Critical Reviews and Perspectives

The structural and functional workings of KEOPS

Jonah Beenstock1,* and Frank Sicheri1,2,3,*

1The Lunenfeld-Tanenbaum Research Institute, Mount Sinai Hospital, Toronto, Ontario, M5G 1X5, Canada, 2Department of Molecular Genetics, University of Toronto, Ontario, M5S 1A8, Canada and 3Department of Biochemistry, University of Toronto, Ontario, M5S 1A8, Canada

Received July 30, 2021; Revised September 09, 2021; Editorial Decision September 12, 2021; Accepted October 04, 2021

ABSTRACT

KEOPS (Kinase, Endopeptidase and Other Proteins of Small size) is a five-subunit protein complex that is highly conserved in eukaryotes and archaea and is essential for the fitness of cells and for animal development. In humans, mutations in KEOPS genes underlie Galloway–Mowat syndrome, which manifests in severe microcephaly and renal dysfunction that lead to childhood death. The Kae1 subunit of KEOPS catalyzes the universal and essential tRNA modification N6-threonylcarbamoyl adenosine (t6A), while the auxiliary subunits Cgi121, the kinase/ATPase Bud32, Pcc1 and Gon7 play a supporting role. Kae1 orthologs are also present in bacteria and mitochondria but function in distinct complexes with proteins that are not related in structure or function to the auxiliary subunits of KEOPS. Over the past 15 years since its discovery, extensive study in the KEOPS field has provided many answers towards understanding the roles that KEOPS plays in cells and in human disease and how KEOPS carries out these functions. In this review, we provide an overview into recent advances in the study of KEOPS and illuminate exciting future directions.

INTRODUCTION

In 2006, a previously unknown five-subunit protein complex was discovered in two independent studies. This complex contained the putative protein kinase Bud32 and the putative endopeptidase Kae1 as well as the three uncharacterized proteins Gon7, Pcc1 and Cgi121. The presence of Kae1 and Bud32 led to the naming of this complex as KEOPS (Kinase, Endopeptidase and Other Proteins of Small size) (1) or EKC (Endopeptidase-like and Kinase Chromatin-associated) (2). For simplicity, we will use KEOPS naming throughout this review. Three key findings illuminated by these landmark studies include:

(i) All KEOPS subunits are conserved across archaeal and eukaryotic organisms apart from Gon7 which appeared specific to eukaryotes.

(ii) The Kae1 subunit is even more well conserved representing one of only ∼60 genes that are universally found in the genomes of all living cells (3).

(iii) The presence of all subunits in KEOPS are required for its ability to support the well-being of cells as measured by growth.

Over the past 15 years, a flurry of studies resolved many mysteries of how KEOPS functions, including its core biochemical role, together with the Sua5 protein, in catalyzing the universal N6-threonylcarbamoyl adenosine (t6A) modification of transfer RNAs (tRNAs), its role in human disease and its structure. However, many critical questions remain unanswered. In this review, we will provide a comprehensive overview into how the KEOPS complex was discovered, the role of KEOPS in cells and in human disease, the atomic structure of KEOPS and how this informs on function and lastly, the pressing frontier questions in the KEOPS field that remain to be addressed.

HOW KEOPS WAS DISCOVERED

The pre-KEOPS period

Bud32 was first identified in Saccharomyces cerevisiae as a gene required for normal cell growth and polarized cell divisions, commonly known as ‘budding’ (hence its name) (4). Sequence analysis predicted that Bud32 is a member of the eukaryotic protein kinase superfamily (5). Protein kinases adopt a bi-lobal architecture with a smaller N-terminal lobe (denoted N-lobe) and a larger C-terminal lobe (denoted C-lobe).
C-lobe). An inter-lobe cleft harbors elements projecting from the two lobes that are required for ATP binding and for the catalytic transfer of the gamma phosphate of ATP to a hydroxyl bearing sidechain in proteinaceous substrates (for a review on the structure of kinases, see (6)). Although the N-lobe of Bud32 harbors all the canonical elements characteristic of a functional protein kinase enzyme, the C-lobe appears abnormally minimalist (see below). Despite the unusually abbreviated C-lobe, Bud32 demonstrated autophosphorylation activity in vitro (5,7–9), a property of most protein kinases (10). As expression of catalytically dead Bud32 mutants did not fully rescue the slow growth phenotype of the bud32Δ yeast strain (11), Bud32’s phospho-transfer function was inferred to be critical for its cellular function. Within the human kinome, Bud32 bears closest resemblance to the Rio kinases (12). Since the Rio kinases and Bud32 are the only two protein kinases conserved across eukaryotes and archaea (12,13), the two may represent primordial members of the eukaryotic protein kinase superfamily from which the numerous other family members in eukaryotes have evolved.

Subsequent studies identified a Bud32 ortholog in humans named Tp53 reactive kinase or p53-related protein kinase (TPRK or PRPK respectively) based on a report assigning p53 as a PRPK substrate (14), which was most likely an in vitro artifact. A yeast two-hybrid analysis using PRPK as bait identified TPRK-binding protein (TPRKB), the human ortholog of yeast Cgi121, as a binding partner (15). Cgi121 was a protein of unknown function described in yeast as a binding partner of Kae1, Bud32 and Cgi121. In some archaeal species such as Methanothermbacter janaschii, Kael and Bud32 are fused into a single polypeptide, hinting that their functions are intertwined.

As noted above, Kael-family enzymes are part of a small group of ~60 genes present in the genomes of all autonomous cells (3). Family members in archaea and eukaryotes are called Kael while the bacterial orthologs are called YgiD/TsAD (18). Similar to Kael, TsAD is essential for bacterial growth (19). Eukaryote genomes also encode the Kael parologue Qri7 (18). The open reading frame (ORF) of Qri7 contains an N-terminal mitochondrial targeting signal peptide. Accordingly, although Qri7 is encoded in the nucleus, it resides exclusively in the mitochondria (20–22). Analogous to the essential role of Kael and TsAD in cellular fitness, Qri7 is essential for the function of mitochondria (20,21).

Very little was known about Pcc1 and Gon7. Deletion of the GON7 gene (as well as BUD32) perturbed the transfer of mannosyl-phosphate groups to cell surface oligosaccharides in S. cerevisiae (23). Pcc1 was completely uncharacterized, in part because of a lack of annotation resulting from its coding by an ORF containing an intron with a non-canonical splice site in S. cerevisiae (24).

Converging roads to the discovery of the KEOPS complex

Two unrelated genetic screens conducted in S. cerevisiae using different strategies led to the discovery of the KEOPS complex. The first study conducted a genome-wide screen for suppressors of the growth defect of cells lacking a functional form of Cdc13, an essential protein that protects telomere ends from erosion (1). The analysis revealed that deletion of the CGI121 gene enabled growth in the absence of Cdc13. Tandem affinity purification experiments of Cgi121 from yeast cells coupled to mass spectrometry revealed that Cgi121 resides in a protein complex with Bud32, Kael and Gon7. The physical interaction between KEOPS proteins was further supported by a large-scale analysis of the yeast interactome (25). Supporting the notion that the newly identified subunits of KEOPS form a functional complex, knockout of any of these genes led to highly similar slow growth and shortened telomeres phenotypes (1), which are most likely an indirect result of the role KEOPS plays in translation fidelity (elaborated on later in this review).

The second study (2) utilized a different starting point, searching for suppressors of splicing defects caused by a U to A mutation at the fifth position of U1snRNP, which yields a cold-sensitive phenotype (26). The analysis revealed that the cold-sensitive phenotype could be overcome by expression of an intron-less version product of the S. cerevisiae YKR095-A gene. This gene encoded a small uncharacterized protein that was required for the transcriptional activation of pheromone and galactose responsive genes and for polarized cell divisions. Chromatin-immunoprecipitation experiments showed the YKR095-A protein associated with chromatin, leading to its naming as polarized growth chromatin-associated controller 1 (Pcc1) (2). Tandem affinity purification coupled to mass spectrometry revealed that Pcc1 resides within a complex with Gon7, Kael, Bud32 and Cgi121.

THE ROLE OF KEOPS IN CELLS AND IN HUMAN DISEASE

KEOPS catalyzes the t6A modification of tRNA

The requirement for KEOPS for cell growth suggested that it fulfills an essential biochemical function. However, the precise role of KEOPS remained elusive until 2011. In this section, we provide an overview on how KEOPS activity is essential for the fidelity of translation through the role KEOPS plays in generating the universal tRNA modification Nε-threonylcarbamoyl adenosine (t6A).

tRNAs undergo a complex and multi-staged biogenesis processes that includes the instalment of multiple posttranscriptional modifications, catalyzed by tRNA-modifying enzymes (27). To date, over 100 tRNA posttranscriptional modifications have been identified (28). Each tRNA is estimated to carry 5 to 15 modifications distributed throughout its structure. These modifications regulate a range of tRNA functions from degradation to roles in translation and modification dysregulation is implicated in human disease (29–32). In particular, the decoding activities of tRNAs are dependent on base modifications at positions 34 and 37 within the tRNA anticodon loop (29), which critically influence the rate and fidelity of translation and the integrity
of the proteome (33). Of all tRNA modifications characterized, approximately 17% are conserved in all domains of life (34). Not unexpectedly, conserved modifications are generated by similarly conserved enzymes.

t6A is a universal tRNA modification discovered in 1969 (35) that is specifically found in tRNAs that decode ANN codons, where N denotes any nucleotide (Figure 1A) (28). t6A occurs exclusively on A37, one position 3' to the anticodon triplet (encompassing positions 34 to 36) and remodels and stabilizes the structure of the tRNA anticodon loop by preventing an inter-loop hydrogen bond between A37 and U33 (Figure 1B) (36–40). This remodeling facilitates enhanced binding to the ribosome (41). In the ribosome, t6A reinforces A1:U36 codon-anticodon binding by inducing base stacking interactions between A37 and A38 of the tRNA and the A at the first position of the codon triplet in the mRNA (Figure 1B) (40,42,43). As a result, t6A is essential for the fidelity of translation and for the functionality of ANN-decoding tRNAs (44–47). t6A can be further modified to five different derivative forms (28). In Escherichia coli and possibly in other species, almost all t6A is cyclized (denoted ct6A) by the CsdL/TsdA enzyme (48). Crystal structures show that ct6A adopts a hydantoin structural form. Like t6A, ct6A stabilizes the structure of the anticodon loop. However, it is not clear if ct6A helps codon-anticodon binding in the ribosome (49). t6A can also be N8-methylated by TRMO (tRNA methyltransferase O) to form m8t6A (50) or N2-methylthiolated by MtaB/CdkIα1l to form m2s2t6A (51). These two specific t6A derivatives modifications provide further tuning to the decoding activities of t6A modified tRNAs.

In 2009, 40 years after the discovery of t6A, the first enzyme in the t6A biosynthesis pathway was discovered through a comparative genomics approach that surveyed universally conserved proteins of unknown function (47) (for a detailed review of the conservation and diversity of t6A biosynthesis enzymes, see (18)). This analysis revealed that Sua5 (Suppressor of upstream AUG 5) (52) is directly involved in t6A biosynthesis. However, since Sua5 in isolation was not sufficient to modify tRNA in vitro (47), the t6A biosynthesis pathway was predicted to involve additional gene products. Interestingly, some bacterial and archaean genomes encode proteins that contain a Sua5-like domain fused to a Kael-like domain (53) hinting at a linked functional role between these distinct classes of enzymes. Rhizobium fredii NoLo, a Kael-Sua5 like fusion protein, functions as a carbamoyl-transferase (54), which raised the possibility that Sua5 and Kae1 could participate in catalyzing reactions with similar chemistry (53), which ultimately proved to be the case.

In 2011, three breakthrough studies identified the Kae1/TsaD/Qri7 family of enzymes as the missing component in t6A biosynthesis (46,55,56). All three studies noted the similarity between Kae1 to the Kael-Sua5 like fusion proteins as an indication that both enzymes might cooperate to generate t6A. The first study (56) noted that the extraordinary conservation of the Kael protein indicated that it would be involved in a universal cellular process. This led to a search for a potential role Kae1 might play in the synthesis of the major macromolecules in cells. While global DNA and RNA levels were only mildly affected in S. cerevisiae cells in the absence of Kael activity, protein synthesis was dramatically reduced, implicating Kae1 in translation. This finding led to the hypothesis that Kae1 works with Sua5 to modify tRNAs. The second study (46) used a comparative genomics approach, reasoning that since t6A is universally conserved the candidate enzymes involved in its biosynthesis would have a universal phylogenetic distribution and an unknown biochemical function. Remarkably, one of the few enzymes that met both criteria were the Kael/TsaD/Qri7 family of enzymes (57) hinting that Kae1 would collaborate with Sua5 in the t6A biosynthesis pathway. Both the first and second studies proved their working hypotheses by showing that mutations in Kael in S. cerevisiae resulted in a loss of t6A. The third study (55) revealed that the knockout of KEOPS genes leads to transcriptional activation of target genes of the transcription factor Gen4 in a Gen2-independent manner. Gen4 protein expression is normally suppressed by upstream ORFs that contain AUG start codons. The Gen4 protein is synthesized when the upstream kinase Gcn2 phosphorylates eIF2α, suppressing the inhibitory effect of the upstream ORFs (for an extensive review of Gen4 translational regulation see (58)). Therefore, increased Gen4 expression in the absence of KEOPS activity hinted that KEOPS would influence the selection of AUG start sites. Because Sua5 (45) and its downstream product t6A (47) are important for translation initiation, this led the authors to hypothesize that KEOPS is involved in generating the t6A modification.

Together, these breakthrough studies unequivocally proved that Sua5 and Kael/TsaD/Qri7 are needed for the t6A modification of tRNA in vivo (46,47,55,56). Subsequent studies with purified proteins (together with their respective binding partners) demonstrated the sufficiency of these enzymes to reconstitute the t6A modification of tRNA in vitro (59–61). The specific role each enzyme plays in this process and how these roles were assigned is described later in this review.

**KEOPS activity is essential for diverse aspects of cell biology and mutations in KEOPS underlie Galloway–Mowat syndrome**

Genetic studies conducted in the eukaryotic species yeast, zebra fish, fruit flies and mice as well as the archaean Haloferax volcanii have established an essential role for KEOPS in normal cell growth and in animal development. KEOPS function has been most extensively characterized in vivo in budding yeast. Knockout of any individual KEOPS gene leads to pleiotropic phenotypes, ranging from slow growth to shortened telomeres and to defects in transcription (1,2,44,55,62,63). These phenotypes are most likely an indirect result of the role KEOPS plays in tRNA modification and translation and of the far-reaching downstream effects that the perturbation of translation has on virtually all cellular processes. The impact of KEOPS activity on translation has been studied using reporter based assays, which showed that the absence of the t6A modification leads to an increase in translation mis-initiation and in ± 1 frame-shift events (46,47,55) as well as defects in stop codon
recognition (45, 64). A higher resolution study using ribosome profiling revealed that the role \( t^6A \) has in translation varies considerably from one ANN codon to another. The translation rates of ANN codons that are decoded by rare tRNAs or tRNAs with weak G34:U3 pairing at the wobble position decreased in the absence of \( t^6A \). In contrast, the translation rates of ANN codons that are dependent on high abundance tRNAs or tRNA with strong I34:C3 pairing at the wobble position were in fact increased by the absence of \( t^6A \) (44). Both types of changes in translation rate likely have a major role in the perturbed translation observed in the absence of \( t^6A \).

The importance of KEOPS activity for the well-being of cells has been confirmed in a variety of other model organisms. Briefly, in the archaeon \( H.\) volcanaii, the \( cgl21 \) and \( kael-bud32 \) fusion genes are essential for cell growth, while knockout of \( pcc1 \) yields a less severe growth defect (65). In \( Drosophila\) melanogaster, only \( Pcc1, Kael \) and \( Bud32 \) (termed \( Prpk \) orthologs have been identified thus far (18, 66). Knockout or knockdown of \( Kael, Bud32, Pcc1 \) or \( Sua5 \) in \( Drosophila\) leads to reduced larval size and transformation to pupa (66–69). The sensitivity to \( Kael \) knockdown varies between different tissues, with mitotic tissues especially sensitive while non-proliferating tissues much less sensitive (68). Interestingly, the rare larvae that manage to transform to pupa despite knockdown of \( Prpk \) develop into flies with reduced cell and organ size but without extreme defects in body patterning (70). This result is reminiscent of mutants in the TOR pathway (71) and hints at a potential role for KEOPS in regulating TOR signaling (67, 72). In agreement with this possibility, the growth defects induced by \( Prpk \) knockdown are reversed by expression of an activated mutant of the S6 kinase (70), a downstream effector of the TOR pathway (73). In \( Danio rerio, osgep \) and \( tprkb \) knockout (\( Kael \) and \( Cgl121 \) orthologs respectively) leads to microcephaly and to a shortened life span (74, 75). Similarly, knockout of either of the \( osgep, lage3, prpk \) or \( tprkb \) (\( Kael, Pcc1, Bud32 \) and \( Cgl121 \) orthologs respectively) genes in \( Mus\) \( musculus\) leads to severe microcephaly and early lethality (74). Why the developmental defects resulting from KEOPS dysregulation localize with great frequency to the brain remains an open question.

In line with studies in model organisms, genetic analysis in humans have identified pathogenic mutations that underlie Galloway–Mowat syndrome (GAMOS) in each of the five KEOPS genes and in \( Sua5 \) (74, 76, 77). GAMOS is a recessive genetic disease that manifests in renal and neurologic dysfunctions with a considerable variety of severity (74, 76–78). As a result, GAMOS patients die at very early stages of life. Since GAMOS inducing mutations are common to KEOPS and \( Sua5 \) encoding genes, it is highly likely that reduced \( t^6A \) levels underpins the GAMOS disease phenotype. Consistent with this notion, introduction of GAMOS disease mutations into yeast \( Kael \) inhibits the modification of tRNA \( in\) \( vivo \) (74, 76). KEOPS pathogenic mutations elicit complex cellular phenotypes that likely stem from perturbed translation including activation of the DNA-damage and unfolded protein response, ER stress and defects in actin regulation (74, 76). Some pathogenic mutations in KEOPS exert their effect by disrupting the formation of KEOPS, while others are pathogenic for unknown reasons (74, 77). Further studies are required for a complete understanding of the molecular mechanisms connecting KEOPS mutations to GAMOS.

\( t^6A \) derivative modifications are also important for cellular processes and their dysregulation is implicated in human disease. Mammalian \( Cdkal1 \) for example catalyzes the methythiolation of \( t^6A \) to \( mS-t^6A \) (51). In the absence of
Cdkα1 activity, insulin synthesis is inhibited (51) due to misreading of lysine codons in pro-insulin (79) leading to development of type 2 diabetes.

**HOW KEOPS FUNCTIONS TO MODIFY tRNA**

*Sua5 makes the t6A modification reaction intermediate threonylcarbamoyl adenylate, which Kae1 uses to modify tRNA*

The mechanism by which *Sua5* and *Kae1* collaborate to modify *tRNA* with t6A was delineated through a series of elegant biochemical and structural studies described below. Briefly, the t6A modification of *tRNA* occurs in a reaction with two major steps. In the first step, *Sua5* utilizes threonine, CO2/, HCO3− and ATP precursors to generate the diffusible intermediate molecule threonylcarbamoyl adenylate (TC-AMP) while releasing PPi from ATP. In the second step, TC-AMP and tRNA are bound by *Kae1/TsaD/Qri7*, which then transfer the threonylcarbamoyl moiety from TC-AMP to the Nα acceptor site of A37 in *tRNA* in a reaction that generates AMP (Figure 2A).

*Sua5 generates TC-AMP.* As early as 1974, t6A biosynthesis was known to require the precursors threonine, ATP and HCO3− (80). Knowing that *Sua5* was involved in the t6A modification process and that *Sua5* has ATP binding properties (81,82), Deutsch et al. and Perrochia et al. examined the effect of threonine and *tRNA* on its ATP hydrolysis activity. They found that ATP consumption by *Escherichia coli*, *Pyrococcus abyssi* and *S. cerevisiae* *Sua5* proteins was dependent on threonine (but not tRNA). This hinted that *Sua5* produces an adenylated threonine containing molecule for use in the t6A modification reaction (59,60). Subsequent studies using *Bacillus subtilis* and *S. cerevisiae* *Sua5* proteins showed this molecule to be TC-AMP (Figure 2B) (83). TC-AMP is a relatively short-lived molecule that breaks down to threonylcarbamoyl and AMP moieties. Its stability is strongly affected by pH and temperature, with a half-life of ~20 seconds at pH 7.5 and at 37°C (83). TC-AMP can be stabilized by its binding to *Sua5*, but not by its binding to *Kae1/TsaD/Qri7* enzymes (84). The structural basis by which *Sua5* generates TC-AMP was delineated by X-ray co-crystal and NMR structures of *Sua5* with threonine, ATP and HCO3− precursors (81,85–87). After the discovery that *Sua5* produces TC-AMP, reanalysis of a previously determined crystal structure of *Sulfolobus tokodaii* *Sua5* bound to ATP, revealed serendipitously that the ATP moiety bound in the *Sua5* active site was actually TC-AMP (81,88).

*Kae1/TsaD/Qri7 uses TC-AMP to make t6A.* The discovery that *Sua5* generates the precursor TC-AMP suggested that the *Kae1/TsaD/Qri7* enzymes would employ TC-AMP and tRNA to catalyze the remainder of the t6A modification reaction. In support, *P. abyssi* KEOPS binds tRNA *in vitro* and co-purifies with tRNA when expressed in *E. coli* cells, suggesting that this complex directly acts on tRNA (60). Evidence that the *Kae1/TsaD/Qri7* enzymes carry out the final step of the t6A modification reaction came from the analysis of *Qri7*, which together with *Sua5* are sufficient to carry out the t6A modification of *tRNA* *in vitro* (61). tRNA modification could be achieved with *Sua5* and *Qri7* in chambers separated by a 2 kDa size cut off membrane but only when tRNA was in the same chamber as *Qri7* (61). *In vitro*, it is currently unclear how the unstable TC-AMP is efficiently shuttled to the active site of the *Kae1/TsaD/Qri7* enzymes before its breakdown.

Remarkably, although all three *Kae1* family orthologs perform the same biochemical reaction, each ortholog functions in a completely different protein complex (Figure 2A). The mitochondrial *Qri7* is the most simplified member of the family and catalyzes tRNA modification without any auxiliary proteins (61,89). The bacterial *TsaD* protein depends on its binding to *YaeZ/TsaB*, which is a structural mimic of *Kae1* enzymes (90–92). *TsaD* bound to *TsaB* can bind *tRNA* but has limited catalytic capacity and likely catalyzes only a single round of tRNA modification (93). For multiple rounds of tRNA modification, *TsaD-TsaB* depend on the activity of the ATPase subunit *YjeE/TseA* (59,92–95) for reasons that are not entirely understood. Notably, *TsaB* and *TsaE* are not similar in structure and apparent function to any KEOPS subunits. The significance of the different complex compositions is one of the most exciting mysteries in the field that remain unresolved (see ‘CONCLUDING REMARKS’ section).

The tRNA modification activity of *Sua5* and *Kae1/TsaD/Qri7* enzymes extends to tRNA modifications other than t6A. *Sua5* can generate adenylated variants of TC-AMP using serine or hydroxy norvaline (hn) in place of threonine. In turn, the bacterial *TsaD* protein can utilize hn-carbamoyl adenylate to generate an hn6A modification on substrate tRNAs (96). Since the hn6A modification is also observed in archaeal species (97,98), this would suggest that KEOPS in both archaea and eukaryotes may similarly utilize hn-carbamoyl adenylate to modify tRNA. However, the functional relevance of this modification remains in question.

**Structure of the individual subunits**

*Bud32* is an atypical protein kinase that lacks the infrastructure for canonical protein substrate recognition. Eight crystal structures of complexes containing *Bud32* have been deposited to the PDB (for a list of the structures of proteins that take part in t6A modifying complexes, see Table 1). These include structures of *Bud32* orthologs from archaea, yeast and human in complex with either *Kae1*, Cgi121 or both. These structures reveal a bi-lobal domain organization characteristic of the eukaryotic protein kinase family as exemplified by PKA, a well-studied prototypical kinase (Figure 3A) (for review on the composition of the kinase domain, see (6)). The kinase N-lobe of Bud32 possesses the canonical antiparallel β–strands, buttressed against a helix αC (Figure 3A) (8,9,99). The C-lobe, which is connected to the N-lobe through a flexible hinge region, is predominantly α-helical and greatly abbreviated (i.e. of smaller size) relative to the C-lobes of other eukaryotic protein kinases. One peculiar feature of the abbreviated kinase C-lobe is the absence of all infrastructure typically required for protein substrate recognition, including helix-αG and the activation segment (100) (Figure 3A). A second unique feature of the kinase C-lobe is the presence of a basic C-terminal tail with no apparent homology in other members of the kinome (5).
Figure 2. t⁶A is universally biosynthesized in a two-step reaction by Sua5 and Kae1/TsaD/Qri7 enzymes. (A) Schematic of the universal t⁶A biosynthesis pathway. Step 1- Sua5 utilizes ATP, threonine and HCO₃⁻/CO₂ to catalyze the formation of threonylcarbamoyl adenylate (TC-AMP). Step 2- TC-AMP is used by Kae1/TsaD/Qri7 enzymes to t⁶A modify ANN-decoding tRNA substrates. (top) TsaD functions with the TsaB and TsaE subunits in bacteria. (middle) Kae1 functions with Cgi121, Bud32, Pcc1 and Gon7 subunits in the cytoplasm in eukaryotes and in archaea. (Bottom) Qri7 functions either as a homo-dimer as shown in figure or as a monomer. (B) Chemical structure of threonylcarbamoyl adenylate (TC-AMP). The threonylcarbamoyl moiety is shown in red and the adenylate is shown in black.

ATP binds to the catalytic cleft of Bud32 situated between the N and C- lobes. The ATP binding mechanism in Bud32 is conserved with that of other protein kinases (6,99,101). Specifically, the adenine base of ATP is wedged towards the kinase hinge. The ribose and phosphate groups are coordinated by the conserved β-3 Lys, αC-helix Glu, Gly rich loop, and Asp side chain of the conserved Asp-Phe-Gly (DFG) motif through a bridging Mg/Mn ion. Phospho-transfer function of protein kinases is mediated in part by a catalytic Asp residue within a conserved His-Arg-Asp (HRD) motif in a region termed the catalytic loop (6). This Asp residue functions as a catalytic base to extract a proton from the target hydroxyl group and is conserved in Bud32, suggesting that Bud32 also shares the ability to catalyze phosphate transfer. Accordingly, Bud32 proteins manifest autophosphorylation activity (5,8,9) and its phospho-transfer function is essential for biological function (9,11). However, the absence of structural elements normally required for protein phosphorylation as well as experiments showing that the autophosphorylation acceptor sites are not required for physiological activity (7,102) suggests an alternate role for its phospho-transfer function. In line with this prediction, the chief function of Bud32 appears directed at ATP hydrolysis (i.e. transfer of phosphate to water) (102).

Kae1 has a bi-lobal ASKHA fold. Nine crystal structures of Kae1 alone or in complex with different KEOPS proteins have been deposited to the PDB (Table 1). These include structures of Kae1 orthologs from archaea, yeast and humans either alone, in complex with Pce1, in complex with Pce1 and Gon7, in complex with Bud32 or in complex with Bud32 and Cgi121. Likewise, multiple structures of the Kae1 orthologs Qri7 and TsaD have been solved in their respective complexes. These structures reveal that Kae1 adopts a bi-lobal domain organization characteristic of the ASKHA (acetate and sugar kinases/Hsc70/actin) family of enzymes (Figure 3B) (103,104). The two sубдо-
Table 1. Structures of KEOPS proteins and sub-complexes. All structures reported are crystal structures unless stated otherwise. f = fusion protein. car-AMP = carboxy-adenylate. BK951 = TC-AMP mimetic.

| t^6A modifying complex | Proteins | Species | Ligand | PDB code |
|------------------------|----------|---------|--------|----------|
| **Eukaryotes (cytoplasm) and archaea** | Bud32-Cgi121 | S. cerevisiae | - | 4WWA |
| | Bud32-Cgi121 | S. cerevisiae | AMP | 4WW7 |
| | Bud32-Cgi121 | S. cerevisiae | ADP | 4WW9 |
| | Bud32-Cgi121 | S. cerevisiae | AMP-PnP | 4WW9 |
| | Bud32-Cgi121 | H. sapiens | - | 6WQX |
| Kae1-Bud32(+) | M. jannaschii | AMP-PnP | 2VWB |
| Kae1-Bud32(+) | M. jannaschii | - | 3EN9 |
| Kae1 | P. abyssi | - | 2IVO |
| Kae1 | P. abyssi | ATP | 2IVP |
| Kae1 | P. abyssi | AMP-PnP | 2IVN |
| Kae1-Pcc1 | T. acidophilus, P. furiosus | AMP-PnP | 6WQX |
| Kae1-Pcc1 | M. jannaschii, P. furiosus | - | 5JMV |
| Kae1-Pcc1 | T. acidophilus, P. furiosus | AMP | 5JMV |
| Kae1-Pcc1 | T. acidophilus, P. furiosus | ATP | 2KSY |
| Kae1-Pcc1 | T. acidophilus, P. furiosus | tRNA | 7KJT |
| **Mitochondria** | Qri7-Qri7 dimer | S. cerevisiae | AMP | 4K2S |
| | Qri7 | S. cerevisiae | AMP | 4WUH |
| **Bacteria** | TsαB dimer | E. coli | - | 10KJ |
| | TsαB dimer | S. enterica | - | 2GEM, 2GEL |
| | TsαB dimer | V. parahaemolyticus | - | 3R6M |
| | TsαB dimer | P. aeruginosa | - | 4Y0W, 5BR9 |
| | TsαB dimer | T. maritima | - | 2A6A |
| | TsE | H. influenzae | - | 1FL9 |
| | TsE | H. influenzae | ADP | 1HTW |
| | TsE | B. subtilis | ADP | 5MVR, 5NP9 |
| TsαD-TsαB | B. coli | BK951, AMP | 6ZB1 |
| TsαD-TsαB | E. coli | ATP | 4WQ4 |
| TsαD(12A)mutant-TsαB | E. coli | ATP | 4WQ5 |
| TsαD(V85E)mutant-TsαB | E. coli | ATP | 4WQ5 |
| TsαD-TsαB | S. typhimurium | AMP | 3ZET |
| TsαD-TsαB | S. enterica | ATP-γ-S | 3ZEU |
| TsαD-TsαB-TsαE | T. maritima | ATP, car-AMP | 6N9A |
| TsαD-TsαB-TsαE | T. maritima | AMPcPP | 6S84 |

...mains of Kae1 bear considerable similarity to each other and contain a five stranded β-sheet core flanked by α-helices. In the cavity between the two subdomains, lies a conserved metal binding triad, which co-ordinates a Zn, Mg or Fe ion. Kae1 enzymes have two unique inserts relative to other ASKHA fold enzymes, termed Kae1 specific insert I and II, which are tightly associated with sub-domain 2 and 1 respectively (Figure 3B) (9,104). Unique inserts in ASKHA-fold enzymes commonly contribute to the unique sub-family functions (103), which also holds true for Kae1 proteins (see below).

How does the active site of Kae1 bind its substrates TC-AMP and tRNA? The unstable nature of TC-AMP hindered the ability to obtain atomic level insight into how it is bound by Kae1-family enzymes. However, recently, Kopina et al. developed a non-hydrolysable TC-AMP analogue (termed BK951) and reported its structure bound to the E. coli TsαD-TsαB heterodimer (84). The structure revealed that the adenine base portion of TC-AMP is ‘sandwiched’ between subdomain 2 and Kae1 specific insert 1, similar to the nucleotide binding mechanism of Kae1 enzymes reported in previous structures (8,84,91,95,105). A cation bound by the metal binding triad serves to coordinate the carboxyethyl group of the threonine moiety of TC-AMP (Figure 3B). This binding mode directs the threonylcarbamoyl group towards a cavity in the active site cleft, presumably guiding it to the binding site for tRNA. How the threonylcarbamoyl group is activated by Kae1 for the transfer reaction to N^6 of A37 remains unclear. To date, there is no structure of a Kae1 family enzyme with a tRNA substrate. The expected position of A37 can at best be inferred from the co-structure of TobZ, a NodU family carbamoyl transferase, with its substrate the antibiotic tobramycin (53,88). Since A37 and tobramycin should occupy a similar position in their respective enzyme’s active site, superimposing the structure of the TobZ-tobramycin complex onto Kae1 places A37 in close proximity to TC-AMP and the metal binding triad (Figure 3B).

What features of substrate tRNAs dictates the substrate selectivity of Kae1 enzymes? ANN-decoding tRNAs universally harbor a conserved 36-UAA-38 motif within the anticodon loop with the A37 site at its center (Figure 3C). This motif partially overlaps with the anticodon at positions 34 to 36. The presence of the 36-UAA-38 motif can be...
Figure 3. Structural characteristics of the individual KEOPS subunits. (A) Bud32 is an atypical protein kinase. (left) The structure of *M. jannaschii* Bud32 (PDB 2VWB) is shown with AMP-PnP in the active site (depicted as ATP for simplicity). Highlighted is the unique C-terminal tail and the location of the binding surfaces for Cgi121 and Kae1. (Right) Structure of the kinase domain of *M. musculus* PKA (PDB 1ATP) is shown with ATP in the active site. Highlighted are the canonical activation segment and helix-αG (red) that are distinctively absent from the structure of Bud32. (B) Kae1 is an ASKHA fold enzyme with two unique insertions. The structure of *M. Jannaschii* Kae1 (PDB 2VWB) is shown with the threonylcarbamoyladenylate (TC-AMP) mimetic BK951 (PDB 6Z81) and substrate analogue tobramycin (tob, PDB 3VET) superimposed on its active site. Highlighted are the Kae1-specific inserts I and II (purple and red, respectively) and the location of the binding surfaces for Bud32 and Pcc1. (C) Kae1/TsaD/Qri7 enzymes substrate tRNAs harbor a universal UAA motif in the anticodon loop. A sequence alignment of representative ANN-decoding and non-ANN-decoding tRNAs from *M. jannaschii* is shown highlighting that ANN-decoding tRNAs harbor a 36-UAA-38 motif (red) within the anticodon loop that partially overlaps with the anticodon triplet (positions 34–36). Note that none of the non-ANN-decoding tRNAs have this motif. (D) Pcc1 can homo-dimerize or hetero-dimerize with Gon7. (top) The structure of *P. furiosus* Pcc1 homo-dimer is shown (PDB 3ENC). The dimerization interface and the location of the Kae1 binding surface are highlighted. (bottom) The structure of human Pcc1-Gon7 hetero-dimer is shown (PDB 6GWJ). Note the similarity between the Pcc1 homo-dimer and Pcc1-Gon7 hetero-dimer configuration. (E) Cgi121 is a single domain protein. The structure of *M. jannaschii* Cgi121 (PDB 3ENH) is shown. The location of the binding surfaces for Bud32 and tRNA are highlighted.
explained in part through the roles these nucleotides play in translation (elaborated on above and shown in the structure presented in Figure 1B). However, the UAA motif also plays a role in the stability of the tRNA to be modified by Kae1. Mutation of U36 or A38 disrupts t^6A modification, demonstrating the necessity of the UAA motif for tRNA recognition by Kae1. Insertion of a 36-UAA-38 motif into the anticodon loop of non-substrate tRNAs can lead to their modification (106–108) demonstrating sufficiency in some instances. For example, mutations of either U36 or A38 in tRNA^ile^E. coli does not modify its t^6A modification (107). In contrast, mtRNA^Ile^ naturally harbors a non-t^6A modification by the human Qri7 orthologue OSGEP1 (106,108). In contrast, mtRNA^ile^ naturally harbors a non-canonical 36-UAG-38 motif and is therefore normally not a substrate for t^6A modification. However, a mtRNA^ile^ G38A pathogenic mutation that leads to Leigh syndrome constitutes a canonical 36-UAA-38 motif, inducing the unnatural t^6A modification of this tRNA (106). As noted below, there are other substrate determinants that are distal to anticodon region (see below).

Pcc1 has homo-dimerization capabilities. Six structures of Pcc1 from archaeal, yeast and human species have been deposited to the PDB (Table 1). These structures include those of the Pcc1 homodimer and of the Pcc1-Gon7, Pcc1-Gon7 complex. These structures reveal that Pcc1 is a small protein (~8 kDa in Pyrococcus furiosus for example) composed of a three-strand antiparallel β-sheet flanked on one side by two α-helices (Figure 3D). Pcc1 alone crystallizes as a homo-dimer with an antiparallel configuration of the two monomers. The dimerization of Pcc1 is mediated through a hydrophobic interface, which is generated mainly by the longer of the two helices in each monomer. Mutations in this interface in P. furiosus Pcc1 yields insoluble proteins, suggesting that archaeal Pcc1 is an obligatory dimer (9). The Pcc1 dimer generates a continuous six anti-parallel β-sheet on one face of the dimer packed against four continuous helices of the other face. Since there are no structures of a eukaryotic Pcc1 protein in its apo form, it is unclear if all Pcc1 proteins are obligatory dimers, but the ability to dimerize has been reported for the human Pcc1 ortholog Lage3 (109).

Gon7 is an intrinsically disordered protein that becomes partially structured in binding to Pcc1. Gon7 alone is an intrinsically disordered protein (109). However, a portion of Gon7 assumes a stable structure upon binding to Pcc1 (77,99). Three structures of Gon7 from yeast and human species have been deposited to the PDB (Table 1). These structures include those of Gon7-Pcc1 and Gon7-Pcc1-Kae1 complexes. The structured region of Gon7 (residues 1 to 20 and residues 25–50, human nomenclature used) form two anti-parallel β-sheets and a single α-helix (Figure 3D). The C-terminus of Gon7 residues 51–100 is enriched with acidic residues, a feature that is conserved among Gon7 orthologs (77,99).

Cgi121 structure. Nine crystal structures and 1 solution structure solved by NMR of Cgi121 alone or in complex with other KEOPS proteins or substrate tRNA have been deposited to the PDB (Table 1). These include the structures of Cgi121 orthologs from archaea, yeast, and human either alone, in complex with Bud32, in complex with Bud32-Kae1 or in complex with tRNA^Lys^UUU. These structures revealed that Cgi121 is a single domain protein containing a core anti-parallel β-sheet of 3 to 5 strands sandwiched between α-helices on either side (9) (Figure 3E).

Basis for KEOPS subunit interaction with each other and with substrate tRNA

Overview: KEOPS subunits adopt a linear binding topology. The architecture and tRNA binding mechanism of KEOPS have been elucidated by solving a series of binary and ternary structures of its subunits (Table 1). In brief, KEOPS adopts a linear binding architecture (Figure 4A). The two enzymes Kae1 and Bud32 bind directly to each other at the center of KEOPS. On opposing ends, Bud32 binds to Cgi121 and Pcc1 binds to Kae1 (Figure 4A). Bound to Kae1, Pcc1 exists as a homo-dimer or as a hetero-dimer with Gon7 (Figure 3D).

The kinase domain of Bud32 ‘wraps’ around subdomain 2 of Kael. Consistent with the initial yeast two hybrid analysis (11), Bud32 binds directly to Kae1. The Kae1 binding surface in Bud32 is composed of regions from both N and C lobes (Figure 4A). The reciprocal binding surface on Kae1 is composed of subdomain 2, including the Kae1-specific insert I. The highly conserved C-terminal tail of Bud32 interacts with the active site of Kae1 (Figure 4A) (8,9). While not essential for binding to Kae1, the C-terminal tail of Bud32 is required the ability of KEOPS to modify tRNA in vitro and for KEOPS functionality in vivo as assessed in yeast (9,102).

Pcc1 binds to subdomain 1 of Kael and can control KEOPS dimerization. The Pcc1 binding surface on Kae1 is composed primarily by helices α1 and α2, which pack against helices α1 and α2 of Pcc1 (Figure 4A). The Kael-Pcc1 binding mechanism is similar to that of TsaD-TsaB and to the dimerization mechanism of Qri7. The significane of this convergent binding mode is currently unclear (see CONCLUDING REMARKS section). The Pcc1 homo-dimer presents two non-overlapping Kael-binding surfaces, raising the enticing possibility that KEOPS might dimerize through Pcc1. In support, archaeal KEOPS with wild-type Pcc1 forms dimers in solution (60,110). In contrast, KEOPS reconstituted with a synthetic asymmetric Pcc1-Pcc1 fusion protein with disruptive mutations on one of two Kae1 binding surfaces is monomeric (110). Since monomeric and dimeric KEOPS complexes have comparable tRNA modification activity in vitro, the role of this dimerization is unclear (110) (for a model of a KEOPS dimer see (9)). Gon7 binds to Pcc1 through the surface that mediates the dimerization of archaeal Pcc1 (Figure 3D), Gon7 was therefore
Figure 4. A model for the KEOPS-tRNA holo-enzyme-substrate complex. (A) Model of KEOPS reveals the basis for the interactions between its subunits and shows that they adopt a linear binding. A cartoon representation with semi-transparent surface of a model of the KEOPS complex. The model was generated using the structures of the Cgi121-Bud32-Kae1, Kae1-Pcc1 and Gon7-Pcc1-Kae1 sub-complexes (PDB 3ENH, 3ENO and 6GWJ respectively, see Table 1). Gon7 (gray) binds to Pcc1 (yellow), which binds to Kae1 (blue). Kae1 binds also to Bud32 (green), which binds Cgi121 (purple). Bud32 (green) binds to subdomain 2 of Kae1 (light blue) through an extensive surface that is composed of the N and C lobes. The C-tail of Bud32 interacts with the active site cleft of Kae1. Pcc1 (yellow) binds to helices α1 and α2 of subdomain 1 of Kae1 (dark blue) in vicinity of the active site cleft of Kae1. Cgi121 (purple) binds to helix αC and β-strand 4 of the N-lobe of Bud32 (dark green) through helix α8. (B) A model of KEOPS-tRNA holoenzyme substrate complex shows the basis for substrate interaction by KEOPS. For clarity, proteins are shown in surface representation and tRNA in cartoon representation and complex is shown in two different orientations. According to the model, Cgi121 (pink) binds the tRNA CCA tail (red) via helices α1, α3 and α4 (purple). Kae1 (blue) and Pcc1 (yellow) interact with the tRNA anticodon loop and Bud32 (green) and Kae1 interact with the tRNA D-arm. Gon7 (gray) does not participate in tRNA binding. This binding mode directs A37 of tRNA (magenta) into the active site of Kae1 and positions the CU motif of tRNA (red) to the interface between Bud32 and Kae1.
proposed to regulate the dimerization function of Pcc1 and thus KEOPS dimerization (99). In support, yeast KEOPS that includes Gon7 is monomeric in solution with a 1:1 binding stoichiometry between its all five subunits (99). Furthermore, 4-subunit human KEOPS (without hGon7) forms dimers in solution while 5-subunit human KEOPS is monomeric (109,111). Lastly hGon7 appears to enhance or stabilize the expression levels of KEOPS in vivo since in the absence of hGon7 overall expression levels of KEOPS subunits are repressed (77). These findings might hint indirectly that KEOPS dimerization could regulate its degradation in eukaryotes, however this possibility requires further exploration.

Cgi121 binds and regulates Bud32 and recruits tRNA to KEOPS. Consistent with the initial yeast two-hybrid analysis (15), Cgi121 binds directly to Bud32. In archaeal protein structures, the Cgi121 binding surface on Bud32 is composed primarily of N-lobe helix–αC and β-strand 4 (Figure 4A). The reciprocal binding surface on archaeal Cgi121 is composed primarily of helix αB. The Cgi121 binding mode in the eukaryotic proteins is essentially the same with slight elaborations. The N-lobe of eukaryotic Bud32 is slightly larger than its archaeal ortholog with an extra helix (termed αA-helix) that participates in binding eukaryotic Cgi121. The binding mode between Bud32 and Cgi121 is reminiscent of cyclin-dependent kinases binding to cyclins (112). Thus, not unexpectedly, Cgi121 potentiates both the ATPase and low level autophosphorylation activity of Bud32 (note that the autophosphorylation activity is unlikely to be physiologically relevant) (9,101,102). Structures of Bud32 in complex with Cgi121 reveal an active-like conformation with productively positioned catalytic elements (99,101). To date, no structures of Bud32 alone have been reported for comparison.

Apart from its ability to bind Bud32, Cgi121 has intrinsic ability to bind tRNA. This ability is essential for the efficient recruitment of tRNA to KEOPS. A co-crystal structure of a Cgi121-tRNA complex revealed that tRNA binding by Cgi121 is directed almost exclusively towards the 3′ CCA tail (positions 74–76, 5′-C74-C75-A76-3′) (Figure 4B) (102). The binding surface on Cgi121 for tRNA is composed primarily of α-helices 1, 3 and 4. The CCA tail is a universal element present in all mature tRNAs that plays multiple essential roles in the functionality and regulation of tRNAs. Perhaps most importantly, the terminal nucleotide of the CCA tail serves for the attachment of amino acids (aminoacylation) by aminoacyl tRNA synthetases (113). The CCA tail also plays an important role in quality control of tRNAs and translation. For example, the CCA tail is excised by ANKZF1 to help the recycling of stalled ribosomes (114) and a second CCA tail is added by the CCA-adding enzyme to structurally unstable tRNAs as a signal for their destruction by the tRNA degradation machinery (115). Cgi121 tRNA binding activity is dependent on the presence of an intact and non-aminocylated CCA tail. Specifically, deletion of the CCA tail, addition of a second CCA or attachment of bulky groups to the 3′ of A76 inhibits tRNA binding by Cgi121 (102). This dependency renders KEOPS activity highly sensitive to the status of the CCA tail of tRNA substrates. The biological role of this dependency is still not understood (see CONCLUDING REMARKS section).

A model for the KEOPS-tRNA holo-enzyme-substrate complex. A model for the KEOPS-tRNA holo-enzyme-substrate complex was generated using the Cgi121-tRNA structure and structures of KEOPS sub-complexes (Figure 4B) (102). According to this model, all four core KEOPS subunits (i.e. Cgi121, Bud32, Kae1 and Pcc1) form an extended tRNA binding surface that is highly complementary to the structure of the tRNA. The tRNA is anchored on both ends to the KEOPS complex with the CCA end bound by Cgi121 and the anticodon loop interacting with Kae1 and Pcc1. At its center, the tRNA D-arm is predicted to contact Bud32 and Kae1 (Figure 4B). Therefore, the tRNA binding mode of KEOPS seems to interrogate the entire ‘inner side’ of the L shape of tRNA. Although all subunits are predicted to take part in contacting tRNA, only mutations on Bud32 and Cgi121 adversely affect tRNA binding affinity. Bud32 does not bind tRNA independently but serves to potentiate the tRNA binding activity of Cgi121. Bud32 achieves this likely by forming secondary contacts with tRNA and by allosterically inducing a high affinity state in the CCA tail binding surface of Cgi121 (102). Despite not contributing to binding affinity, the predicted tRNA binding surface in Kae1 and Pcc1 are none-the-less vital for tRNA modification activity (102).

The ATPase activity of Bud32 is regulated by tRNA binding to KEOPS. The ATPase activity of Bud32 is essential for the ability of KEOPS to modify tRNA substrates in vitro and is required for Bud32’s essential function in vivo (9,102). Interestingly, activation of Bud32 ATPase activity is strongly potentiated by tRNA, but only in the context of the Cgi121-Bud32-Kae1 sub-complex or the full KEOPS complex (102). Mutations in any of the KEOPS subunits on the predicted tRNA binding surfaces inhibit the ability of tRNA to activate Bud32 ATPase. How tRNA substrates regulate Bud32 ATPase activity remains an unsolved question. However, in archaea this ability is not correlated with the anticodon loop UAA motif but rather the presence of a di-nucleotide CU motif located at positions 10–11 within the tRNA D-arm (102). This motif is almost completely conserved in ANN-decoding tRNAs and absent from almost all non-ANN-decoding tRNAs with the sole exception of tRNAAla. In the KEOPS-tRNA complex, the CU motif is predicted to localize at the interface between Bud32 and Kae1 in proximity to the active site of Bud32 (Figure 4B) (102). The role this element has in the activation of Bud32 and in the ability of KEOPS to modify tRNA awaits further investigation.

CONCLUDING REMARKS AND QUESTIONS REMAINING TO BE ANSWERED

Although many questions about how KEOPS functions have been answered over the past 15 years, many more remain to be addressed, as outlined below.
Why has KEOPS evolved added complexity?

Fifteen years of studying the structure and function of KEOPS have revealed an intricate mechanism of action. Unlike Qri7, which functions without binding partners, the activity of Kae1 is dependent on at least three binding partners. This raises the question of what function does this added complexity serve when in principle Kae1 could function like Qri7?

We consider two tantalizing possibilities. First, the machinery built around Kae1 could enable the regulation of its tRNA modifying activity. What benefit could there be from KEOPS downregulation? Although the total absence of $t^\delta A$ is harmful for cells, temporal control of $t^\delta A$ levels could be beneficial for cells as a means to control translation under specific cellular conditions. For example, Ser and Arg can be coded by $t^\delta A$-dependent (AGT/C and AGG/A respectively) and $t^\delta A$-independent codons (TCT/C/A/G and CG/T/C/A/G respectively). As such, KEOPS downregulation would favor the translation of mRNA transcripts enriched with Ser and Arg codons that are not dependent on $t^\delta A$ modification. Thaiville et al. reported that in yeast mRNAs encoding $\sim$300 proteins exclusively contain Arg codons that are $t^\delta A$-dependent (44). These would be prime candidates for downregulation in response to KEOPS inhibition.

Translation is regulated by modulation of other types of tRNA modifications at positions 34 and 37 (for examples, see references (116–119)) and mRNAs that are differentially translated due to the presence or absence of tRNA modifications have been termed modification tunable transcripts (MoTTs) (120). If the temporal regulation of $t^\delta A$ is exploited for regulatory reasons, the added complexity of KEOPS provides multiple opportunities to achieve this end. For example, the tRNA binding mechanism of KEOPS renders its function sensitive to Cgi121 expression levels. The short half-life of Cgi121 in yeast ($\sim$30 min) (9) and its sub-stoichiometric level relative to the other subunits (121), make it an excellent restriction point for regulation.

A second possibility is that the complexity of KEOPS addresses a specific challenge or need of modifying cytoplasmic tRNAs that is not required for mitochondrial tRNAs. The cytoplasm contains a much larger number of tRNAs, both ANN and non-ANN-decoding, which synthesizes a far larger number of proteins relative to mitochondria. The human mitochondrial genome transcribes only 22 different tRNAs (122), of which 5 are substrates for $t^\delta A$ modification (28,123). These tRNAs are needed for the synthesis of only 13 mitochondrial proteins (122). The human nuclear genome in contrast contains approximately 500 tRNA genes, including hundreds that are not KEOPS substrates, and these participate in the synthesis of 20 000–30 000 different proteins. Perhaps Kae1 requires its auxiliary subunits to deal with the greater diversity of tRNAs and the greater demand for modified tRNAs in the cytoplasm relative to those observed in mitochondria. In support of this possibility, while Qri7 and KEOPS display comparable catalytic rates in purified systems, massive overexpression of cytoplasmic Qri7 is required for its ability to rescue the growth defects resulting from the knockout of KEOPS (61). This in turn raises the question of what specific advantage the added complexity of KEOPS actually provides relative to Qri7? Possible advantages include the ability to read-out tRNA features beyond the anticodon loop (102). This enhancement of substrate binding mechanism could serve as a substrate quality control mechanism that would help discriminate between functional and non-functional tRNA substrates, which might perturb the translation machinery. A second advantage may be the ability to coordinate $t^\delta A$ modification with other tRNA modification and processing events. tRNAs undergo many processing steps from their initial transcription to their degradation, and many of these steps occur in a hierarchical manner (124,125). In support of this possibility, since a subset of tRNAs contain an intron that is located between U36 and A37, the $t^\delta A$ modification by KEOPS can only occur after splicing. Likewise, the tRNA binding mechanism entails that $t^\delta A$ modification will occur after CCA tailing and before aminoacylation (102). How the activity of KEOPS is affected by other tRNA modifications and processing events and what role this might play awaits further study.

How does the ATPase activity of Bud32 promote the $t^\delta A$ modification of tRNA by Kae1?

The ATPase activity of Bud32 is required for KEOPS activity in vitro (102) and in vivo (9). However, it is still unclear how this activity exerts a specific effect on the tRNA $t^\delta A$ modification reaction. Since the ATPase activity of Bud32 precedes the tRNA modification activity of Kae1 (102) it is reasonable to infer that it serves to activate Kae1. A candidate region that could mediate an activating effect on Kae1 is the conserved C-terminal tail of Bud32. This region is dispensable for the activation of ATPase activity by tRNA (and thus exerts its essential effect downstream of the ATPase activating event) but is required for the tRNA modification activity of Kae1. How could this region of Bud32 mediate the activation of Kae1? Structures of Kae1-Bud32 complexes reveal that the C-terminus of Bud32 resides near the Kae1 active site and the Pcc1 binding surface on Kae1 (Figure 4A). From this perch, the C-terminal tail of Bud32 is well placed to help remodel the active site of Kae1 into a productive conformation. Alternatively, its position could serve in some capacity to collaborate with Pcc1. Either way, since ATP binding and hydrolysis by protein kinases is known to influence inter-lobe closure (126), Bud32’s catalytic function could generate movements of the C-tail of Bud32 to enable its supporting role in Kae1 catalytic function.

What are the precise roles of Pcc1 and Gon7 in the activity of KEOPS?

One of the main unanswered issues in the KEOPS field is the precise role of Pcc1. Pcc1 binds close to the active site of Kae1. Interestingly, the absence of Pcc1 has the same effect on KEOPS as the deletion of the C-tail of Bud32, namely the absence of both has minimal effect on the tRNA binding activity of KEOPS or the activation of Bud32 ATPase activity in response to binding tRNA (102). However, both are necessary for the tRNA modification activity of KEOPS in vitro and in vivo (9,102). The proximity of Pcc1
to the active site of Kae1 indicates that Pcc1 might function to activate Kae1, similar to the C-tail of Bud32. Providing an initial clue of Pcc1’s involvement in the regulation of the \( t^6A \) catalytic function of KEOPS, tRNA binding to KEOPS induces perturbations to the binding interface between the Kae1 and Pcc1 subunits as measured by hydrogen deuterium exchange mass-spectrometry (102).

Another possible role for Pcc1 in \( t^6A \) modification could be to guide the anticodon loop of the tRNA into the active site of Kae1. Pcc1 is predicted to directly contact tRNA in close vicinity of the anticodon loop harbouring the A37 modification site (Figure 4B). Whatever the exact role of Pcc1, mutation of an invariant Arg residue on its inferred tRNA binding surface (Arg 63 in \( P. furiosus \) Pcc1) disrupts \( t^6A \) modification function (102).

The role of the Gon7 subunit in KEOPS also remains elusive. Since Gon7 proteins have not been identified in archaeal species (18) they appear to be an adaptation specific to eukaryotic KEOPS. Thus, Gon7 does not likely fulfill an essential role in the biochemical activity of KEOPS. In support, human KEOPS is active in vitro in the absence of Gon7 (102,109). In vivo however, the absence of Gon7 yields a strong phenotype that is similar to the absence of other KEOPS subunits (1) and in humans mutations in Gon7 can cause GAMOS (77). Interestingly, Gon7 possesses a highly conserved acidic C-terminus (residues 51–100 in human Gon7) that does not participate in binding to Pcc1 and remains unstructured in the Pcc1-Gon7 complex. Uncovering what this region does could provide insight into the essential role of Gon7 in cells.

Conclusion and relation to other \( t^6A \) modifying complexes

A great amount has been learned over the past 15 years of intensive KEOPS study. While not covered here in detail, parallel research efforts on the structure (Table 1) and function of the orthologous tRNA modification machinery in bacteria (for examples, see references (59,93–95) by the Van Tilbeurgh, Creyc-Lagard, Iwata-Reuyl and Swairjo labs) have been similarly informative. These efforts have shed light into functions shared across the Kae1/TsaD/Qri7 family, and functions specific to the divergent protein complexes each enzyme operates in (see Figure 2A for schematic representations of the complexes). Comparing KEOPS with these orthologous systems raises two interesting points for consideration.

First, as noted above, a striking common feature of the three different \( t^6A \) modifying enzymes is the conserved binding mode between Kae1/TsaD/Qri7 and their respective binding partners Pcc1/TsaB/Qri7. Each of these interactions (i.e. Kae1-Pcc1, TsaD-TsaB and Qri7-Qri7) is essential for tRNA modification. The precise function of this conserved binding mode remains elusive, but once assigned for Kae1, TsaD or Qri7, it will likely be informative for all three systems. A second striking feature is the dependency of Kae1 and TsaD on the ATPase subunits Bud32 and TsaE respectively for efficient tRNA modification. Despite this apparent similarity, Bud32 and TsaE play distinct roles in their respective complexes. In KEOPS, Bud32 promotes tRNA binding and its ATPase activity is required for tRNA modification (102) while in the TsaD-TsaB-TsaE complex, TsaE competes with tRNA binding (93) but its ATPase activity is required for maximal tRNA modification (92,93,95). How precisely the ATPase activities of Bud32 and TsaE exert their effects on Kae1 and TsaD respectively and if there are similarities in these effects remains to be determined.

No doubt, future studies will illuminate answers to these remaining questions and others that unify and differentiate the mitochondrial, bacterial and eukaryotic/archaeal \( t^6A \) modification machineries and their important biological relevance.

DATA AVAILABILITY

The structures used for generating Figures 1B, 3A, 3B, 3D, 3E and 4A can be found on the PDB website. The model presented in Figure 4B can be provided upon request.

ACKNOWLEDGEMENTS

We wish to thank Ms. Susan Kelso, Dr Salima Daou, professor Ute Kothe, professor Mark Bayfield, Ms. Jennifer Porrat and professor Daniel Durocher for their critical reading of this manuscript and their useful comments. We wish to thank Ms. Samara Ona for her help in generating the KEOPS-tRNA model presented in Figure 4B.

FUNDING

CIHR [FDN 143277 to F.S.]. Funding for open access charge: CIHR [FDN 143277].

Conflict of interest statement. None declared.

REFERENCES

1. Downey, M., Houlsworth, R., Maringele, L., Rollie, A., Brehme, M., Galicia, S., Guillard, S., Partington, M., Zubko, M. K., Krogran, N. J. et al. (2006) A genome-wide screen identifies the evolutionarily conserved KEOPS complex as a telomere regulator. Cell, 124, 1155–1168.
2. Kisseleva-Romanova, E., Lopreiato, R., Baudin-Baillieu, A., Rousselle, J. C., Ilan, L., Hofmann, K., Namane, A., Mann, C. and Libri, D. (2006) Yeast homolog of a cancer-testis antigen defines a new transcription complex. EMBO J., 25, 3576–3585.
3. Koonin, E. V. (2003) Comparative genomics, minimal gene-sets and the last universal common ancestor. Nat. Rev. Microbiol., 1, 127–136.
4. Ni, L. and Snyder, M. (2001) A genomic study of the bipolar bud site selection pattern in Saccharomyces cerevisiae. Mol. Biol. Cell, 12, 2147–2170.
5. Stocchetto, S., Marin, O., Carignani, G. and Pinna, L. A. (1997) Biochemical evidence that Saccharomyces cerevisiae YGR262c gene, required for normal growth, encodes a novel Ser/Thr-specific protein kinase. FEBS Lett., 414, 171–175.
6. Komev, A. P. and Taylor, S. S. (2015) Dynamics-driven allosterie in protein kinases. Trends Biochem. Sci., 40, 628–647.
7. Facchin, S., Lopreiato, R., Stocchetto, S., Arrigoni, G., Cesaro, L., Marin, O., Carignani, G. and Pinna, L. A. (2002) Structure-function analysis of yeast PD261/Bud32, an atypical protein kinase essential for normal cell life. Biochem. J., 364, 457–463.
8. Becker, A., Lopreiato, R., Grellle, M., Collin, B., Forrer, P., Libri, D. and van Tilbeurgh, H. (2000) Structure of the archaeal Kae1/Bud32 fusion protein MJ1130: a model for the eukaryotic EKC/KEOPS subcomplex. EMBO J., 27, 2340–2351.
48. Miyauchi, K., Kimura, S. and Suzuki, T. (2013) A cyclic form of N6-threonylcarbamoyladenosine as a widely distributed RNA hypermodification. Nat. Chem. Biol., 9, 105–111.

49. Matyszewski, M., Wojciechowski, J., Miyauchi, K., Gdaniec, Z., Wolf, W.M., Suzuki, T. and Sochacka, E. (2017) A hydantoin isomer of cyclic N6-threonylcarbamoyladenine (ct6A) is present in tRNAs. Nucleic Acids Res., 45, 2137–2149.

50. Kimura, S., Miyauchi, K., Ikeuchi, Y., Thiaville, P.C., Crecy-Lagarde, V. and Suzuki, T. (2014) Discovery of the beta-barrel-type RNA methyltransferase responsible for N6-methylation of N6-threonylcarbamoyladenosine in tRNAs. Nucleic Acids Res., 42, 9350–9365.

51. Wei, F.Y., Suzuki, T., Watanabe, S., Kimura, S., Kaitisuka, T., Fujimura, A., Matsui, H., Atta, M., Michiue, H., Fontecave, M. et al. (2011) Deficit of tRNA(Lys) modification by Cdkk1 causes the development of type 2 diabetes in mice. J. Clin. Invest., 121, 3598–3608.

52. Na, J.G., Pinto, I. and Hampey, M. (1992) Isolation and characterization of SUA5, a novel gene required for normal growth in Saccharomyces cerevisiae. Genetics, 131, 791–801.

53. Aravind, L. and Koonin, E.V. (1999) Gleaning non-trivial structural, functional and evolutionary information about proteins by iterative database searches. J. Mol. Biol., 287, 1023–1040.

54. Jabbouri, S., Relic, B., Hanin, M., Kamalapriya, P., Burger, U., Prom, D., Prom, J.C. and Broughton, W.J. (1998) nolO and nol (ManII) of Rhizobium sp. NGR284 are involved in 3-O-carmaboylation and 2-O-methylation of nod factors. J. Biol. Chem., 273, 12047–12055.

55. Daugeron, M.C., Lenstra, T.L., Frizzarin, M., El Yacoubi, B., Liu, X., Baudin-Bailleau, A., Linzjaard, P., Decourty, L., Saveanu, C., Jacquier, A. et al. (2011) Gcn4 misregulation reveals a direct role for the evolutionary conserved EKC/KEOPS in the t6A modification of tRNAs. Nucleic Acids Res., 39, 6148–6160.

56. Srinivasan, M., Mehta, P., Yu, Y., Pruger, E., Koonin, E.V., Karzai, A.W. and Sternglanz, R. (2011) The highly conserved KEOPS/ECK complex is essential for a universal tRNA modification, t6A. EMBO J., 30, 873–881.

57. Galperin, M.Y. and Koonin, E.V. (2004) ‘Conserved hypothetical’ proteins: prioritization of targets for experimental study. Nucleic Acids Res., 32, 5452–5463.

58. Hinnebusch, A.G. (2005) Translational regulation of GNC4 and the general amino acid control of yeast. Annu. Rev. Microbiol., 59, 407–450.

59. Deutsch, C., El Yacoubi, B., de Crey-Lagarde, V. and Iwata-Reuy, D. (2012) Biosynthesis of threonylcarbamoyladenosine (t6A), a universal RNA nucleoside. J. Biol. Chem., 287, 13666–13673.

60. Perrochia, L., Crozat, E., Hecker, A., Zhang, W., Bareille, J., Collinet, B., van Tilbeurgh, H., Forterre, P. and Basta, T. (2013) In vitro biosynthesis of a universal t6A RNA modification in Archaea and Eukarya. Nucleic Acids Res., 41, 1953–1964.

61. Wan, L.C., Ma, D.Y., Neculai, D., Strecker, J., Chiotti, D., Kurinov, I., Poda, G., Thevakumaran, N., Yuan, F., Szilard, R.K. et al. (2013) Reconstitution and characterization of eukaryotic N6-threonylcarbamoylation of tRNA using a minimal enzyme system. Nucleic Acids Res., 41, 6332–6346.

62. Pollo-Oliveira, L., Klassen, R., Davis, N., Cifcici, A., Bacusmo, J.M., Martinelli, M., DeMott, M.S., Begley, T.J., Dedon, P.C. and Schaffrath, R. et al. (2020) Loss of Elongator- and KEOPS-Dependent tRNA Modifications Leads to Severe Growth Phenotypes and Protein Aggregation in Yeast. Biomolecules, 10, 322.

63. Peng, J., He, M.H., Duan, Y.M., Liu, Y.T. and Zhou, J.Q. (2015) Inhibition of telomere recombination by inactivation of KEOPS subunit Cty121 promotes cell longevity. PLoS Genet., 11, e1005071.

64. Yoshi, K., Bhatt, M.J. and Farabaugh, P.J. (2018) Codon-specific effects of tRNA anticond loop modifications on translational misreading errors in the yeast Saccharomyces cerevisiae. Nucleic Acids Res., 46, 10531–10539.

65. Naoa, A., Thiaville, P.C., Altman-Price, N., Cohen-Or, I., Allers, T., de Crey-Lagarde, V. and Gophna, U. (2012) A genetic investigation of the KEOPS complex in halophilic Archaea. PLoS One, 7, e43013.

66. Rojas-Benitez, D., Eggers, C. and Glavic, A. (2017) Modulation of the proteostasis machinery to overcome stress caused by diminished levels of t6A-modified tRNAs in Drosophila. Biomolecules, 7, 25.
102. Beenstock, J., Kasai, T., Akasaka, R., Higashijima, K., Terada, T., Kigawa, T., Shinkin, A., Besho, Y. and Yokoyama, S. (2011) Crystal structure of Sulfolobus tokodaii Suu5 complexed with L-threonine and AMPPNP. *Proteins*, 79, 2065–2075.

103. Hurley, J.H. (1996) The sugar kinase/heat shock protein 70/actin superfamily: implications of conserved structure for mechanism. *Annu. Rev. Biophys. Biomol. Struct.*, 25, 137–162.

104. Hecker, A., Leulliot, N., Gadelle, D., Graille, M., Justome, A., Dorlet, P., Brochier, C., Quevillon-Cheruel, S., Le Cam, E., van Tilbeurgh, H. et al. (2007) An archaean orthologue of the universal protein Kael is an iron metalloprotein which exhibits atypical DNA-binding properties and apurinic-endonuclease activity in vitro. *Nucleic Acids Res.*, 35, 6042–6051.

105. Tomina, T., Kobayashi, K., Ishii, R., Ishitani, R. and Nureki, O. (2014) Structure of Saccharomyces cerevisiae mitochondrial Qr7 in complex with AMP. *Acta Crystallogr. F Struct. Biol. Commun.*, 70, 1009–1014.

106. Lin, H., Miyaiuchi, K., Harada, T., Okita, R., Takeshita, E., Komaki, H., Fujioka, K., Yagasaki, H., Goto, Y.I., Yanaka, K. et al. (2018) CO2-sensitive tRNA modification associated with human mitochondrial disease. *Nat. Commun.*, 9, 1875.

107. Morin, A., Auxilien, S., Senger, B., Tewari, R. and Grosjean, H. (1998) Structural requirements for enzymatic formation of threonylcarbamoyl-AMP. *Nucleic Acids Res.*, 26, 3126–3132.

108. Parthier, C., Gorlich, S., Jaenecke, F., Breithaupt, C., Brauer, U., Fandrich, U., Clausnitzer, D., Wehmeier, U.F., Bottcher, C., Scheel, D. et al. (2012) The O-carboxamidotransferase TobZ catalyzes an ancient enzymatic reaction. *Angew. Chem. Int. Ed. Engl.*, 51, 4046–4052.

109. Zhou, J.B., Wang, Y., Zeng, Q.Y., Meng, S.X., Wang, E.D. and Zhou, X.L. (2020) Molecular basis for t6A modification in human mitochondria. *Nucleic Acids Res.*, 48, 3181–3194.

110. Nichols, C.E., Johnson, C.C., Lockyer, M., Charles, I.G., Lamb, H.K., Hawkins, A.R. and Stammers, D.K. (2006) Structural characterization of Sulfolobus thermophilum YeaZ, an M22 O-sialoglycoprotein endopeptidase homolog. *Proteins*, 64, 111–123.

111. Nichols, C.E., Lamb, H.K., Thompson, P., Olami, K., Lockyer, M., Charles, I., Hawkins, A.R. and Stammers, D.K. (2013) Crystal structure of the dimer of two essential Salmonella typhimurium proteins, YgiD & YeaZ and caloriometric evidence for the formation of a ternary YgiD-YeaZ-YjeE complex. *Protein Sci.*, 22, 628–640.

112. Zhang, W., Collinet, B., Perrochia, L., Durand, D. and van Tilbeurgh, H. (2015) The ATP-mediated formation of the YgiD-YeaZ-YjeE complex is required for the biosynthesis of tRNA t6A in Escherichia coli. *Nucleic Acids Res.*, 43, 1804–1817.

113. Luthra, A., Swinehart, W., Bayoos, S., Phan, P., Stec, B., Iwata-Reuyl, D. and Swain, D.A. (2018) Structure and mechanism of a bacterial t6A biosynthesis system. *Nucleic Acids Res.*, 46, 1395–1411.

114. Missoory, S., Planqueel, S., Li de la Sierra-Gallay, I., Zhang, W., Liger, D., Durand, D., Damnak, R., Collinet, B. and van Tilbeurgh, H. (2018) The structure of the TsaB/TsaD/TsaE complex reveals an unexpected mechanism for the bacterial t6A tRNA-modification. *Nucleic Acids Res.*, 46, 5850–5860.

115. Luthra, A., Paranagama, N., Swinehart, W., Bayoos, S., Phan, P., Quach, V., Schiffer, J.M., Stec, B., Iwata-Reuyl, D. and Swain, D.A. (2019) Conformational communication mediates the resut step in t6A biosynthesis. *Nucleic Acids Res.*, 47, 6551–6567.

116. Swinehart, W., Deutsch, S., Sarachan, K.L., Luthra, A., Bacuso, J.M., de Crecy-Lagard, V., Swain, D.A., Agris, P.P. and Iwata-Reuyl, D. (2020) Specificity in the biosynthesis of the universal tRNA nucleoside N (6-threonylcarbamoyl adenosine (t6A)-TsaD is the gatekeeper. *RNA*, 26, 1094–1103.

117. Wolff, P., Villette, C., Zunz, J., Heintz, D., Antoine, L., Chane-Woon-Ming, B., Droogmans, I., Grosjean, H. and Westhof, E. (2020) Comparative patterns of modified nucleotides in individual tRNA species from a mesophilic and two thermophilic archaea. *RNA*, 26, 1957–1975.

118. Yu, N., Jora, M., Solivio, B., Thakur, P., Acedo-Rocha, C.G., Rueda, L.L., de Crecy-Lagard, V., Aoki, M., Yari, B. and Limbach, P.A. (2019) tRNA Modification Profiles and Codon-Decoding Strategies in Methanococcos jannaschii. *Microbiology*, 201, e00690-18.

119. Zhang, W., Collinet, B., Graille, M., Daugeron, M.C., Lazar, N., Libri, D., Durand, D. and van Tilbeurgh, H. (2015) Crystal structures of the Gon7/Pcc1 and Bus32/Cgi121 complexes provide a model for the complete yeast KEOPS complex. *Nucleic Acids Res.*, 43, 3358–3372.

120. Dar, A.C., Dever, T.E. and Sicheri, F. (2005) Higher-order substrate recognition of eIF2alpha by the RNA-dependent protein kinase PKR. *Cell*, 122, 887–900.

121. Li, J., Ma, X., Banerjee, S., Chen, H., Ma, W., Bode, A.M. and Dong, Z. (2021) Crystal structure of the human PRPK-TPK-RBK complex. *Commun Biol.*, 4, 167.

122. Beenstock, J., Osa, S.M., Porat, J., Orlicky, S., Wan, L.C.K., Cuccarelli, D.F., Maisonneuve, P., Szlak, R.K., Yin, Z., Setiaputra, D. et al. (2020) A substrate binding model for the KEOPS tRNA modifying complex. *Nat. Commun.*, 11, 6233.

123. Hurley, J.H. (1996) The sugar kinase/heat shock protein 70/actin superfamily: implications of conserved structure for mechanism. *Annu. Rev. Biophys. Biomol. Struct.*, 25, 137–162.

124. Hecker, A., Leulliot, N., Gadelle, D., Graille, M., Justome, A., Dorlet, P., Brochier, C., Quevillon-Cheruel, S., Le Cam, E., van Tilbeurgh, H. et al. (2007) An archaean orthologue of the universal protein Kael is an iron metalloprotein which exhibits atypical DNA-binding properties and apurinic-endonuclease activity in vitro. *Nucleic Acids Res.*, 35, 6042–6051.

125. Tomina, T., Kobayashi, K., Ishii, R., Ishitani, R. and Nureki, O. (2014) Structure of Saccharomyces cerevisiae mitochondrial Qr7 in complex with AMP. *Acta Crystallogr. F Struct. Biol. Commun.*, 70, 1009–1014.
121. Ghaemmaghami, S., Huh, W.K., Bower, K., Howson, R.W., Belle, A., Dephoure, N., O’Shea, E.K. and Weissman, J.S. (2003) Global analysis of protein expression in yeast. *Nature*, **425**, 737–741.

122. D’Souza, A.R. and Minczuk, M. (2018) Mitochondrial transcription and translation: overview. *Essays Biochem.*, **62**, 309–320.

123. Suzuki, T., Yashiro, Y., Kikuchi, I., Ishigami, Y., Saito, H., Matsuzawa, I., Okada, S., Mito, M., Iwasaki, S., Ma, D. *et al.* (2020) Complete chemical structures of human mitochondrial tRNAs. *Nat. Commun.*, **11**, 4269.

124. Han, L. and Phizicky, E.M. (2018) A rationale for tRNA modification circuits in the anticodon loop. *RNA*, **24**, 1277–1284.

125. Barraud, P., Gato, A., Heiss, M., Catala, M., Kellner, S. and Tisne, C. (2019) Time-resolved NMR monitoring of tRNA maturation. *Nat. Commun.*, **10**, 3373.

126. Masterson, L.R., Cheng, C., Yu, T., Tonelli, M., Kornev, A., Taylor, S.S. and Veglia, G. (2010) Dynamics connect substrate recognition to catalysis in protein kinase A. *Nat. Chem. Biol.*, **6**, 821–828.