The Hsp90 molecular chaperones are ATP-dependent enzymes that maintain protein homeostasis and regulate many essential cellular processes. Higher eukaryotes have organelle-specific Hsp90 paralogs that are adapted to each subcellular environment. The mitochondrial Hsp90, TNF receptor–associated protein 1 (TRAP1), supports the folding and activity of electron transport components and is increasingly appreciated as a critical player in mitochondrial signaling. Calcium plays a well-known and important regulatory role in mitochondria where it can accumulate to much higher concentrations than in the cytoplasm. Surprisingly, we found here that calcium can replace magnesium, the essential enzymatic cofactor, to support TRAP1 ATPase activity. Anomalous X-ray diffraction experiments revealed a calcium-binding site within the TRAP1 nucleotide-binding site located near the ATP α-phosphate and completely distinct from the magnesium-binding site adjacent to the β- and γ-phosphates. In the presence of magnesium, ATP hydrolysis by TRAP1, as with other Hsp90s, was noncooperative, whereas calcium binding resulted in cooperative hydrolysis by the two protomers within the Hsp90 dimer. The structural data suggested a mechanism for this cooperative behavior. Because of the cooperativity, at high ATP concentrations, ATPase activity was higher with calcium, whereas the converse was observed at low ATP concentrations. Integrating these observations, we propose a model in which the divalent cation choice can control switching between noncooperative and cooperative TRAP1 ATPase mechanisms in response to varying ATP concentrations. This switching may facilitate coordination between cellular energetics, mitochondrial signaling, and protein homeostasis via alterations in the TRAP1 ATP-driven cycle and its consequent effects on different mitochondrial clients.

Enzymatic studies of ATPases generally include magnesium as the physiological co-factor required for both nucleotide binding as well as catalysis. Except in specific cases, other divalent ions are generally not considered as even calcium concentrations are kept quite low (nanomolar) in the resting cytoplasm. By contrast with the cytoplasm, subcellular organelles can have vastly different ionic environments. It is now well appreciated that mitochondria serve as a major cellular calcium reservoir in nonmuscle cells. Calcium-phosphate granules form inside the mitochondrial matrix facilitating calcium storage and buffering, and if completely dissolved, would reach molar concentrations (1). Many studies show that calcium levels within mitochondria are tightly linked to both mitochondrial and cellular signaling pathways, including cell death via the mitochondrial permeability transition pore (2). Unsurprisingly, several mitochondrial enzymes are regulated by calcium ions (3). In particular, mitochondrial dehydrogenases involved in ATP production are stimulated by calcium ions, suggesting a direct link to cellular energetics. Given the prevalence of calcium in the mitochondrial matrix, it is likely that calcium ions play important roles in regulating other mitochondrial proteins. In this study, we look at how a mitochondrial homolog of the molecular chaperone Hsp90 (TRAP1)2 may be regulated by calcium binding via a novel mechanism.

Hsp90 molecular chaperones facilitate the folding and activation of ~10% of cellular proteins (4). Increasingly, it is appreciated that Hsp90s can support function throughout a protein’s lifetime, not just during initial synthesis or in response to cellular stress. Although Hsp90s are well conserved from bacteria to mammals, distinct orthologs can be found within organelles: Gpr94 in the endoplasmic reticulum, TRAP1 in mitochondria, and Hsp90.5 in chloroplasts. Beyond their role in supporting individual “client” substrate proteins, cytosolic Hsp90s are critically important for maintaining protein homeostasis and modulating certain pro-apoptotic kinases.

All Hsp90s are homodimers with each protomer composed of 3 globular domains: an ATP-binding N-terminal domain (NTD), a middle domain, and C-terminal dimerizing domain (CTD). Differences serve to adapt the Hsp90s for their various cellular roles, interactions with co-chaperones (cytoplasm only) and localizations. Hsp90s go through complex conformational rearrangements (domain rotations and open-closed transitions) throughout their ATP hydrolysis cycle (5) and this
ATPase activity is absolutely essential for its function in eukaryotic cells (6, 7). Despite being structurally conserved, Hsp90 homologs differ in their ATPase kinetics (8–10), likely reflecting a distinct yet precisely tuned conformational equilibrium between the open and closed state for each species (11).

Although the in vivo roles of cytosolic Hsp90s have been the most studied because of their importance for disease and evolution (12), recently attention is being focused on the mitochondrial isoform, TRAP1. TRAP1 has been implicated in regulating mitochondrial energetics (13), the opening of the mitochondrial transition pore (14) and kinase signaling (15) suggesting linkages between mitochondrial protein homeostasis, energetics, and broader aspects of cell function. Also relevant, have been clear connections to neurodegeneration (16, 17) and cancer (18).

TRAP1 structural studies have revealed the presence of a remarkably asymmetric closed state in the presence of nonhydrolyzable ATP analogs (19) that leads to a two-step sequential ATP hydrolysis mechanism, with the first ATP leading to a flip in asymmetry, and the second leading to re-opening (Fig. 4) (20). As the primary client-binding site maps to the region of maximal asymmetry, this suggests a strong coupling between the asymmetry flip and client remodeling. Another unique feature of TRAP1 is an N-terminal extension that renders its open-closed equilibrium particularly sensitive to temperature variations, possibly connecting its folding and signaling role to mitochondrial thermogenesis (21).

Although ATP is required to stabilize the closed N-terminally dimerized state in all Hsp90s, uniquely for TRAP1 closure (but not hydrolysis) will occur in the absence of any divalent cation co-factor (20, 21). Although the mechanism and purpose of this phenomenon are unknown, there have been reports of free ATP (not complexed to Mg$^{2+}$) existing in the mitochondrial matrix (22).

To obtain a better understanding how divalent cations may regulate TRAP1, experimental systems with purified protein can bypass the complexity of cell-based models; providing useful mechanistic insights that may be important in the cellular context. To this end, we investigated how the ATPase activity of TRAP1 is regulated by two divalent cations: magnesium or calcium. We found that unlike most other Hsp90 homologs, the TRAP1 ATPase can efficiently utilize calcium alone to support hydrolysis, and that the relative preference for magnesium or calcium depends upon the concentration of free ATP. Moreover, calcium induced a profound change in the cooperativity of ATP hydrolysis, indicating a change in the underlying mechanism. Crystallographic analysis revealed a calcium-binding site within the NTD distinct from that used by magnesium. These observations lead to a model of how calcium binding within TRAP1 can influence the TRAP1 catalytic cycle in an ATP-level dependent manner.

**Results**

**TRAP1 has unique enzyme kinetics depending on magnesium or calcium**

Given the role of mitochondria as a cellular calcium reservoir, we decided to test whether calcium might affect ATP hydrolysis by human TRAP1. ATP hydrolysis by Hsp90 happens in the closed state when the NTDs are dimerized. This closed state can be stabilized by nonhydrolyzable ATP analogs, or in the case of TRAP1, simply by omitting magnesium. Under these conditions, the rate of ATP turnover is much slower than the usually rate-limiting dimer closure, resulting in a build-up of the closed state (20, 21). We have previously taken advantage of this phenomenon to decouple dimer closure from ATP hydrolysis, which can be initiated by addition of Mg$^{2+}$. The closed dimer reopens once ATP hydrolysis resumes, and this can be readily monitored by a FRET assay (Fig. 1A). Because the steady-state assay only reads out the slow step of dimer closure, we decided to test whether calcium alone can trigger ATP hydrolysis from the closed state. As previously observed, addition of magnesium causes loss of FRET signal due to hydrolysis-triggered dimer re-opening. The addition of calcium can also cause dimer opening (Fig. 1B), and with a ~10-fold faster kinetics than magnesium. This suggests that in the absence of magnesium, calcium can fully support ATP hydrolysis by TRAP1 once it is closed.

Next, we assessed whether calcium alone can support both closure and ATP hydrolysis. As previously observed (9), ATPase activity in the presence of magnesium is noncooperative and reaches a plateau with as little as ~50 μM ATP. By contrast, ATPase activity using calcium follows the Hill equation with n = 1.4 and an ATPase rate that is 1.6-fold faster than in magnesium at saturating ATP (Fig. 1C, see Table 1 for fitted parameters). While fitting the data, we noted that the Michaelis-Menten equation is a poor fit to the magnesium ATPase activity (Fig. S1). This behavior also appears to be conserved in zebrafish TRAP1 (zTRAP1) (Fig. S2). The “bump” on the curve between 0 and ~200 μM ATP can be explained by a model in which the ATPase activity of the singly bound ATP state is greater than when ATPs are bound by both protomers. We confirmed this by measuring the ATPase activity of a heterodimeric mixture of hTRAP1 in which one ATP-binding site was inactivated via a D158N mutation. In the presence of magnesium, the ATPase activity curve of the heterodimer no longer has the bump and follows standard single-site Michaelis-Menten kinetics (Fig. 1d, blue line). The apparent affinity to ATP (K$_{M,ATP}$ defined as the concentration at half-maximum of activity) measured in this heterodimer (Table 2) is close to that estimated by the WT experiment (Table 1) (58 versus 73 μM). As predicted, this half-occupied ATP heterodimeric mixture has a dramatically higher ATPase activity (3.8 min$^{-1}$) than the fully occupied WT dimer (0.37 min$^{-1}$) (Fig. 1D, Tables 1 and 2). This is consistent with previous heterodimer experiments where a related point mutant, which should also abolish ATP binding (D158A), appeared to stimulate ATPase activity (23). The mechanism underlying this nontrivial rate enhancement is currently unknown, and it highlights yet another unique aspect of TRAP1 regulation by nucleotide. In contrast to the WT enzyme, the heterodimer ATPase activity in the presence of calcium is significantly reduced (0.6-fold), and consequently calcium is no longer preferred. To further examine whether the differential effects of magnesium versus calcium are a unique feature of TRAP1, ATPase activities of several Hsp90 orthologs were also measured.
Beyond the general trend that ATPase activity is lower when calcium is used, none of the tested homologs exhibited the distinct ATP concentration-dependent activity as seen in TRAP1 (Fig. S3).

### Table 1

|                     | 37 °C | 30 °C |
|---------------------|-------|-------|
|                     | hTRAP1, MgCl₂ | hTRAP1, CaCl₂ | zTRAP1, MgCl₂ | zTRAP1, CaCl₂ |
| Vₘₐₓ                | 0.67 ± 0.01  | 0.82 ± 0.01  |             |               |
| Vₛₜₜₜ,ATP           | 0.68 ± 0.05  | 1.50 ± 0.24  |             |               |
| Vₚₜₚₜ,ATP           | 0.37 ± 0.02  | 0.56 ± 0.03  |             |               |
| Kₛₜₜₜ,ATP (µM)      | 56.1 ± 2.2   | 66.2 ± 2.5   |             |               |
| Kₛₚₚₚₚ,ATP (µM)    | 58.0 ± 7.8   | 126.3 ± 30.5 |             |               |
| Kₛₚₚₚₚₚ,ATP (µM)   | 38.0 ± 19.9  | 25.2 ± 12.7  |             |               |
| nₜₜₜ,ATP            | 1.4 ± 0.0    | 1.8 ± 0.1    |             |               |

### Table 2

|                     | MgCl₂  | CaCl₂  |
|---------------------|--------|--------|
| Vₘₐₓ                | 3.77 ± 0.06 | 2.31 ± 0.15 |
| Kₛₚₚₚₚ,ATP (µM)    | 72.5 ± 4.4  | 273.2 ± 39.2 |

Beyond the general trend that ATPase activity is lower when calcium is used, none of the tested homologs exhibited the distinct ATP concentration-dependent activity as seen in TRAP1 (Fig. S3).
Modulation of TRAP1 ATPase activity by calcium and magnesium

Table 3
Fitted parameters for divalent cation titrations in low versus high ATP concentrations for zebrafish and human TRAP1

|                  | Low ATP (50 μM) | High ATP (400 μM) |
|------------------|-----------------|-------------------|
|                  | zTRAP1, MgCl₂   | zTRAP1, CaCl₂     |
|                  | zTRAP1, MgCl₂   | zTRAP1, CaCl₂     |
|                  | zTRAP1, MgCl₂   | zTRAP1, CaCl₂     |
| Vₘₐₓ, nM         | 0.47 ± 0.09     | 0.39 ± 0.01       |
| Kₘₐₓ, μM         | 38.6 ± 15.0     | 66.1 ± 5.0        |
| nₜₘₐₓ           | 0.9 ± 1.1       | 1.2 ± 1.0         |

|                  | Low ATP (35 μM) | High ATP (350 μM) |
|------------------|-----------------|-------------------|
|                  | hTRAP1, MgCl₂   | hTRAP1, CaCl₂     |
|                  | hTRAP1, MgCl₂   | hTRAP1, CaCl₂     |
|                  | hTRAP1, MgCl₂   | hTRAP1, CaCl₂     |
| Vₘₐₓ, nM         | 0.24 ± 0.00     | 0.16 ± 0.00       |
| Kₘₐₓ, μM         | 32.9 ± 1.8      | 91.9 ± 3.7        |
| nₜₘₐₓ           | 1.0 ± 0.0       | 1.1 ± 0.0         |

**ATP-site occupancy biases preference for magnesium versus calcium**

To better understand the inter-relationship between ATPase activity and divalent cation concentrations, we examined the effects of cation concentration under high and low ATP concentrations (hTRAP1 in Fig. 1, e and f; zTRAP1 in Fig. 5a). The apparent affinity (Kₘₐₓ) and occupancy (nₜₘₐₓ) of the divalent cations can be approximated by fitting the Hill equation to initial rates from titrations at fixed ATP concentrations. At low ATP concentrations, hTRAP1 has a 2.8-fold tighter apparent affinity for magnesium than calcium (33 μM Mg²⁺ versus 92 μM Ca²⁺), whereas zTRAP1 has a 1.7-fold higher affinity (39 μM Mg²⁺ versus 66 μM Ca²⁺), and also ~1.5-fold higher activity in both species (Table 3, low ATP). At low ATP concentrations, the ion-binding Hill coefficient is near 1.0 for both magnesium and calcium, suggesting that the cations preferentially bind to the protomer having one ATP bound. By contrast at high ATP concentrations, calcium binds ~4.5-fold more tightly than magnesium in hTRAP1 (59 μM Mg²⁺ versus 13 μM Ca²⁺) and ~7.5-fold in zTRAP1 (260 μM Mg²⁺ versus 37 μM Ca²⁺; see also Table 3, high ATP). The observed differences in apparent affinities between species most likely reflect intrinsic properties of each protein. At high ATP concentrations, n remains close to 1.0 for magnesium while indicating very significant cooperativity (n = 1.7) for calcium.

**Calcium binds to the NTD of TRAP1**

Crystal structures of the closed state of TRAP1 clearly reveal a single magnesium-binding site on each protomer, coordinating the β- and γ-phosphates of ATP as is commonly observed in many ATPases. The unitary Hill coefficient for both ATP and magnesium likely reflects the obligatory sequential (noncooperative) nature of ATP hydrolysis previously observed (20). Identifying where calcium binds would greatly facilitate understanding how calcium imparts an apparent cooperativity in TRAP1 ATPase activity, which has never been observed in other Hsp90 homologs.

As TRAP1 does not contain any apparent calcium-binding motifs, one possibility is that calcium binds at the same site used by magnesium, for example, as found in several other ATPases (Hsp70/DnaK/actin family) (24, 25). To determine where calcium binds in the closed state and any consequences on the structure, we replaced the magnesium in our standard zTRAP1 crystallization conditions (19) with calcium. The structure was solved to a resolution of 2.6-Å using molecular replacement (Protein Data Bank code 6D14), and reveals a virtually identical asymmetric closed state (19) (Fig. 2a). Despite the similarity in overall structure, no density coordinating the β- and γ-phosphates, where magnesium would be, was observed (Fig. S5, a and b). To unambiguously determine where calcium binds, we exploited the anomalous X-ray scattering signal from calcium and collected two datasets: one at 7450 eV (1.664 Å) and 11111 eV (1.116 Å). Although the anomalous difference at these energies is considerably weaker than what would be required for experimental phasing, significant peaks can be reliably observed for a well-determined structure using ANODE (26).

Using phases from the solved structure without the contribution of calcium atoms, the resulting anomalous difference density map reveals one strong peak within the nucleotide pocket of each protomer, located just above the α-phosphate (Fig. 2A, inset). The density is tightly coordinated by six oxygen atoms in an octahedral arrangement near the “lid” of the NTD (4 backbone carbonyl oxygens, a tyrosine hydroxyl, and the α-phosphate) (Fig. 2b). As a control, anomalous data were collected from crystals lacking both calcium and magnesium prepared for another study (20). In these crystals excess EDTA was used and the dimer was closed with ATP instead of AMPNP. No significant anomalous peaks were detected near the α-phosphate (Fig. S5c), again providing strong support that the peak observed in Fig. 2a is indeed from a calcium atom.

**Discussion**

Enzymatic studies of Hsp90 have almost exclusively used magnesium as a co-factor. Following up on an unexpected observation that calcium alone was sufficient to support TRAP1 ATPase activity, here we show that TRAP1 has a higher ATPase activity in the presence of calcium than in magnesium. This contrasts sharply with other Hsp90 orthologs, which are significantly less active with calcium. Strikingly, there are also significant differences in ATPase behavior as a function of ATP and cation concentration, with calcium inducing a cooperative dependence on ATP. Furthermore, at high ATP levels, activity cooperatively depends on calcium concentration, whereas magnesium acts noncooperatively. Consequently, calcium can differentially regulate TRAP1 activity in an ATP-concentration dependent manner. Together this reveals that these ions must act by quite distinct underlying mechanisms.

Rather than being a magnesium mimic positioned to stabilize negative charge build-up on the β- and γ-phosphates at the catalytic transition state, direct visualization of the calcium-binding site by crystallography shows that calcium occupies a site near the α-phosphate (Fig. 2B). In the closed state, the calcium is octahedrally coordinated by backbone carbonyl oxygens from the ATP lid, Tyr-221 hydroxyl, and the ATP α-phosphate oxygen. Unlike the magnesium site, this site is configured quite differently in the apo state versus the closed state, thus providing a unique coupling between open-closed conformation and calcium binding (Fig. 3, Movie S1). TRAP1’s ability to close does not require divalent cations, because ATP alone is sufficient to stabilize the closed state and reach an identical asymmetric conformation (20, 21). Moreover, because dimer closure is rate-limiting in the ATPase cycle, any decrease in
catalytic efficiency caused by the absence of a magnesium-stabilized transition state is not expected to dramatically alter the overall reaction rate. Whether there is an alternative hydrolysis pathway where solvent can stabilize the charge build-up on the $\beta$- and $\gamma$-phosphates while the calcium activates a different attacking water is not clear. Structural data supports that the calcium-binding cavity near the $\alpha$-phosphate presented here is conserved in the GHKL (Gyrase, Hsp90, histidine Kinase and MutL) ATPase superfamily. Hearnshaw et al. (27) observed a potassium ion occupying the same pocket with a similar octahedral coordination in DNA gyrase, and they pointed out that in yeast topoisomerase II, a lysine $\epsilon$-ammonium group occupies the same pocket. For Hsp90s, perhaps this pocket is responsible for the mild ATPase stimulation by monovalent cations previously observed for yeast Hsp90 (28).

The remarkable change in cooperativity observed with calcium indicates that TRAP1 can adopt a switch-like response to calcium at high ATP levels, providing a strong connection between calcium concentration in the mitochondrial matrix and TRAP1 ATPase-driven cycling. Although the physiological consequences have yet to be determined, there are at least several scenarios where this could modulate cell function: (i) by participating in the calcium-dependent switch between glycolysis and oxidative phosphorylation; (ii) by modulating TRAP1’s role in opening the mitochondrial transition pore via altered sequestration of cyclophilin D binding, (iii) by switching TRAP1 functionality or substrate specificity from client remodelling to more of a holdase behavior. Indeed, all of these could be a consequence of short-circuiting the normal, two-step ATP hydrolysis mechanism (20), going directly from closure to full reopening, and skipping the flip in asymmetry believed to be coupled to client remodeling (Fig. 4). TRAP1’s ability to suppress client aggregation by a capture and release mechanism (holdase activity) would likely be unaffected. Related to this, it...
Modulation of TRAP1 ATPase activity by calcium and magnesium

![Diagram of TRAP1 ATPase cycle showing the normal sequential two-step ATP hydrolysis mechanism observed with magnesium (solid blue arrows and text) and an alternative path (dashed green arrow and text) when using calcium.](image)

would be interesting to see whether TRAP1 has a distinct subset of client proteins under high versus low concentrations of ATP.

In principle, because calcium ions bind at an orthogonal site from magnesium, simultaneous binding of both divalent cations is possible. Preliminary experiments showed only a mild synergism when calcium is added to magnesium-ATPase reaction or vice versa. The ATP activities converge to roughly the average between calcium- and magnesium-ATPase rates alone as concentrations of Mg:Ca approach 1:1 stoichiometry (Fig. S6). Although the physiological context for this calcium and magnesium activity modulation is not yet known, the findings in this work provide a molecular model to inform future work in elucidating the role of TRAP1 in the mitochondria.

Materials and methods

Protein purification

Full-length WT hTRAP1 and zTRAP1 (without mitochondrial signal sequence) was cloned into a pET151DTOPO vector as described previously (19). Point mutations were introduced via site-directed mutagenesis by PCR. For FRET assays, single cysteines (G151C and K428C) were introduced into a cysteine-free zTRAP1 construct (21) to enable site-specific labeling with maleimide derivatives of Alexa Fluor 555/647 (Life Technologies). Proteins were expressed in BL21(DE3)-RIL Escherichia coli cells grown in TB media by addition of 0.5 mM isopropyl-β-D-galactopyranoside at A_{600} ~0.8, then incubated with shaking for 12–18 h overnight at 16 °C. All of the TRAP1 constructs have an N-terminal His6 tag with a cleavable tobacco etch virus site. Purification follows a standard protocol for nickel-nitrilotriacetic acid chromatography using 40 mM Tris buffer at pH 8.0, followed by an anion exchange with a MonoQ column, and a final size-exclusion step with a Superdex S200 16/60 (GE Healthcare) column.

FRET assays

FRET measurements were taken on a FluoroMax-4 (HORIBA Jobin Yvon) using excitation wavelength 532 nm and emission wavelengths 567/668 nm for Alexa Fluor 555/647, respectively. Temperature was maintained at 20 °C by a circulating water bath connected to the sample chamber. Kinetics were recorded every 4 s with an integration time of 0.3 s. To initiate dimer closure, 10 mM ATP was added to the reaction. Hydrolysis was then triggered by addition of 10 mM MgCl2 or CaCl2.

ATPase assays

Most of the ATPase assays performed in this study uses the phosphate release assay as previously described (20) using a coumarin-labeled phosphate-binding protein (29). In all assays, 0.5 μM TRAP1 dimer was used per reaction. Heterodimeric hTRAP1 was formed by equilibrating a mixture of the WT protein with 15-fold molar excess of the D158N point mutant (the D158N homodimer is inactive). Fluorescence was measured using a SpectraMax M5 plate reader by exciting at 385 nm and measuring emission at 475 nm with a cutoff filter at 455 nm at “low” sensitivity setting. The ATPase assays with zebrafish TRAP1 also monitors phosphate release via a chromogenic substrate 7-methyl-6-thioguanosine (7-MESG) in the presence of an E. coli PNase (purine nucleoside phosphorylase). In these assays, 0.14 mM 7-MESG and 2 mM PNase was used per reaction. The net absorbance of 7-MESG was measured by subtracting absorbance at 500 nm from 354 nm. Both the phosphate-binding protein and PNase used in this study was expressed and purified as previously described (20). Initial rates from phosphate release assays were from obtained from fitted slopes of a linear function, y = mx + b, to the most linear region of each kinetic trace. The activity versus concentration curves were analyzed by a standard Michaelis-Menten equation with the Hill coefficient, n, in the exponent,

$$V_0 = V_{max} \frac{[ATP]^n}{K_{ATP}^n + [ATP]^n} + \text{baseline}$$  \hspace{1cm} (Eq. 1)

where [ATP] is the ATP concentration and $K_{0.5}$ is the ATP concentration at which activity is half-maximum, $V_{max}$. The two-population ATPase activity model of WT TRAP1 in magnesium used the following equation,

$$V_0 = \left( V_{single,ATP} \times \frac{2 \times [ATP]}{K_{single,ATP}} + \frac{[ATP]^2}{K_{single,ATP} \times K_{double,ATP}} \right) / d$$

$$d = 1 + \frac{2 \times [ATP]}{K_{single,ATP}} + \frac{[ATP]^2}{K_{single,ATP} \times K_{double,ATP}}$$  \hspace{1cm} (Eq. 2)

where $K_{single,ATP}$ and $K_{double,ATP}$ corresponds to apparent affinity constants at which half-maximum activity is achieved for half (1-ATP bound) or fully occupied (2-ATP bound) popula-

References

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Table 4
Data collection and refinement statistics for zebrafish TRAP1 bound to AMPPNP (Protein Data Bank code 6D14) and calcium

| Zebrafish TRAP1 bound to AMPPNP and calcium |
|-------------------------------------------|
| Wavelength | 1.116 |
| Resolution range | 48.4–2.5 (2.591–2.5) |
| Space group | C 1 2 1 |
| Unit cell | 176.252 95.751 124.546 90 134.663 90 |
| Unique reflections | 49,471 (4,870) |
| Completeness (%) | 97.6 |
| Mean I/σ(I) | 10.36 |
| Wilson B-factor | 61.0 |
| Rfree | 11.3% |
| CC1/2 | 99.6% |
| Reflections used in refinement | 47,254 (4,868) |
| Reflections used for Rfree | 2,488 (251) |
| Rwork | 0.230 (0.4170) |
| Rfree | 0.253 (0.4471) |
| Number of non-hydrogen atoms | 9,794 |
| Macromolecules | 9,553 |
| Ligands | 64 |
| Solvent | 177 |
| Protein residues | 1,185 |
| Root mean square (bonds) | 0.016 |
| Root mean square (angles) | 1.31 |
| Ramachandran favored (%) | 93.67 |
| Ramachandran allowed (%) | 5.12 |
| Ramachandran outliers (%) | 1.21 |
| Rotamer outliers (%) | 9.65 |
| Clashscore | 11.10 |
| Average B-factor | 73.67 |
| Macromolecules | 74.35 |
| Ligands | 42.12 |
| Solvent | 48.43 |
| Number of TLS groups | 7 |

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