Maize Ribosome-Inactivating Protein Uses Lys158–Lys161 to Interact with Ribosomal Protein P2 and the Strength of Interaction Is Correlated to the Biological Activities

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Abstract

Ribosome-inactivating proteins (RIPs) inactivate prokaryotic or eukaryotic ribosomes by removing a single adenine in the large ribosomal RNA. Here we show maize RIP (MOD), an atypical RIP with an internal inactivation loop, interacts with the ribosomal stalk protein P2 via Lys158–Lys161, which is located in the N-terminal domain and at the base of its internal loop. Due to subtle differences in the structure of maize RIP, hydrophobic interaction with the ‘FGLFD’ motif of P2 is not as evidenced in MOD-P2 interaction. As a result, interaction of P2 with MOD was weaker than those with trichosanthin and shiga toxin A as reflected by the dissociation constants \( K_D \) of their interaction, which are \( 103 \pm 6.75 \mu M, 61.70 \pm 28.13 \mu M \) and \( 194.84 \pm 9.47 \mu M \) respectively. Despite MOD and TCS target at the same ribosomal protein P2, MOD was found 48 and 10 folds less potent than trichosanthin in ribosome depurination and cytotoxicity to 293T cells respectively, implicating the strength of interaction between RIPs and ribosomal proteins is important for the biological activity of RIPs. Our work illustrates the flexibility on the docking of RIPs on ribosomal proteins for targeting the sarcin-ricin loop and the importance of protein-protein interaction for ribosome-inactivating activity.

Introduction

Ribosome-inactivating proteins (RIPs) comprise more than one hundred \( \alpha \)-glycosidases [1] which cleave the \( \alpha \)-glycosidic bond of adenine-2660 at 23S rRNA or adenine-4324 at 28S rRNA of ribosome specifically and irreversibly [2–3]. The corresponding adenine is situated at the \( \alpha \)-sarcin ricin loop (SRL) which belongs to the GTPase centre of ribosome and actively involved in the interaction with the elongation factors [4]. Upon depurination of the SRL by RIPs, elongation factors fail to form a stable complex with ribosome and thus delivery of aminoacyl-tRNA to ribosome and translocation of peptide are prohibited, inevitably causing the cease of protein synthesis and cell death [5–7].

At present, about 30 RIPs have their structures solved and it is found that the structures and catalytic residues of RIPs are highly similar. Typically, the globular RIP is divided into a large N-terminal domain and a small C-terminal domain, creating an active cleft holding the five invariant catalytic residues [8–9]. Despite sharing highly conserved structure and catalytic centre, the potency and specificity of RIPs on ribosome vary [10–12].

Ricin has been demonstrated to depurinate both 23S and 28S rRNA isolated from the ribosome at the conserved SRL with similar \( k_m \) and \( k_{cat} \) values although it cannot inactivate the prokaryotic ribosome. In addition, the \( k_{cat} \) value of depurination using eukaryotic ribosome as substrate is over 80,000 folds higher than using naked 28S rRNA [13]. These indicate ribosomal proteins are involved in the biological activity of RIPs.

Among the studies aiming at finding the interacting ribosomal proteins, human P-proteins have been found in several occasions. Trichosanthin (TCS), shiga-like toxin 1 (SLT-1) and ricin A chain (RTA) have been shown to interact with the last eleven residues of human P-proteins [14–16]. Also, RTA has been crosslinked to human L10e [17] which is identical to the P-protein P0.

On the contrary, pokeweed antiviral protein (PAP) has been found to interact with yeast ribosomal protein L3 [18–19] and not require the C-terminus of P-proteins for its ribosome depurinating activity [20]. It has also been found that yeast L3 with Trp-255 and Pro-257 mutated is resistant to PAP while those deprived of P1 and P2 are less vulnerable to the deleterious effect of RTA and shiga toxin [18,21–22].

To understand why RIPs exhibit different ribosome specificity despite their highly similar structure, several regions of major structural difference between PAP and RTA were swapped and tested. The swapping weakened but did not totally abolish the
prokaryotic ribosome deaminating activity of PAP, suggesting the dissimilar regions alone are not responsible for the ribosome specificity [23]. Conversely, substituting Gly209-Lys225 in PAP with the corresponding region of karasurin-A greatly reduced the prokaryotic ribosome deaminating activity of PAP [24]. Alignment of amino acid sequences of PAP and karasurin-A has revealed a higher charge density in PAP at this region but whether the charges contribute to the prokaryotic ribosome specificity of PAP remains unknown.

Our group has previously solved the structure of TCS complexed with a 11 amino acid peptide (C11-P) corresponding to the conserved C-terminus of P-proteins. This C11-P peptide could fit to the corresponding groove of RTA, STL-1 and saporin but not PAP, implying PAP may get access to the ribosome through a different route that does not require any P-protein [25].

Recently, we have also solved the crystal and solution structures of maize RIP [26–27]. Maize RIP is an atypical RIP that contains a 25-amino acid internal peptide in the precursor form (pro-RIP). The internal peptide is absent in most RIPs and represses the activity of maize RIP by at least 600 folds [28–29] and pro-RIP displays a lower affinity to rat liver ribosome in comparison to the active form [26]. We postulated the internal peptide creates a steric hindrance causing a crash between the protein and rRNA, thereby prohibiting the protein to inactivate the ribosome and reducing its activity [26]. By chemical shift perturbation NMR experiment, we have shown that amino acid residues at the N-terminal domain of MOD, the active form of maize RIP, are perturbed upon the addition of ribosomal protein P2.

In this report, we demonstrate that among all the positively charged residues on the surface of MOD, only Lys158-Lys161 take part in the interaction with P2. The P2-binding site of MOD is very different from other known RIP-P2 interaction. The affinity of MOD to P2 is weaker than those of TCS and shiga toxin A chain by 1.7 folds and 5.3 folds respectively and is likely due to the lack of hydrophobic interaction with P2. Additionally, MOD was found less potent than TCS in N-glycosidase activity and cytotoxicity by 48 and 10 folds respectively. Our findings illustrate RIPs can use various regions to dock on the same ribosomal target in response to the subtle differences in the structures and the affinity between RIPs and ribosome plays a role in the activity of the RIPs.

Materials and Methods

Cloning and site-directed mutagenesis of RIPs

Maize [A1–16, Δ287–300]-Pro-RIP (Pro-RIP-WT in short) and [A1–16, Δ163–164, Δ167–189, Δ287–300]-Pro-RIP (MOD-WT in short) were prepared as described [26]. MOD [K158A-K161A] was constructed by polymerase chain reaction (PCR) using mutagenic primers and subsequently cloned into pET3a (Novagen). For His-Myc-tagged MOD (Myc-MOD), nucleotides coding for the myc tag were inserted at the 3’ of the gene of MOD by PCR. The construct was then cloned into pET28a (Novagen). Trichosanthin (TCS) was cloned into pET3d (Novagen) as described [30]. His-SUMO-tagged shiga toxin A chain inactive mutant (StxA [E167A, R170A], StxA in short) was cloned into pRSETA (Invitrogen) pre-cloned with DNA encoding His-SUMO. C-terminal His-tagged MOD and TCS were cloned into pET28a while P2 and its variants P2 [ΔC5], P2 [ΔC10] were cloned into pET11c similarly as described [14].

Protein expression and purification

Maize RIP variants, TCS and P2 variants were expressed and purified as described before [14,26,30]. His-myc tagged MOD and C-terminal His-tagged MOD and TCS were expressed in E. coli BL21(DE3)pLysS under the induction of 0.4 mM IPTG. The bacterial pellets were resuspended and sonicated in His buffer A (20 mM sodium phosphate buffer, 800 mM NaCl, 50 mM imidazole, pH 7.4), the soluble lysate was then subject to HisTrap High Performance column (GE Healthcare) for affinity purification. The proteins were then eluted using His buffer B (20 mM sodium phosphate buffer, 300 mM NaCl, 300 mM imidazole, pH 7.4).

His-SUMO-StxA was expressed in E. coli BL21(DE3)pLysS at 25°C for 5 hours under the induction of 0.4 mM IPTG and was firstly isolated as mentioned above. Eluate was then treated with His-SUMO-protease for removing the His-SUMO tag and dialysed against His buffer A at 4°C for 16 hours before loaded to HisTrap to recover the protein from the flow through.

Purification of rat liver ribosome

Rat liver ribosome was prepared according to the protocol in Spedding [31] and a brief summary was given in Mak et al. [26]. The quality of isolated ribosome was checked by measuring its absorbance at 260 nm and 280 nm and a ratio of A260/A280 higher than 1.8 indicates a clean ribosome preparation was achieved.

In vitro pull-down assay with ribosome

MOD, pro-RIP and MOD variants were diluted in coupling buffer (200 mM NaHCO3, 500 mM NaCl, pH 8.3) before immobilization. 150 μl suspension of NHS-activated Sepharose 4 Fast Flow (GE Healthcare) was rinsed with 10 bead-volumes of ice-cold 1 mM hydrochloric acid (HCl) and 2 mg MOD variant was added to the beads immediately after removal of HCl. After incubating for an hour, the protein solution was aspirated and unreacted succinimide ester on the beads was inactivated according to the instruction of the manufacturer. The beads were then washed with 1 ml elution buffer (20 mM sodium phosphate buffer, 1 M NaCl, pH 7.4) and equilibrated in phosphate buffer saline (PBS) for 15 minutes. 0.3 mg ribosome in 800 μl was then loaded to the beads for interaction with maize RIP variants for an hour. Following washing the beads extensively with PBS, the retained proteins were eluted using elution buffer and examined with SDS-PAGE.

Crosslinking assay and western blotting

All RIPs and rat liver ribosome had their buffer exchanged to PBS using Amicon Ultra-15 prior to the crosslinking. Myc-MOD and a buffer control were labelled with 5-molar excess of crosslinker NHS-SS-Diazirine (SDAD, Pierce) for an hour. Then the unreacted succinimide ester of SDAD was quenched with 50 mM Tris and uncoupled SDAD was removed by Amicon Ultra-0.5. Labelled RIPs were then mixed with ribosomes at 90 μM and 0.8 μM respectively in PBS and allowed to interact at room temperature for 30 minutes in dark. Controls with either RIP or ribosome supplemented with SDAD-pre-treated buffer were set up at equal protein amount as stated above. The reaction mixtures were irradiated for an hour under a UV lamp (Spectrolite) emitting light of 365 nm to activate the diazirine group to crosslink the interacting partners.

For MOD-P2 crosslinking, P2 was labelled by 2-molar excess of SDAD and incubated with MOD at equal concentration of 0.4 μM for 30 minutes and followed by an irradiation of an hour. Controls with MOD or P2 alone were set up as well.

After crosslinking, the samples were concentrated and subject to western blot of which the membrane was probed with monoclonal mouse anti-Myc (Cell Signaling Technology), in-house polyclonal
rabbit anti-MOD and anti-P proteins (anti-P) antibodies and horseradish peroxidase conjugated goat anti-mouse (Bio-rad) and anti-rabbit (Invitrogen) antibodies.

**Surface plasmon resonance**

BLAcore 3000 surface plasmon resonance biosensor (GE healthcare) was used for the kinetic study of RIP-P2 interaction. CM5 sensor chip was activated using the amine coupling kit (GE healthcare). In brief, P2 was diluted to 50 μg/ml with 10 mM sodium acetate, pH 3.8 and 40 μl of the sample was injected to the chip at a flow rate of 5 μl/min to obtain a final response unit (RU) of 1300. Ethanolamine was then injected to block the unreacted succinimide ester. A control flow cell was prepared with identical treatments but excluding P2 immobilization.

**In vitro pull-down assay of RIPS with P2 and variants**

2 mg of P2, P2 [AC5] or P2 [AC10] were immobilized to the NHS-activated Sepharose and MOD, pro-RIP, MOD [K158A-K161A], TCS or StxA was added at 55 μM in binding buffer (20 mM Tris, 30 mM NaCl, 0.005% Tween 20, pH 7.4) and diluted to appropriate concentration for the kinetic analysis. Samples were injected at 30 μl/min for 3 to 5 minutes to attain equilibrium and running buffer was injected for 3 minutes to allow dissociation. The chip was then regenerated using 2 M NaCl. Triplicate was performed for every concentration of sample to obtain three independent RU. The RU values at equilibrium of every concentration were taken for fitting into the equation modified from the Michaelis-Menten equation by KalcidaGraph (Synergy software):

\[ y = \frac{R_{\text{max}} \times x}{K_D + x} \]

where \( R_{\text{max}} \) is the maximum response unit and \( K_D \) is the dissociation constant.

**N-glycosidase activity assay and quantitative PCR**

50 μg rat liver ribosome was incubated with 5 nM MOD, pro-RIP, MOD [K158A-K161A] or TCS for 20 minutes at 30°C in the assay buffer A [20 mM Heps (pH 7.6), 150 mM KCl, 10 mM MgCl2, 40 mM NH4Cl] at a final volume of 25 μl. 400 μl DEPC-treated water, 500 μl Trizol (Invitrogen) and 200 μl chloroform were added for phase separation of rRNA and proteins. After centrifugation, rRNA in the upper aqueous phase was precipitated with isopropanol and washed with 75% ethanol. The RNA pellet was then resuspended with DEPC-treated water and 500 ng of RNA was taken for cDNA synthesis using SuperScript II Reverse Transcriptase Kit (Invitrogen). The cDNA products were diluted for 10 folds for subsequent quantitative PCR (qPCR) using primers and protocol as described before [32–33] except 1 μl but not 4 μl of diluted cDNA was added. The relative N-glycosidase activity: Number of depurinated 28S rRNA at SRL/Number of total 28S rRNA, could be calculated from the threshold cycle (Ct) values. The data were illustrated and statistically analyzed using unpaired t-test by Prism 5 (GraphPad Software).

### Results

**Results**

All four residues of Lys158–Lys161 are involved in ribosome interaction

To examine whether ionic interaction is important for MOD-ribosome interaction, we have screened all the positively charged residues at the surface of MOD for their ability in binding ribosome.

A total of 17 variants were constructed with cluster of closely located basic residues mutated to alanine. Pull-down assay showed that all variants were able to interact with ribosome at similar extent as the wild type except MOD [K158A-K161A] (Figure 1), suggesting these four continuous lysine are responsible for the interaction with ribosome.

**MOD interacts with P2 on rat liver ribosome**

By chemical perturbation experiment, we have shown that P2 may interact with the N-terminal domain of MOD [27]. Here, we examined if maize RIP really targets at P2 on the ribosome. SDAD-labelled Myc-tagged MOD was incubated with rat liver ribosome for interaction and permanent crosslinking was elicited by activation of the diazirine group of SDAD upon UV irradiation. Crosslinker SDAD (spacer-arm of 13.5 Å) was employed since it has two different chemical groups, N-hydroxysuccinimide ester and photoactivatable diazirine, at two ends which allow a two-step crosslinking to avoid extensive self-
crosslinking of ribosomal proteins that increases the complexity of western analysis.

Diazirine group of the crosslinker can react with any amino acid side chain or peptide backbone and thus inevitably generates a small amount of non-specific crosslinked bands which lead to undesired background on the western blot. Therefore, a C-terminal Myc-tagged variant of MOD was prepared for reducing the noise of the western blot, as the monoclonal antibody against the Myc-tag is more specific than the polyclonal anti-MOD antibody.

The membranes were probed to detect MOD and P-proteins (Figure 2). In lane 2, a band of about 42 kDa could be detected on both blots, but absent in the negative controls (lanes 1 and 3). The size of the product coincided with the complex composed of one Myc-MOD and one P2 (42.3 kDa). Alternatively, crosslinking was also performed on mixture of MOD and P2 which yielded a band of slightly smaller size on both blots (MOD and P2 give a total size of 39 kDa), proposing MOD likely interacts with P2 when it encounters the ribosome.

As sizes of P-proteins P1 and P2 are similar (11.5 and 11.7 kDa), mass spectrometric analysis was employed to confirm whether the crosslinked partner is P2. Crosslinking was repeated on MOD immobilized to the resin and the crosslinked partner was collected by reducing the disulphide bond of the crosslinker and identified by mass spectrometry. The m/z spectrum of the product was found to match with that of P2 and thus MOD was crosslinked to P2 instead of P1 (data not shown).

Lys158–Lys161 on Maize RIP are responsible for binding to P2

We next examined if maize RIP establishes a direct interaction with P2 via Lys158–Lys161. With same amount of P2 immobilized to the sepharose, identical amount of maize RIP variants were added for binding. The eluates were collected and analyzed by SDS-PAGE (Figure 3A) which showed that MOD but not pro-RIP could retain P2. The interaction between MOD [K158A–K161A] and P2 was highly diminished, implying that alanine substitution of these residues significantly impaired the interaction.

Surface plasmon resonance was employed to quantitatively compare the interaction between maize RIP and P2 (Figure 3B). MOD gave the highest response among the analytes tested with a $K_D$ of $1037.50^{\pm}65.75$ M while the response of pro-RIP and MOD [K158A–K161A] were negligible and comparable to the buffer control.

### Table 1

|        | Myc-MOD | MOD |
|--------|---------|-----|
| MOD    | +       | +   |
| Ribosome | -       | +   |
| P2     | -       | -   |

| Protein | 4 | 5 | 6 |
|---------|---|---|---|
| MOD     | + | + | - |
| Ribosome | - | - | - |
| P2      | - | + | + |

Figure 1. Screening of basic residues on MOD that are responsible for ribosome binding. The indicated residues were mutated to alanine and screened for their abilities to bind ribosome. W and E denote last wash and elution respectively.

doi:10.1371/journal.pone.0049608.g001

Figure 2. MOD can be crosslinked to rat liver ribosome and P2. Crosslinking reactions were carried with individual proteins (lanes 1 and 3, 4 and 6) or mixtures of two proteins (lanes 2 and 5) and subject for western analysis. Protein bands were detected by anti-myc, anti-MOD and anti-P antibodies.

doi:10.1371/journal.pone.0049608.g002
Pro-RIP and MOD [K158A–K161A] were less potent in ribosome depurination and cytotoxicity to 293T cells

Next, we measured the relative depurinating activities of maize RIP and its variants on naked 28S rRNA and rat liver ribosome by estimating the relative amount of depurinated 28S rRNA and total 28S rRNA.

It is observed that MOD, pro-RIP and MOD [K158A–K161A] cleaved naked 28S rRNA to a similar extent. Conversely, pro-RIP and MOD [K158A–K161A] negligibly depurinated 28S rRNA embedded in the ribosome (Figure 4A and Table 1). These suggested the three forms of maize RIP possess comparable catalytic activity and the difference in ribosome depurinating activity was due to their different affinity to P2. Consistently, pro-RIP and MOD [K158A–K161A] were much less toxic to 293T cells than MOD, as the former two variants did not show cytotoxicity up to 15 μM, while the IC₅₀ of MOD was 7.88±1.57 μM (Figure 4B).

Interaction of RIP-P2 under various ionic strengths

Respecting MOD, TCS and StxA share the same partner P2, the next interest is to verify whether these RIPs interact with P2 with comparable affinity. Pull-down experiments on P2-column using buffers of different salt concentrations were carried out by loading equal amount of RIPs to the column. It was found that the amount of RIPs bound on P2-column was reduced with increasing salt, implying the interaction between RIP and P2 is highly electrostatic. MOD was least resistant to ionic challenge as it failed to bind to P2 at 100 mM or higher concentration of NaCl while TCS and StxA could still bind to P2 at 150 mM NaCl (Figure 5A).

Figure 3. Lys158–Lys161 on MOD are responsible for the interaction between maize RIP and P2. A) MOD does not bind to P2 when Lys158–Lys161 are mutated to alanine. SDS-PAGE of the last wash (W) and elution (E) obtained from the pull-down assay of maize RIP and P2. Input indicates the purified proteins loaded to the column. B) Sensorgram showing MOD but not pro-RIP and MOD [K158A–K161A] interact with sensor chip-immobilized P2. 500 nM of maize RIP variants or running buffer were injected to the sensor chip for three minutes for association, followed by a dissociation using running buffer. Response unit (RU) was monitored in a real-time manner.
doi:10.1371/journal.pone.0049608.g003
Maize RIP does not bind to the hydrophobic residues at the P2 tail

To examine the role of the highly consensus C-terminus of P-proteins for interacting with maize RIP and other RIPs, P2 variants lacking its last five or ten conserved residues were constructed for the pull-down assay.

It was observed that removing the last five residues ‘FGLFD’ of P2 heavily reduced the band intensities for TCS and StxA but imposed no effect on MOD-P2 interaction, implying MOD does not rely on the last five hydrophobic residues for interaction (Figure 5B). When the last ten residues of P2 were truncated, interactions of all tested pairs were abolished, indicating that the hydrophilic motif DDDMG on P2 is indispensable for RIPs to bind P2.

MOD interacts with P2 at lower affinity than TCS and StxA

Surface plasmon resonance was carried out to quantify the strength of the interaction of each RIP-P2 pair. The $K_D$ values of StxA-P2, TCS-P2 and MOD-P2 interaction were estimated to be $194.84 \pm 6.94$ nM, $611.70 \pm 28.13$ nM and $1037.50 \pm 65.75$ nM respectively. The affinity between MOD and P2 is the weakest among three RIPs.

### Table 1. Relative N-glycosidase activities of maize RIP and its variants on 28S rRNA and rat liver ribosome.

| RIPs            | rRNA N-glycosidase activity (Mean ± S.D.) | Folds increased to untreated | Ribosome N-glycosidase activity (Mean ± S.D.) | Folds increased to untreated |
|-----------------|------------------------------------------|------------------------------|-----------------------------------------------|------------------------------|
| MOD             | 6.41 ± 5.52                              | 69.48                        | 54.26 ± 15.63                                | 14.28                        |
| Pro-RIP         | 4.51 ± 2.18                              | 48.94                        | 6.06 ± 5.92                                  | 1.59                         |
| MOD [K158A-K161A] | 3.58 ± 3.85                             | 38.85                        | 3.24 ± 2.28                                  | 0.85                         |
| Untreated       | 0.09 ± 0.06                              | /                            | 3.80 ± 2.94                                  | /                            |

doi:10.1371/journal.pone.0049608.t001
MOD is less active than TCS

Regarding the binding site and affinity of interaction of MOD and TCS with P2 are very different, we would like to further test on their biological activities to verify if the activities are related to their affinities with P2.

The relative N-glycosidase activities on ribosome of MOD and TCS were estimated to be $54.26 \pm 6$ and $2607.21 \pm 711.35$ respectively, reflecting MOD was about 48 folds less potent than TCS in depurinating the ribosome. On the other hand, the relative N-glycosidase activities on naked rRNA of MOD and TCS were found to be $6.41 \pm 5.52$ and $0.35 \pm 0.18$ respectively. This reflected TCS was 18 folds less active on rRNA than MOD. The lower potency of MOD on ribosome is consistent to its lower affinity to P2.

The cytotoxicity to 293T cells also demonstrated a higher potency of TCS than MOD. For a fair comparison of the cytotoxicity, a hexa-His tag was incorporated into the C-termini of MOD and TCS so that equal amount of purified RIPs was loaded to the P2-sepharose column and proteins were eluted using buffer with 1M NaCl. The left panel of Figure S1 illustrates the same amount (20 ng) of RIPs was loaded. The right panel shows comparable intensity for MOD and TCS and hence a similar degree of cellular uptake of these proteins could be assumed. As determined from the viability curves, the effective protein concentration that killed 50% of cells (EC$_{50}$) of MOD was $7.88 \pm 1.57$ \(\mu\)M while that of TCS was $0.63 \pm 0.32$ \(\mu\)M, implicating MOD was at least 10 folds less toxic to 293T than TCS.

Discussion

RIPs interact with the SRL on the 28S rRNA and remove a specific adenine residue. It is believed RIPs target at specific ribosomal protein that is in proximity to their rRNA target SRL. Upon binding to RIPs, conformation of ribosomal proteins alters and subsequently increases the accessibility of the RIPs to SRL. Thus, ribosomal proteins can enhance the activity of RIPs by facilitating the latter to bind to the ribosome as well as to locate their target.

As proposed earlier, the interaction between RTA and ribosome can be characterized by a two-step binding model, consisting of a faster and stronger interaction involving particular ribosomal protein(s) and a slower and weaker interaction between the ribosomal surface and RIPs which may involves multiple binding sites on a single ribosome [34]. Ribosomal protein is believed to facilitate the depurinating activity of RIPs by guiding them to the SRL [14,34]. Unable to bind to a target ribosomal protein(s) would inevitably compromise the interaction with ribosome as well as the activity of RIPs.

In the case of maize RIP, the P2-binding site on maize RIP was located at Lys158–Lys161, which is not at the C-terminus as in...
other RIPs, and their mutation severely compromised the interaction with ribosome. Our findings show the crucial role of electrostatic interaction in RIP-ribosome interaction and highlight the importance of interacting with specific ribosomal protein for RIPs to attain a good accessibility to ribosome.

The interacting site on maize RIP for P2 was found to be different from other RIPs. Several pronounced structural differences were found between MOD and the typical topology of RIPs [26]. The P2-binding site on TCS is located at the interface of strands β7 and β8 and helices αG and αJ at which TCS establishes extensive hydrophilic and hydrophobic bonding with C11-P [25]. However, the antiparallel β7 and β8 are replaced by a short helix αI in MOD (Figure 6A). It has also been found that StxA and C11-P interact through hydrophilic and hydrophobic interaction [16]. We

Figure 6. Residues on MOD that correspond to the C11-P interacting residues on TCS. A) Superimposed structures of MOD (pdb: 2PQI) and TCS-C11-P complex (pdb: 2JDL). The beta strands β7 and β8 in TCS (wheat) are replaced by the helix αI in MOD (cyan) while helices αG and αJ are conserved. P2-binding residues on MOD and TCS are distant apart as indicated in red and green respectively. B) Comparison of C11-P interacting residues on TCS and the corresponding residues on MOD. Structures of MOD and TCS are superimposed to locate the residues on MOD (colored in cyan and labelled in black and italic) corresponding to the C11-P interacting residues on TCS (colored in wheat and labelled in red). Many residues in these two RIPs are different, especially those at the hydrophobic patch of TCS (Phe-166, Val-223, Ile-225, Gly-231, Val-232 and Asn-236), suggesting MOD may interact with C11-P at a different site. C) Stereo image zooming in the hydrophobic pocket of MOD (colored in cyan) with C11-P (colored in yellow). The model reveals Arg-275 on MOD may crash with Leu-9 on C11-P while Glu-272 confronts directly Asp-11. Residues on TCS that interact with C11-P are highlighted in wheat for reference.

doi:10.1371/journal.pone.0049608.g006
superimposed the structures of MOD and TCS-C11-P complex and found that the hydrophobic region Gln-170, Lys-173, Arg-174 and Lys-177 in TCS for interacting with C11-P are substituted by aliphatic or polar Thr-217, Ala-220, Gly-221 and Ser-224 in MOD that appear less favorable for the electrostatic interaction with the DDD motif of C11-P (Figure 6B).

Alternatively, the hydrophobic and slightly polar patch on TCS is replaced by a charged and highly polar surface on MOD. Phc-166, Val-223, Ile-225, Gln-231/Val-232 and Asn-236 in TCS are replaced by Thr-213, Lys-264, Lys-268, Ghu-272 and Arg-275 in which the housing of hydrophobic FGLFD motif of C11-P is highly discouraged. Particularly, we noticed Arg-275 of MOD may crash with Leu-9 of C11-P while Ghu-272 is facing the Asp-11 of C11-P (Figure 6C).

While MOD, TCS and StxA commonly rely on the tail of P2 for interaction, we observed that MOD docks at the DDDMG motif of P2 without the involvement of the FGLFD motif. This explains why the interaction between MOD and P2 is weaker and more salt-dependent. Examination of the hydrophobic amino acids around Lys158-Lys161 of MOD showed that they form a shallow cavity which may not be appropriate for the housing of the hydrophobic motif of C11-P. Alternatively, the positively charged Lys158-Lys161 in MOD are replaced by hydrophobic and polar residues Tyr-141, Tyr-142, Asn-143 and Ala-144 in TCS and less polar residues Ser-144, His-145, Ser-146 and Gly-147 in StxA which further supports MOD binds P2 very differently from those of TCS and StxA.

We also noticed that the DDDMG motif on P2 may contribute a more substantial role in the interaction for some RIPs. In the case of StxA, there was still some interaction with P2 with the deletion of the last five residues (Figure 5B).

Amino acid sequences of RIPs show that their ribosome binding motifs are highly variable and the ability to interact with the ribosomal stalk proteins is believed to attain independently through convergent evolution [35]. P-protein binding residues in TCS, StxA and MOD are not mutually conserved as seen from their superimposed structures and also not consensus in their respective homologous RIPs. Our findings further illustrate the flexibility on docking of RIPs on ribosomal proteins for the targeting of the sarcin-ricin loop.

Previous studies demonstrated RIPs could not elicit their full activity when the ribosome-binding residues were mutated or the ribosome is deficient in their ribosomal interacting partner [14,18,21,22,26], pointing out the importance of interaction with ribosomal proteins to the activity of RIPs. In the present study, we observed that the ρ1-glycosidase activity and cytotoxicity of MOD were much lower than TCS, further ascertaining the role of ribosomal proteins for the enzymatic activity of RIPs and implying the strength of their interaction with P2 may play a role on the activity of RIPs.

In summary, we have shown that RIPs may use different regions to dock on the same ribosomal target depending on their structural features. In the case of maize RIP, it interacts with P2 at a novel binding site Lys158-Lys161 which only permits hydrophilic interaction with the DDDMG motif but not the hydrophobic tail of C11-P.

We have further illustrated that the strength of interaction between RIPs and ribosomal proteins is an important factor for the activity of RIPs. Lastly, our work provides an example on the use of protein-protein interaction but not the catalytic centre to modulate enzymatic activity and illustrates the flexibility on the docking of RIPs on ribosomal proteins for activity.

**Supporting Information**

Figure S1 Cellular uptakes of MOD and TCS are similar. Same amount of RIPs was added to cells. Protein taken up by cells was detected by antibody against His-tag. Beta-actin was used as the loading control to ensure similar amount of cells was subject to the western analysis. (TIF)

**Acknowledgments**

We thank Prof. R.S. Boston for providing the plasmid DNA encoding MOD and pro-RIP.

**Author Contributions**

Conceived and designed the experiments: YTW KBW PCS. Performed the experiments: YTW YMN ANSM. Analyzed the data: YTW KBW PCS. Contributed reagents/materials/analysis tools: KHS KBW PCS. Wrote the paper: YTW PCS.

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