Small GTPases were incubated with rRRSP proteins at equimolar concentrations (10 μM) in 10 mM Tris pH 7.5, 500 mM NaCl, 10 mM MgCl₂ at 37°C for 10 min. The time course cleavage assay the reaction was conducted at 10 min and 30 min, 1, 2, 4, and 8 hours; in

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addition, for rKRas and rRalA:S1:Ras the cleavage reaction was monitored even at shorter time points such as 1, 3 and 5 min. The time course cleavage assay with tagless GDP-rKRas and also GppNHp-rKRas was performed using 10 μM of GTPases and 10 nM of RRSP and monitored after 2, 4, 8, 16, 32 and 64 min. Reactions were stopped by adding 6X Laemmli sample buffer and incubating the sample at 90°C for 5 min. Proteins were separated on 18% SDS–polyacrylamide gels and visualized using Coomassie stain.

**SDS-PAGE analysis and quantification**

Cleavage products of GTPases were quantified from scanned SDS–polyacrylamide gels using NIH ImageJ 1.64. The percentage of cleavage product was calculated by the following formula:

\[ \text{% Cleavage product} = \frac{\text{cleaved band}}{\text{cleaved GTPase} + \text{ uncleaved GTPase}} \times 100 \]

where `cleaved GTPase` is the band running at 18 KDa. Data were fit in GraphPad Prism 6.0 to a one-phase exponential association curve to determine the \( k_{obs} \). Each point of the time course cleavage assay represents the average and standard deviation of results from three independent experiments.

**Peptide synthesis and in vitro peptide cleavage assay**

Peptides corresponding to Switch I sequence of Ras (FVDEYDPTIE), Rap1 (FVEKYDPTIE), Ral (FVEDYEPTKA) and Ras Switch I Y32A/D33A (FVDEAAPTIE) were synthesized by fast flow Fmoc solid phase peptide synthesis on H-Rink Amide-ChemMatrix resin\(^4\). Each peptide was purified using an Agilent Zorbax 300SB-C\(_{18}\) column (5 μm, 9.4 x 250mm) at 7 mL/min and 1–31% acetonitrile (0.1% TFA) over 60 minutes. 10 μM of each peptide was then incubated in the presence or absence of RRSP (10 μM) in 20 mM Tris pH 7.5 with 150 mM NaCl in a 37°C water bath. Time points were pulled at 0, 1, 2, 5, and 8 hours, quenched with 1:1 water (0.1% TFA):acetonitrile (0.1% TFA), and analyzed by Agilent 6520 ESI-Q-TOF mass spectrometer with an Agilent Zorbax 300SB-C\(_3\) column (5 μm, 2.1 x 150 mm) at 0.8 mL/min and 1–41% acetonitrile (0.1% formic acid) over 8 minutes.

**RESULTS**

**Y32 and D33 residues of Ras Switch I are important for the recognition by RRSP**

RRSP was previously shown\(^2\) to effectively process Ras isoforms (H/N/K) and Rap1 within the Switch I loop; the scissile bond in Ras and Rap1 was identified between Y32 and D33 (Fig. 1).

To define the impact of the scissile pair amino acids on RRSP activity, full-length recombinant KRas4B (rKRas) was purified, mixed at equimolar concentration with rRRSP, incubated at 37°C, and the reaction products were analyzed by SDS-PAGE. As shown previously, rKRas is efficiently cleaved by RRSP within 10 minutes reducing the electrophoretic mobility of rKRas by ~5 kDa.\(^2\) rKRas variants with Y32A and D33A amino acid substitutions were also generated and purified. About 50% of rKRas Y32A and
D33A variants were cleaved, indicating that both residues contribute to RRSP specificity, but neither is essential (Fig. 2A). By contrast, a rKras variant with both Y32A and D33A mutations was found to be resistant to RRSP processing, demonstrating an additive effect of both Y32 and D33 on RRSP specificity (Fig. 2A).

To quantify the processing defect, rKras and rKras Y32A, D33A and Y32A/D33A mutant variants were each tested as substrates in a time course experiment. As shown in Fig. 1B and Table 1, the catalytic rates of rRRSP for rKras Y32A (rRRSP $k_{\text{obs}} 0.031 \text{ min}^{-1}$) and D33A (rRRSP $k_{\text{obs}} 0.017 \text{ min}^{-1}$) were slower than for rKras (rRRSP $k_{\text{obs}} 2.35 \text{ min}^{-1}$), although complete cleavage of both did occur after 4 h. By contrast, rKras Y32A/D33A presented no cleavage products at any time points (Fig. 2B).

The loss of activity of RRSP for rKras Y32A/D33A could be due to either dramatically decreased substrate specificity or the direct involvement of both Y32 and D33 to the cleavage mechanism of RRSP. To define the role of the phenol group of Y32 or the carboxyl group of D33 for RRSP activity, rKras Y32F and rKras Y32F/D33A were produced and tested with rRRSP at equimolar concentration. Recombinant rKras Y32F, showed partial cleavage after 10 minutes, reaching a complete cleavage after 4 hours incubation, similar to the results obtained with the Y32A variant (Fig. 2C). In addition, RRSP $k_{\text{obs}}$ with rKras Y32F as substrate was 0.094 min$^{-1}$, about 25-fold lower than for rKras (Table 1), indicating that the reduced RRSP activity for both rKras Y32A and Y32F was due to the absence of the hydroxyl group. Thus, while the presence of the hydroxyl group is not essential for catalysis, and this group might play an important role for the interaction between RRSP and the Switch I of Ras.

However, rKras Y32F/D33A showed almost 35% cleavage only after 8 hours and RRSP catalytic activity was 0.00073 min$^{-1}$ (Fig. 2C,D and Table 1). This further suggests that the phenyl group of Y32F helps in the recognition of the Switch I by RRSP, but the absence of the carbonyl group in position 33 dramatically reduces the cleavage of the Switch I. Overall, these results show that residues Y32 and D33 play a crucial role for RRSP substrate recognition and interaction, but are not fundamental for RRSP catalysis.

**Residues downstream of Y32 are partially involved in RRSP substrate recognition**

It was also previously shown$^{22}$ that RRSP does not cleave RalA, a close relative of Ras and Rap in the Ras subfamily (Fig. 1). If Y32-D33 amino acids are the sole contributors to RRSP specificity, it seems unlikely that RRSP cannot cleave RalA, which has a conservative Y-E amino acid flanking its potential Switch 1 scissile bond (Fig. 1). Thus, recombinant mutant protein rKras D33E was purified and tested to assess whether RRSP can cleave a Y-E scissile bond. As shown in Fig. 3B rRRSP cleaved almost 90% of rKras D33E in 10 minutes, demonstrating that the presence of E44 within the RalA Switch I does not explain why RalA is not a substrate of RRSP.

RalA Switch I amino acids upstream of the scissile bond are highly conserved with Ras or Rap1, while downstream of D33 the RalA Switch I has K47 and A48 instead of the Ras or Rap1 I36 and E37. These amino acid differences change the overall charge downstream of the scissile bond (Fig. 1). To evaluate their contribution to RRSP substrate specificity, RalA
Switch I residues were introduced in the sequence of the rKRas Switch I region, generating the protein variant rKRas I36K and the double mutant protein rKRas I36K/E37A (Fig. 3A). Purified rKRas I36K and rKRas I36K/E37A were each incubated for 10 minutes with RRSP. About 90% of rKRas I36K was cleaved by RRSP, showing that the mutation I36K did not affect the interaction between KRas and RRSP (Fig. 3B). However, rKRas I36K/E37A showed only about 50% of cleaved product, suggesting that there was an effect when both residues were mutated, probably due to the change of the net charge from negative (I36-E37) to positive (K36-A37) (Fig. 3B). Moreover, rKRas D33E/I36K/E37A protein was also purified and tested with RRSP and similar to rKRas I36K/E37A, this protein was also cleaved about 50% (Fig. 3B). Taken together, these results suggest Switch I residues downstream of the scissile bond are important for RRSP specificity, but are not entirely essential.

The total sequence of Ras Switch I is necessary for RRSP specificity

To further investigate the contribution of the entire Switch I, the KRas Switch I sequence between 30–37 residues was entirely replaced with corresponding amino acids of non-substrate RalA Switch I (residues 41–48), creating rKRas D30E/E31D/D33E/I36K/E37A (rKRas:S1:Ral) (Fig. 4A,B). In 10 min, neither rRalA nor rKRas:S1:Ral was efficiently processed by RRSP (Fig. 4C), indicating that it is the RalA Switch I sequence that is poorly recognized by RRSP. When quantified, the change of Ras Switch I to RalA Switch I decreased the \( k_{\text{obs}} \) of RRSP from 2.35 min\(^{-1}\) (KRas) to 0.0041 min\(^{-1}\) (KRas:S1:Ral), a reduction of 573-fold (Fig. 4D and Table 1). This resulted in about 40% processing of rKRas:S1:Ral after 8 h (Fig. 4C). However, this residual cleavage of rKRas:S1:Ral compared to a complete absence of cleavage of rRalA reveals that while RRSP can at very low efficiency cleave the RalA Switch I in vitro, this occurs only when appropriately presented to RRSP in the context of the KRas protein, suggesting there may be a contribution of the backbone to RRSP substrate recognition.

RalA but not RhoA backbone interactions contribute to processing by RRSP

To further explore the potential contribution of sequences outside of the Switch I to RRSP substrate specificity, an comparable chimera mimicking the Ras Switch I sequence (rRalA:S1:Ras) was also generated by substituting RalA residues 41–48 with corresponding residues of Ras residues 30–37 ((E41D/D42E/E44D/K47I/A48E)(Fig. 4A,B). Remarkably, rRalA:S1:Ras was efficiently cleaved in 10 minutes, similar to rKRas, demonstrating that swapping the entire Ras Switch I into a non-substrate GTPase converted it to a suitable substrate for RRSP (Fig. 4C). A time course experiment revealed that rrRRSP had about 2.6-fold reduced catalytic activity (\( k_{\text{obs}} \)) for the Ras Switch I when presented by RalA (rRalA:S1:Ras) compared to the natural rKRas substrate (Fig. 4D, Table 1). Thus, a RalA backbone has only a minimal effect on processing efficiency and further confirms that RalA is not cleaved due its Switch I sequence, not due to the backbone interactions with RRSP.

RalA is a close relative of Ras within the Ras subfamily perhaps accounting for its ability to serve as a presenter of the Ras Switch I to RRSP if backbone interactions impart some specificity of RRSP for Ras and Rap1. To further test the importance of Ras Switch I compared to the backbone, the RhoA Switch I sequence was swapped with the Ras Switch I...
sequence (RhoA:S1:Ras) (Fig. 5A). As previously shown, rRhoA was not cleaved by RRSP. Furthermore, rKRas:S1:Rho was not cleaved, indicating that RRSP does not recognize RhoA Switch I even when presented by KRas. By contrast, when the corresponding amino acid positions of RhoA Switch I (residues 31–41) were substituted into KRas (Fig. 5A) RRSP processed rRhoA:S1:Ras about 25% after 10 minutes (Fig. 5B). Interestingly, this shows that compared to rRalA:S1:Ras, which was effectively cleaved in 10 minutes, RRSP is less able to recognize the Ras Switch I when presented in the context of RhoA. Indeed, in a time course experiment, rRhoA:S1:Ras was slowly processed, reaching 90% of cleavage product formation only after 8 hours (Fig. 5C and Table 1). In addition, RRSP for rRhoA:S1:Ras ($k_{\text{obs}}$ 0.0049 min$^{-1}$), was ~185-fold and ~480-fold lower than rRalA:S1:Ras and rKRas, respectively. The partial cleavage of rRhoA:S1:Ras suggests that the RhoA structural region outside of Switch I does not contribute to RRSP specificity, while the more closely related GTPase RalA does provide these interactions. Thus, RRSP substrate recognition is predominantly driven by the Switch I sequence but the efficiency of processing is also influenced by additional protein-protein interactions.

**RRSP cleaves Ras in the active (GTP-bound) and inactive (GDP-bound) states**

The structural conformation of Switch I when bound to nucleotide is known to influence the recognition of downstream effectors and bacterial toxins. In particular, structural studies of Ras revealed residues Y32 and D33 dramatically change their orientation, depending on whether Ras is bound to GDP or GTP (Fig. S1). To define whether RRSP specificity is nucleotide dependent explaining in part the sequence specificity of Switch I, rKRas proteins loaded with GDP and GppNHp, a non-hydrolysable GTP analog were used. Both GDP- and GppNHp-rKRas were mixed at different molar ratios with increasing amounts of recombinant RRSP (rRRSP). At equimolar ratio GDP-rKRas and GppNHp-rKRas were efficiently cleaved by rRRSP (Fig. 6A-inset). Also, at molar ratio 1:10 and 1:100 (rRRSP:rKRas), rRRSP cleaved both GppNHp- and GDP-bound KRas (Fig. 6A). At molar ratio 1:1000 partial processing was observed of about 80% for GppNHp-rKRas and only 50% for GDP-rKRas (Fig. 6A), indicating a slight preference for GppNHp-KRas as substrate. To quantify this slight difference at molar ratio 1:1000, a time course experiment was conducted to determine the cleavage rates of RRSP for both nucleotide-bound forms. For a reaction with only 0.1 μM rRRSP, GppNHp-rKRas was processed 3-fold faster than GDP-rKRas, although after one hour both rKRas forms were completely cleaved. Thus, the nucleotide state and the orientation of Y32 and D33 on Switch I had only a minor effect on rRRSP cleavage kinetics, demonstrating that RRSP is able to process both GDP- and GTP-bound forms.

**RRSP cleaves peptides corresponding to Ras and Rap1 Switch I**

All total, our data suggest that RRSP has sequence specificity for the Ras and Rap1 Switch I with GTPase backbone and bound nucleotide contributing to only efficiency. If this is the case, then a peptide alone should be processed by RRSP without a backbone or nucleotide. Peptides corresponding to the Switch I (residues 28–38) of Ras and Rap1 (peptides Ras-S1 and Rap1-S1, respectively) were synthesized and incubated with recombinant RRSP (rRRSP). Each peptide was incubated at equimolar ratio with 10 μM rRRSP at 37°C,
reactions were quenched after 0, 1, 2, 5 and 8 h and the cleavage products were analyzed by liquid chromatography-mass spectrometry (LC-MS). Both Ras-S1 and Rap1-S1 were completely processed after 2 h incubation with rRRSP. The reaction products of Ras-S1 in the presence of rRRSP revealed the formation of a smaller peptide (Fig. 7A, peak 1) with an observed mass of 671.3 Da, corresponding to the N-terminus cleavage product peptide FVDEY. Similarly, a product with observed mass of 684.3 Da (FVEKY) was detected for Rap1-S1 when incubated with rRRSP (Fig. 7B, peak 3). Since RalA Switch I is not a substrate of RRSP (Fig. 5A) a peptide corresponding to Ral Switch I (Ral-S1) was also tested as a substrate. As expected, no cleavage product was detected for Ral-S1 even after 8 hours at 37°C (Fig. 7C, peak 5). Further, exchange of the Ras scissile bond residues to Y32A/D33A generated a non-cleavable peptide. Thus, sequences that are not substrates in the context of intact Ras are also not substrates as peptides. This demonstrates that amino acid differences in the Switch I are sufficient to determine RRSP specificity, discriminating between substrate and non-substrate GTPases. However, the lower efficiency of peptide processing compared to processing of the whole GTPase proteins underscores that the nature of the GTPases and its nucleotide state can contribute to efficiency.

**DISCUSSION**

Small GTPases regulate a variety of cellular mechanisms that span from cell proliferation and differentiation to vesicle trafficking and apoptosis. Because of their central role in major cellular pathways, small GTPases are naturally involved in cell defense mechanisms of the immune system, in particular in the activation of the innate immune response against external pathogens. Host cells use pattern recognition receptors, such as Toll-like receptors (TLRs), to sense pathogen-associated molecular patterns and to activate the immune system. Bacteria are recognized through TLR2, TLR4 and TLR5 by binding lipoproteins, lipopolysaccharide and flagellin, respectively. This activates the innate and adaptive immune responses, eliminating the pathogens and establishing protective immunity against them. However, bacterial pathogens, through the use of secreted toxins, have developed specific ways to destroy the eukaryotic cell cytoskeleton and to reduce the host response against bacterial infection by inhibiting or completely inactivating the small GTPases. Among these toxins, some have broad specificity targeting many small GTPases, while others show more limited substrate preference.

RRSP was recently identified as the first member of an uncharacterized endopeptidase family able to selectively cleave the Switch I of Ras and Rap1 small GTPases, blocking Ras-MAPK signaling. In this study, we found that theSwitch I of either Ras or Rap1 was the minimal sequence required for RRSP recognition. The final demonstration of cleavage of Ras-S1 or Rap1-S1 peptides by RRSP demonstrated that the primary sequence of Switch I between residues 28–38 was sufficient for RRSP to discriminate substrates and identify the scissile bond between Y32 and D33. Since the Switch I sequence is directly involved in the interaction with downstream effectors, the specific recognition of RRSP to this region might indicate similar surface contact. In fact, Ras Switch I residues 30 and 31 are known to be critical for the specific binding to BRaf, a downstream effector of Ras. Interestingly, changing residues 30 and 31 in Rap1 to those found in the corresponding positions of Ras lead to favorable interaction between Rap1 and BRaf in vitro. We found that Y32 and D33
play important roles in RRSP activity, as their substitution with alanine residues entirely inhibits the cleavage of Switch I. Although Y32 and D33 at the scissile bond are highly but not completely conserved among other GTPases, RRSP has been shown to cleave only Ras and Rap1\textsuperscript{22}. Indeed, the scissile pair does not fully account for substrate specificity, as the Y-E bond found in RalA was a suitable bond when substituted into Ras. In addition, swapping individual residues found in RalA into Ras failed to identify a single residue that would result in complete RRSP loss of activity. Rather, we found that it is the whole Switch I primary sequence, comprised of residues 28–38 that defines the RRSP specificity.

This recognition of the Switch I sequence is a distinct strategy compared to other toxins that target the Switch I of GTPases. The well-characterized toxins TcdA and TcdB of the pathogen \textit{Clostridium difficile} use UDP-glucose as the sugar donor to mono-O-glucosylate Rho proteins, including RhoA, RhoB, RhoC, RhoG, Rac1 and Cdc4\textsuperscript{23}. However, recently Zaiser \textit{et al} using a proteomic approach have found that TcdA and TcdB also glucosylate Ras proteins (K/H/N) and other Ras subfamily proteins like Rap and Ral\textsuperscript{51}. \textit{Clostridium sordellii} Lethal Toxin (LT or TcsL) uses mono-O-glucosylation to modify GTPases from both the Rho proteins, like Rac1 and Ras subfamily proteins, including Rac, Ras, Rap and Ral\textsuperscript{43,52}. The TpeL toxin of \textit{Clostridium perfringens} is more stringent in its recognition and targets Ras proteins by mono-O-GlcNAcylation using UDP-N-acetylglucosamine (UDP-GlcNAc) as the sugar donor\textsuperscript{53}. All these clostridial toxins mono-O-glucosylate the conserved threonine in the Switch I (T35 or T37 of Rho or Ras proteins, respectively). Glucosylation of T35 and T37 impair GTPase activity, blocking their signaling functions\textsuperscript{54–56}. Since the targeted threonine residue within the Switch I is 100% conserved across all GTPases\textsuperscript{23}, sequence specificity of the modified residue cannot account for the substrate preference of these clostridial toxins.

Instead, for these other toxins, specificity is driven in part by conformation change due to the bound nucleotide. LT preferentially modifies Rho and Ras in GDP-form, as the target hydroxyl group of the threonine residue directly coordinates the magnesium cation and binds the γ-phosphate of the GTP\textsuperscript{43}. By contrast, PaTox of \textit{Photorhabdus asymbiotica} GlcNAcylation of GDP-loaded RhoA is poor while GTP-RhoA is efficiently modified in the Switch I. Interestingly, PaTox mono-O-glucosylates Y34 of RhoA, the equivalent residue Y32 of Ras, indicating that Y34 is in a favorable and exposed conformation to be modified when RhoA is in its active form\textsuperscript{42}. This precedence established for the glucosyltransferase toxins to discriminate GTP- and GDP-bound forms of target GTPases is found to be quite different for RRSP. In this work, we showed that RRSP efficiently cleaved both GDP- and GTP-KRas, demonstrating that differences in the orientations of Ras Y32 due to nucleotide binding had only a minor 2.8-fold effect on Switch I recognition. In this way, RRSP is more similar to the \textit{Clostridium botulinum} C3 exoenzyme, which catalyzes the ADP-ribosylation of N41 of RhoA/B/C. C3 is capable of modifying both GTP- and GDP-bound forms of Rho proteins. Similarly to RRSP, the nucleotide occupancy does not affect the enzyme recognition but only the kinetics\textsuperscript{57}. However, since the nucleotide state of any small GTPase is crucial for association with downstream effectors or guanosine nucleotide dissociation inhibitors in the cell, it is possible RRSP might have different substrate accessibility in the cellular environment due to binding of other proteins.
Among the many toxins that target small GTPases, substrate specificity is also dictated by interactions outside of the Switch I region, but structurally nearby. For TcdB and LT, a single amino acid modification within the Switch II region of RhoB and RhoD was sufficient to change the target specificity of both toxins. The adenyltransferase IbpA of *Histophilus somni* also targets the Switch I, by catalyzing the addition of adenosine monophosphate to Y34 of RhoA and to Y32 of both Cdc42 and Rac1. Similarly to TcdB and LT, IbpA also recognizes Rho-family GTPases by binding to the Switch II region. Moreover, C3 exoenzyme from *Clostridium botulinum*, which specifically modifies RhoA/B/C but not other members of either Rho or Ras subfamilies, recognizes Switch II and interswitch regions in addition to Switch I.

Unlike these other toxins, the specificity of RRSP is largely driven by the sequence of Switch I as shown when the entire Ras Switch I was exchanged into RalA, RalA became a substrate for RRSP and the rate of processing decreased only slightly. Yet, when exchanged into RhoA, the processing rate declined dramatically. This suggests that, in addition to a different presentation of the Ras Switch I by RhoA, there is a contribution of the protein backbone to substrate recognition. However, this result also indicates that the Ras Switch I is strictly required for substrate specificity. Notably, the complete cleavage of both Ras-S1 and Rap-S1 peptides by RRSP required two hours, while ten minutes was sufficient for cleavage of intact KRas protein. These data further demonstrate that although the sequence recognition of the Switch I region is mainly responsible for the RRSP specificity, possible additional interactions with the whole GTPase structure increases the recognition and efficiency of RRSP. Indeed, the next question that will need to be addressed is how these other regions of Ras and Rap1 outside the Switch I can contribute to a second level of specificity.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

**Acknowledgments**

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**ABBREVIATIONS**

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tr>
<td>RRSP</td>
<td>Ras/Rap1-specific endopeptidase</td>
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</table>
**References**


Figure 1.
Switch I sequence alignment and RRSP cleavage site. A Alignment of amino acid sequences of the Switch I sequence of several small GTPases by Clustal Omega and ESPript 3. The black arrow between Y32 and D33 indicates the RRSP processing site. The RRSP protein substrates (Ras/Rap1) are located above the black dashed line. Identical amino acid residues are highlighted in red. The accession numbers of amino acid sequences were obtained from NCBI: KRas (NP_004976), HRas (NP_001123914), NRas (NP_002515), Rap1A (CAA00804), Rap1B (CAA00805), RalA (NP_005393), RalB (AAA60250), RhoA (NP_001300870) and RhoB (NP_004031). B Crystal structure of KRas-GDP (PDB 4OBE) prepared using PyMOL Molecular Graphics System, Version 1.8 (Schrödinger, LLC). Switch I region is colored in blue and the amino acid Y32 and D33 next to RRSP processing site are shown as stick models.
Figure 2.
Scissile bond residues in KRas Y32 are important for RRSP specificity. A–C SDS-PAGE analysis of *in vitro* cleavage products of 10 μM rKRas as indicated incubated for 10 min at 37º C with and without 10 μM rRRSP. B–D Time course experiment to assess RRSP activity for rKRas as indicated. Each curve represents the average and standard deviation of three independent biological replicates. WT KRas curve is the same in panel B and D because experiments were conducted simultaneously.
Figure 3.
Residues downstream of Y32 are important but not essential for RRSP substrate recognition. A Illustration of the residues of the Ras Switch I region substituted with amino acid corresponding to Ral Switch I. KRas-GDP switch I is colored blue and residues substituted with RalA Switch I amino acids are colored yellow and shown as stick models. B SDS-PAGE analysis of in vitro cleavage products of 10 μM rKRas mutants incubated for 10 minutes at 37°C with and without 10 μM rRRSP, representing at least 3 independent experiments.

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Figure 4.
Ras Switch I drives the specificity of RRSP. A Superimposition of crystal structures of KRas-GDP (blue) (PDB 4OBE 63) and RalA-GDP (yellow) (1U8Z 64). Switch I residues are shown as stick models and annotated accordingly to their positions. B Scheme of KRas/RalA chimera constructs designed, expressed and purified. C SDS-PAGE analysis of in vitro cleavage products of 10 μM of rKRas, rRalA, KRas:S1:RalA and RalA:S1:Ras incubated for 10 min at 37°C with and without 10 μM rRRSP. D Time course experiment to assess RRSP activity for recombinant protein substrates, as described for Fig. 2.
Figure 5. RRSP cleaves RhoA with Ras Switch I. A Scheme of KRas/RhoA chimera constructs designed, expressed and purified. B SDS-PAGE analysis of in vitro cleavage products of 10 μM of rKRas, rRhoA, KRas:S1:Rho and RhoA:S1:Ras incubated for 10 min at 37°C with and without 10 μM rRRSP. C Time course experiment to assess RRSP activity for recombinant protein substrates, as described for Fig. 2.
Figure 6.
RRSP cleaves Ras in the active (GTP-bound) and inactive (GDP-bound) states. A In vitro cleavage of 10 μM rKRas with 10 μM rRRSP (inset) or concentration indicated. B Time course experiment to assess RRSP activity for GDP-rKRas and GppNHp-rKRas as indicated. Each curve represents the average and standard deviation of three independent biological replicates. Each curve present in A and B represents the average and standard deviation of three independent biological replicates.
Figure 7.
RRSP cleaves Ras and Rap1 Switch I peptides. In vitro cleavage assay using LC-MS of Switch I peptides was performed at equimolar concentration with rRRSP (10 μM) at different time points. A Results of Ras Switch I peptide (Ras-S1); B Results of Rap1 Switch I peptide (Rap1-S1); C Results of Ral Switch I peptide (Ral-S1). Upper and middle panels show the retention times of the cleavage products Switch I peptides incubated with and without rRRSP. Incubation times are color-coded: time zero (black), 1 h (red), 2 h (green), 4 h (violet) and 8 h (purple). Lower panels indicate the observed and calculated monoisotopic mass values for each Switch I peptides and their cleavage product. D In vitro cleavage assay of Ras Y32A/D33A Switch I peptide was performed at equimolar concentration with 10 μM rRRSP for time indicated.
Table 1
Catalytic activity of RRSP ($k_{\text{obs}}$) for different substrates.

<table>
<thead>
<tr>
<th>Substrate</th>
<th>RRSP $k_{\text{obs}}$ (min$^{-1}$)</th>
<th>RRSP activity reduction (fold-change)</th>
</tr>
</thead>
<tbody>
<tr>
<td>rKRas</td>
<td>2.35 ± 0.14</td>
<td>-</td>
</tr>
<tr>
<td>rKRas Y32A</td>
<td>0.031 ± 0.0054</td>
<td>75.8</td>
</tr>
<tr>
<td>rKRas D33A</td>
<td>0.017 ± 0.0019</td>
<td>138</td>
</tr>
<tr>
<td>rKRas Y32F</td>
<td>0.094 ± 0.0079</td>
<td>25.0</td>
</tr>
<tr>
<td>rKRas Y32A/D33A</td>
<td>undetectable</td>
<td>-</td>
</tr>
<tr>
<td>rKRas Y32F/D33A</td>
<td>0.00073 ± 0.0014</td>
<td>2350</td>
</tr>
<tr>
<td>rRalA</td>
<td>undetectable</td>
<td>-</td>
</tr>
<tr>
<td>rKRas:S1:Ral</td>
<td>0.0041 ± 0.0027</td>
<td>573</td>
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<tr>
<td>rRalA:S1:Ras</td>
<td>0.91 ± 0.095</td>
<td>2.58</td>
</tr>
<tr>
<td>rRhoA</td>
<td>undetectable</td>
<td>-</td>
</tr>
<tr>
<td>rKRas:S1:Rho</td>
<td>undetectable</td>
<td>-</td>
</tr>
<tr>
<td>rRhoA:S1:Ras</td>
<td>0.0049 ± 0.0074</td>
<td>479</td>
</tr>
</tbody>
</table>

RRSP $k_{\text{obs}}$ values were determined from time course experiments plots as shown in Fig. 2, 4, 5 and 7. and are presented as the average and standard deviation from three different experiments. All values reported are for 10 μM RRSP incubated with 10 μM indicated substrate for up to 8 hr.