The Structure of the Karrikin-Insensitive Protein (KAI2) in Arabidopsis thaliana

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**Abstract**

KARRIKIN INSENSITIVE 2 (KAI2) is an α/β hydrolase involved in seed germination and seedling development. It is essential for plant responses to karrikins, a class of butenolide compounds derived from burnt plant material that are structurally similar to strigolactone plant hormones. The mechanistic basis for the function of KAI2 in plant development remains unclear. We have determined the crystal structure of Arabidopsis thaliana KAI2 in space groups P2₁ 2₁ 2₁ (a = 63.57 Å, b = 66.26 Å, c = 78.25 Å) and P2₁ (a = 50.20 Å, b = 56.04 Å, c = 52.43 Å, β = 116.12°) to 1.55 and 2.11 Å respectively. The catalytic residues are positioned within a large hydrophobic pocket similar to that of DAD2, a protein required for strigolactone response in Petunia hybrida. KAI2 possesses a second solvent-accessible pocket, adjacent to the active site cavity, which offers the possibility of allosteric regulation. The structure of KAI2 is consistent with its designation as a serine hydrolase, as well as previous data implicating the protein in karrikin and strigolactone signalling.

**Introduction**

Bushfires present organisms with both hardships and opportunities. For plants, fires deliver a flux of nutrients to the soil and reduce competition for light and water. Karrikins are a class of compound present in bushfire smoke that stimulate the germination of dormant seeds [1,2]. Karrikins are characterised by a butenolide moiety that is also common to strigolactones, endogenous plant hormones that were originally identified as factors that promote germination of seeds of parasitic weeds of the Orabanchaceae such as *Striga* and *Orobanche* species [3,4]. Strigolactones stimulate hyphal branching of arbuscular mycorrhizal fungi [5–7], promoting a symbiotic interaction with more than 80% of vascular plants that enhances nutrient uptake. In addition, strigolactones regulate several aspects of plant shoot and root development and thus influence overall plant architecture [8–12].

Discovering the molecular mechanisms of karrikin and strigolactone action has recently taken a step forward with the isolation of plant mutants. A family of α/β hydrolases has emerged as central players in mediating the responses of plants to these butenolide compounds. The *dwarf14* mutant of rice is insensitive to strigolactones, as are orthologous mutants in Arabidopsis and petunia [13–15]. DAD2, the D14 orthologue in petunia, has slow hydrolytic activity towards the synthetic strigolactone GR24 (~3 molecules of GR24 hydrolysed per DAD2 molecule per hour [15]) and this activity has been proposed to be essential for strigolactone perception [15]. Furthermore, the direct hydrolysis of the butenolide ring has recently been proposed as the mode of action for strigolactones and extended to karrikins [16]. In Arabidopsis, the DWARF14 parologue KAI2 ([KARRIKIN INSENSITIVE 2]) is required for responses to karrikins, but does not appear to be essential for strigolactone responses, at least with respect to the regulation of shoot branching [14]. Nevertheless, genetic studies indicate that KAI2 can mediate activity of compounds besides karrikins, including GR24 and other butenolides with strigolactone-like functionality [14,17]. Structural specialisation between DWARF14/DAD2 and KAI2 proteins may explain their functional specialities towards different butenolides.

The precise role that KAI2 plays in the karrikin and strigolactone signalling pathways remains unclear. Here we report the crystal structure of KAI2, which may provide valuable insight into its involvement in these pathways, especially with regard to substrate or ligand specificity. In particular, a comparison with the recently elucidated structure of DAD2 and its postulated role as both a strigolactone receptor and hydrolase is discussed.

**Materials and Methods**

**Synthesis of KAR₂**

KAR₂ was prepared according to the method of Goddard-Borger et al. [18].

**Cloning and expression of KAI2**

The native Arabidopsis thaliana KAI2 coding sequence (At4g37470) was amplified by PCR using seedling-derived cDNA template and primers 5’-GGGGACAAAGTTTGTACAAAAAGCCAGCTCATGGGTTGAGTGAAGAACG-3’ and 5’-GGGGACCACTTTGTACAGAAAGCTGCGTTTCAACATAGCAT-3’.
were frozen in dry ice and stored at 
-18 hours at 16°C. Growth proceeded for a further 
cooled to 16°C in 150 mM NaCl. Screening for the buffer 
with optimal protein solubility was performed according to 
Jancarik et al [20], which is the same format as a hanging drop 
vapour diffusion crystallisation experiment. Briefly, drops containing 
5 µL of protein solution were mixed with 5 µL of buffer solution 
and equilibrated against 500 µL of buffer solution at 293 K. After 4 weeks, the drops containing 0.1 M Tris pH 8.5 
and 9.0 supplemented with 75 and 150 mM NaCl had remained 
clear. These buffers were used in subsequent purifications of 
KAI2.

KAI2 purification

Pellets from 400 mL of expression culture were resuspended
with 15 mL of 100 mM Tris 300 mM NaCl 20 mM imidazole
pH 9.0, supplemented with 15 units of Benzonase (Novagen,
Darmstadt). Cells were lysed with 4 passes through an Emulsiflex
C5 high-pressure homogeniser at 10,000–15,000 kPa (Avestin,
Ottawa). Lysates were clarified by centrifugation at 16,100 rcf for
1 h at 4°C. All chromatography steps were performed at room 
temperature. The clarified lysate was filtered through a 0.22 µm
filter and applied manually to a pre-equilibrated 5 mL HisTrap column (GE Healthcare). The loaded column was washed with 
four column volumes of lysis buffer at 2 mL/min. Protein was eluted 
using a ten column volume linear gradient from 0–100% 
buffer +250 mM imidazole). KAI2 eluted at approximately 
100 mM imidazole. KAI2-containing fractions were pooled,
concentrated to 3.5 mL and applied to a 120 mL Superdex 200 
column (GE Healthcare) for size exclusion chromatography in
100 mM Tris 150 mM NaCl pH 9.0. Pure KAI2 typically eluted 
as one peak, which was collected and concentrated using a 10 kDa 
molecular weight cut off centrifugal ultrafilter (Sartorius) to 
10.2 mg/mL, yielding approximately 16 mg of highly purified 
KAI2 protein. In an attempt to obtain protein bound to 
phenylmethyl sulfonyl fluoride (PMSF), KAI2 was lysed in 
20 mM Tris pH 7.0, 150 mM NaCl, 1 mM PMSF. The lysate 
was gently rocked at 4°C for 1 h before 5 mL of 200 mM
TRIS pH 9.0 was added and the purification was continued as
previously outlined. Purified protein was further incubated with 
1 mM PMSF at 4°C for 16 hours. All protein samples were 
divided into aliquots, flash-cooled in liquid nitrogen and stored at 
-80°C.

Protein crystallisation

Sparse matrix vapour diffusion crystallisation screens were 
performed using Hampton Index Screen (Hampton Research) 
with a Phenix liquid handling robot (Art Robbins) in sitting drop 
format in 96 well plates (Hampton Research, HR8-149). Drops 
between 0.4 and 0.6 µL were equilibrated against 90 µL of 
reservoir solution. A harvestable, diffracting protein crystal grew in 
a condition with 0.2 µL of protein solution (10.2 mg/mL KAI2) 
and 0.4 µL of reservoir solution (1.4 M sodium/potassium 
phosphate pH 8.4). Crystals were optimised by screening buffer 
and concentration in 24 well sitting drop format where 2.5 µL 
of protein solution (3.0 mg/mL) was added to 2.5 µL reservoir 
solution and equilibrated against 500 µL of reservoir solution. A
32 µm×6 µm×6 µm multiple crystal grew in 1.5 M sodium/
potassium phosphate pH 7.2 (KAI2b; Figure 1) and single crystals

Figure 1. KAI2 crystal. A multiple KAI2 crystal (32 µm×6 µm×6 µm) 
that was split for data collection (KAI2b).
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The Structure of KAI2

Optimisation of protein solubility

The instability of KAI2 at concentrations greater than 1 mg/
M in 20 mM sodium phosphate buffer, 150 mM NaCl pH 8.0, 
prompted optimum solubility screening. The buffer solutions 
screened against were citric acid pH 3.0–6.0, sodium acetate 
3 µm, sodium carbonate, bis-Tris pH 5.3, Bis-Tris pH 3.0–7.0, MOPS pH 6.5–7.4, HEPES pH 6.5–8.0 or Tris pH 6.5–9.0, at 0.1 M concen-
tration with either 75 or 150 mM NaCl. Screening for the buffer 
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clear. These buffers were used in subsequent purifications of 
KAI2.
grew in 1.4 M sodium/potassium phosphate pH 7.3. (KAI2c). KAI2 seed stock was prepared from a crystal in the same drop that yielded KAI2b. Attempts to co-crystallise KAI2 with the karrikin molecule KAR2 were unsuccessful, but yielded higher resolution diffracting crystals (KAI2a). Crystals that grew in the presence of KAR2 were crystallised from a drop containing 1.8 µL protein sample (6 mg/mL), 0.6 µL KAI2 seed stock, 0.4 µL 30% glycerol and 1.2 µL reservoir solution of 1.4 M sodium/potassium phosphate pH 7.3. Prior to crystallisation, KAR2 dissolved in 100% (v/v) DMSO was added to the protein solution to a final concentration of 1 mM KAR2 and 2.5% DMSO. The N-terminal leader sequence was not removed prior to crystallisation experiments.

**Figure 2. Crystal structure of KAI2.** A. Stereoscopic ribbon diagram of KAI2 coloured from amino (blue) to carboxy (red) terminus. Every twentieth Cα is shown as a labelled sphere. B. Stereoscopic ribbon diagram of all three models of KAI2 (KAI2a blue, KAI2b orange, KAI2c black) and the models of DAD2 (purple) and RsbQ (brown) superposed. C. Stereoscopic cartoon diagram of KAI2. The α/β hydrolase domain is shown in blue and the cap domain shown in red. doi:10.1371/journal.pone.0054758.g002
Data collection and processing

Where necessary, single crystals were split from the multiple crystals. Crystals mounted in a nylon loop were briefly immersed in mother liquor containing 20% glycerol then frozen in liquid nitrogen for data collection. Complete X-ray data (KAI2a –180° in 0.5° rotations, KAI2b and KAI2c –360° in 1.0° rotations) were collected at the Australian Synchrotron beamlines MX1 or MX2. Data were integrated with XDS [21] and scaled using SCALA [22] from the CCP4 software suite [23]. The structure was solved by molecular replacement with MOLREP [24] using the crystal structure of the monomeric Bacillus subtilis protein RsbQ (PDB code 1WOM) [25] as the search model. Model building was performed with COOT [26]. Initial rigid-body and restrained refinement was performed using REFMAC [27]. Final rounds of refinement were performed with BUSTER [28]. Root-mean-square deviation (RMSD) values were calculated with LSQMAN [29]. Cavity volumes were calculated using VOIDOO [30] on the highest resolution structure KAI2a, using a primary grid spacing of 0.2 Å. Molecular graphics were generated using PYMOL [31]. Structures were analysed using MOLPROBITY [32]. Atomic coordinates and structure factors have been deposited in the Protein Data Bank under accession codes 4HRY (KAI2a), 4HTA (KAI2b) and 4HRX (KAI2c).

| Table 1. Data collection and refinement statistics for Arabidopsis thaliana KAI2. |
|--------------------------------------|-----------------|-----------------|-----------------|
| Structure               | KAI2.a          | KAI2.b          | KAI2.c          |
| Space group             | P2,2,2         | P2,2,2         | P2             |
| Unit-cell parameters (Å, °) | a = 63.57, b = 66.26, c = 78.25, α = 90, β = 116.12 | a = 63.39, b = 66.06, c = 77.62, α = 90, β = 116.12 | a = 50.20, b = 56.04, c = 52.43, β = 116.12 |
| Temperature (K)         | 100             | 100             | 100             |
| X-ray source            | MX1, Australian Synchrotron | MX2, Australian Synchrotron | MX2, Australian Synchrotron |
| X-ray wavelength (Å)    | 0.95370         | 0.95390         | 0.95390         |
| Detector                | ADSC Quantum 210r CCD | ADSC Quantum 315r CCD | ADSC Quantum 315r CCD |
| Resolution (Å)          | 27.50–1.55 (1.64–1.55) | 77.62–1.90 (2.00–1.90) | 56.04–2.11 (2.23–2.11) |
| Rmerge (%)              | 9.3 (54.2)      | 11.5 (68.0)     | 16.2 (65.9)     |
| Rmax (%)                | 10.2 (59.1)     | 12.3 (70.6)     | 17.4 (71.1)     |
| Rb_conv (%)             | 4.0 (23.1)      | 3.2 (18.6)      | 6.3 (26.4)      |
| No. of unique reflections | 48612 (6723)   | 26147 (3575)    | 14659 (1896)    |
| Average multiplicity    | 6.5 (4.0)       | 14.3 (13.9)     | 7.4 (6.7)       |
| (|I|/|<I>|)               | 10.8 (3.0)      | 22.5 (4.9)      | 11.1 (2.8)      |
| Rwork                   | 18.10           | 15.52           | 16.09           |
| Rfree                   | 20.65           | 17.40           | 20.63           |
| Mean B value (Å²)       | 20.04           | 22.48           | 22.19           |
| R.m.s.d. from ideal geometry | -             | -               | -               |
| Bond lengths (Å)        | 0.010           | 0.010           | 0.010           |
| Bond angles (°)         | 1.010           | 0.96            | 1.070           |
| No. of protein residues | 268             | 271             | 266             |
| Water/solvent atoms     | 405             | 272             | 194             |
| Estimated coordinate error (Luzzati) (Å) | 0.164 | 0.159 | 0.205 |
| Poor rotamers%           | 0              | 0               | 0               |
| Ramachandran%            | -              | -               | -               |
| Favoured (%)             | 97.7%           | 98.5%           | 98.1%           |
| Allowed (%)              | 2.3%           | 1.5%            | 1.9%            |

Values in parentheses correspond to the highest resolution shell. * Values obtained using MOLPROBITY [32].

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Protein sequence analysis

Protein sequence alignments were performed using ALINE [33]. KAI2 protein sequences used in sequence alignments were from Arabidopsis thaliana (NCBI GI: 15255567), Ricinus communis (NCBI GI: 255567989), Populus trichocarpa (NCBI GI: 224071259), Solanum lycopersicum (NCBI GI: 225311281), Vitis vinifera (NCBI GI: 225458830), Brachypodium distachyon (PlantGDB: Brachypodium_distachyon-10841), Hordeum vulgare (NCBI GI: 326500818), Zea mays (NCBI GI: 226530032), Sorghum bicolor (NCBI GI: 242035387) and Oryza sativa cv. japonica (NCBI GI: 115453689). D14 protein sequences used in sequence alignments were from Petunia hybrida (DAD2) (NCBI GI: 404434487), Arabidopsis thaliana (NCBI GI: 18396732), Ricinus communis (NCBI GI: 255538072), Vitis vinifera (NCBI GI: 225458830), Populus trichocarpa (NCBI GI: 224129864), Solanum lycopersicum (Sol genomics network: Solyc04g077860.2.1), Hordeum vulgare (NCBI GI: 326496392), Oryza sativa cv. japonica (NCBI GI: 115453689), Sorghum bicolor (NCBI GI: 242041843) and Zea mays (NCBI GI: 226501208).

Results and Discussion

A. thaliana KAI2 was cloned into pDEST17 and expressed in E. coli resulting in protein samples of approximately 30 kDa.
molecular mass as determined by SDS-PAGE. An optimal buffer for solubility was identified as 0.1 M Tris pH 9.0, 150 mM NaCl. KAI2 was successfully crystallised from sodium/potassium phosphate in two crystal forms, one orthorhombic, \( P2_1 2_1 2_1 \) (\( a = 64 \text{ Å}, b = 66 \text{ Å}, c = 78 \text{ Å} \)) and the other monoclinic, \( P2_1 \) (\( a = 50 \text{ Å}, b = 56 \text{ Å}, c = 52 \text{ Å}, \beta = 116^\circ \)), each with one protein molecule in the asymmetric unit. Full details of data collected from three crystals (KAI2a, KAI2b and KAI2c) are presented in Table 1. The processed KAI2a data showed a high signal-to-noise ratio at high resolution indicating that the crystal diffracted beyond the collected limit of 1.55 Å. Unfortunately, the crystal had deteriorated before higher-resolution data could be collected, and other crystals did not diffract so well. KAI2a is presented here as the highest resolution structure (1.55 Å) (Figure 2A). Although KAI2b diffraction less well (1.90 Å), a portion of the N-terminal linker sequence (from Ser2 to Lys25) could be clearly observed in the electron density. Monoclinic KAI2c diffracted to 2.11 Å. No conspicuous conformational differences were observed between KAI2 structures (Figure 2B; maximum RMSD between KAI2a and KAI2c, 0.4 Å [262/269 Cα atoms]). The overall fold of KAI2 consists of an \( \alpha/\beta \) hydrolase domain and a four helix cap domain (Figure 2C). KAI2 has essentially identical topology to DAD2 [15] and the signalling protein RsbQ from \textit{Bacillus subtilis} [25] [all missing the first \( \beta \)-strand of the canonical \( \alpha/\beta \) hydrolase domain [34]], with RMSD values of 0.9 Å [262/269 Cα atoms] and 1.2 Å [254/269 Cα atoms] from KAI2a respectively.

The protein surface

Functional specialisation between KAI2 and D14 proteins might be mediated by each of the proteins’ respective interaction partners, so we scrutinised the conserved amino acid differences of surface exposed residues within each of the protein families. No conspicuous regions of conserved differences were detected which would indicate sites of interaction with different protein partners.

The active site cavity

The active site is present in a largely hydrophobic pocket of volume 336 Å³, with a classical serine hydrolase catalytic triad at its base (composed of Ser95-His246-Asp217). The KAI2 catalytic pocket is smaller than the catalytic pocket of DAD2 (448 Å³), but it is still sufficiently large to accommodate the synthetic strigolactone GR24. The two pockets are highly similar in terms of overall shape and amino acid composition (Figures 3A and 3B). No obvious active site cavity features account for a difference between KAI2 and DAD2 [15] and the signalling protein RsbQ from \textit{Bacillus subtilis} [25] [all missing the first \( \beta \)-strand of the canonical \( \alpha/\beta \) hydrolase domain [34]], with RMSD values of 0.9 Å [262/269 Cα atoms] and 1.2 Å [254/269 Cα atoms] from KAI2a respectively.
and D14 in terms of karrikin binding. Six of the seven cavity-lining phenylalanine residues are conserved between the two proteins (Figures 3A and 3B). The non-conserved residue within these is Tyr124, which replaces Phe125. The hydroxyl group of this side chain occludes a small pocket (41 Å³) proximal to the catalytic residues. In DAD2, this small pocket is connected to the main cavity, helping to explain the discrepancy in size between the KAI2 and DAD2 main cavity volumes. This specific Tyr/Phe substitution is conserved within KAI2 and D14 protein families (Figure 3C) suggesting that the two proteins may differ in their natural substrates/ligands.

Catalytic residues

The catalytic triad of KAI2 is observed with either a Tris buffer molecule (KAI2a) or glycerol molecule (KAI2b) nearby. As the structure of a PMSF-adduct of RsbQ had previously been reported [25], we attempted extensive incubation of KAI2 with PMSF, but were unable to detect any covalent modification at Ser95 either in crystals or by electrospray mass-spectrometry. This observation is not unprecedented for catalytically active serine hydrolases [35,36]. In an attempt to rationalise the inactivity of KAI2 towards PMSF, we scrutinised the conformation of the catalytic residues. The Ser-His-Asp triad residues are hydrogen bonded in a classical conformation for active hydrolysis (Figure 4A) [34]. However, the imidazole side-chain of His246 is in a different plane to that observed in DAD2 and RsbQ, coordinating to Oδ1 of Asp217 as opposed to Oδ2. Furthermore, a discrepancy in the coordination of Ce1 of His246 in KAI2 compared to that of active serine hydrolases, including DAD2 and RsbQ, was observed. Ce1 of the catalytic histidine in serine hydrolases is acidic and acts as a CH…O hydrogen bond donor to a carbonyl oxygen [37] in catalytically active serine hydrolases [37,38]. Consistent with this, Ce1 of His246 in DAD2 forms a 3.34 Å hydrogen bond to the amide O of Gly120 which deviates 9° from the Ce-H bond. In RsbQ, Ce1 of His247 forms a 3.04 Å hydrogen bond to the amide O of Gly120 which deviates 22° from the Ce1-H bond, while Ce1 of His246 in KAI2 is 3.65 Å from the amide O of Ser119 and deviates 53° from the Ce1-H bond (Figure 4B). Steric hindrance from the side chain of Ser119 in KAI2 (as opposed to Gly119 in DAD2 and Gly120 in RsbQ) alters the arrangement of the catalytic histidine, potentially explaining the inability of the protein to react with PMSF. This amino acid substitution between KAI2 and DAD2 at position 119 is conserved within KAI2 and D14 protein families (Figure 3C). The structure of DAD2 is described as having an inactive conformation at the active site (Oγ of Ser96 is oriented away from His246) [15], yet retains hydrolytic activity on GR24 [15]. Therefore some rearrangement must occur within the catalytic residues of DAD2 prior to catalysis. Similarly, the catalytic residues of KAI2 are observed in an inactive conformation and may require perturbation for activity. This may be facilitated by the natural substrate for the enzyme or some other regulatory mechanism, which cannot be mimicked by PMSF.
A secondary pocket

There is a second solvent-accessible pocket within KAI2, adjacent to the primary pocket containing the active site residues but separated from it internally by the side-chain of Phe26 (Figure 5A). The secondary pocket (137 Å³) is significantly smaller than the primary pocket but is large enough to accommodate a karrikin-sized molecule. The same pocket is present within the DAD2 structure (Figure 5B), although it is smaller (30 Å³) and blocked from the solvent by the side chain of Phe188. The equivalent residue in KAI2 is Ser188, which permits solvent accessibility of the pocket. This Ser/Phe substitution is conserved within KAI2 and D14 protein families (Figure 3B). The difference in size between the two pockets is accounted for by another conserved difference between the proteins at residue Gly53 of KAI2/Cys54 of DAD2, where in DAD2 the cysteine side chain protrudes in to the second pocket, reducing its volume. The solvent accessibility, volume, and close proximity of this secondary pocket to the active site pocket offer the possibility that KAI2 activity may be regulated by an allosteric co-factor. The conserved differences between KAI2 and D14 protein families suggest that this secondary pocket may play a role in the functional specialisation of these two proteins.

Conclusion

The X-ray crystal structure of KAI2 has provided us with insight into the possible molecular function of the enzyme. The conformation of active site residues specifies the design of the enzyme as a serine hydrolase, although the conditions under which it is active and its native substrate remain unknown. The KAI2 active site cavity is large enough to accommodate a strigolactone molecule and is highly similar in structure and amino acid composition to the strigolactone hydrolase DAD2. This observation is consistent with genetic studies that indicate that KAI2 and AdDWARF14 (the Arabidopsis DAD2 orthologue) can both mediate seedling responses to the synthetic strigolactone GR24 [14]. However, AdDWARF14 is unable to mediate karrikin signalling, and karrkins cannot function as inhibitors of shoot branching, unlike GR24 [39]. There are no conspicuous features within the KAI2 structure that explain the difference in karrakin perception between KAI2 and D14 proteins. An inspection of the conserved differences in surface amino acid composition between KAI2 and D14 yielded no conspicuous regions of divergence. The unexpected finding of a second solvent-accessible pocket, distinct from, yet in close proximity to the active site pocket suggests that the activity of KAI2 might be regulated by an allosteric cofactor. This second pocket is larger than the equivalent pocket in DAD2 which is not solvent accessible, suggesting that this second pocket may contribute to functional specialisation between KAI2 and D14 proteins. The elucidation of the KAI2 structure now opens up avenues for investigating this possibility.

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Author Contributions

Conceived and designed the experiments: RBD MTW SMS CSB. Performed the experiments: RBD MTW AS GRF SMS. Analyzed the data: RBD MTW AS GRF SMS. Contributed reagents/materials/analysis tools: RBD MTW AS. Wrote the paper: RBD MTW SMS CSB.

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