

FEATURE ARTICLE

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The purinosome, a multi-protein complex involved in the *de novo* biosynthesis of purines in humans

Cite this: *Chem. Commun.*, 2013, **49**, 4444

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Received 25th February 2013,
Accepted 2nd April 2013

DOI: 10.1039/c3cc41437j

www.rsc.org/chemcomm

Purine nucleotides are ubiquitous molecules that play vital roles in all kingdoms of life, not only as components of nucleic acids, but also participating in signaling and energy storage. Cellular pools of purines are maintained by the tight control of several complementary and sometimes competing processes including *de novo* biosynthesis, salvage and catabolism of nucleotides. While great strides have been made over the past sixty years in understanding the biosynthesis of purines, we are experiencing a renaissance in this field. In this feature article we discuss the most recent discoveries relating to purine biosynthesis, with particular emphasis upon the dynamic multi-protein complex called the purinosome. In particular we highlight advances made towards understanding the assembly, control and function of this protein complex and the attempts made to exploit this knowledge for drug discovery.

1. Introduction

Purines are essential molecules that serve a variety of functions and are utilized by all forms of life. Purines are components of a myriad of biomolecules that are vital for many cellular

processes such as genetic transfer (DNA), translation and transcription (RNA), energy storage and transfer (ATP and GTP), signaling (cyclic AMP and GMP) and also act as cofactors (NADH, NADPH and coenzyme A) in varied biochemical reactions.^{1–3} While much is known about the metabolism of purines,^{1–3} particularly from work on prokaryotic pathways, and despite several decades of research, new and surprising findings are regularly being reported.^{4–11} The recent discoveries have led to renewed interest in the study of purine biosynthesis, its regulation and the relationship this pathway and its intermediates have with other fundamental cellular processes.

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Jarrod B. French

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Cells access purine nucleotides through two separate mechanisms, *de novo* biosynthesis of inosine monophosphate (IMP) from 5-phosphoribosyl-1-pyrophosphate (PRPP, Fig. 1A) or through a salvage pathway that utilizes hypoxanthine, adenine or guanine (Fig. 1B). IMP is the point of convergence for the two pathways, as the purine nucleotides adenine and guanine are both separately synthesized from IMP in two additional steps (Fig. 1C). When the level of hypoxanthine is sufficient for cell growth (30 μM in one case¹²), purine nucleotides are synthesized preferentially by the salvage pathway. The regulatory capacity of the *de novo* pathway, however, is significant larger than that of the salvage pathway and has a much greater effect upon growth rate.^{12,13} This increased production of purines *via* the *de novo* pathway is intricately linked to cell growth and malignant transformation.^{14,15} The level of purine *de novo* biosynthesis varies widely amongst different human tissues. While liver and skeletal muscle tissue were reported to have relatively high rates of biosynthesis, the activity of the pathway enzymes in brain cells is only 25–30% of that in liver. Bone marrow cells, however, are believed to have limited capacity for *de novo* biosynthesis of purines.^{16–19} Those tissues that have inherently limited biosynthetic capacity would be expected to rely upon the purine salvage pathway or obtain purines from other tissues.

There are numerous disorders of purine and pyrimidine metabolism that affect humans and result, primarily, from genetic deficiencies of metabolic enzymes. There have been at least thirty different defects of purine and pyrimidine metabolism identified and seventeen of these are known to cause human disease.²⁰ One of the most common of these, stemming from a deficiency of the salvage enzyme hypoxanthine phosphoribosyl transferase (HPRT), leads to hyperuricemia, a condition where the level of uric acid in the blood is higher than the normal range (360 $\mu\text{mol L}^{-1}$ for women, 400 $\mu\text{mol L}^{-1}$ for men), and a disorder called Lesch–Nyhan disease.^{21,22} A similar hyperuricemic metabolic disorder results from mutation of phosphoribosylpyrophosphate synthetase (PRPPS). In this case, the overactivity of this enzyme that delivers purine precursors to the pathway causes the hyperuricemia.²² Hyperuricemia can

also lead to debilitating inflammatory arthritis, called gout.^{23,24} The hyperuricemia characteristic of gout is believed to stem from a variety of metabolic defects, including HPRT deficiency and accelerated purine biosynthesis or transport.^{24,25} Several disorders of purine metabolism can also lead to immunodeficiency. Multiple mutations on the gene encoding adenosine deaminase have been identified and are believed to be the major cause of severe combined immunodeficiency disease (SCID)²⁶. A deficiency of purine nucleoside phosphorylase can also lead to a clinical syndrome similar to SCID.²⁷ Mutations in the enzyme that catalyzes the eighth step in purine biosynthesis, adenylosuccinate lyase (ASL), can lead to mental retardation and seizures.²⁸ Similarly, a more recently discovered disorder, which results from a buildup of aminoimidazole carboxamideribonucleotide (AICAR), is caused by a defect in the transformylase activity of ATIC and can lead to impaired development or neurologic disease.²⁹

Despite the serious health implications that can result from defects, the redundancy in purine metabolism resulting from the presence of both a *de novo* and salvage pathway has precipitated the development of many clinically important drugs that target these pathways. In fact, greater than 20% of all approved oncology drugs are purine or pyrimidine antimetabolites.³⁰ Examples of these include thiopurines, used to treat acute lymphocytic leukemia,³¹ and purine analogues such as fludarabine, nelarabine, cladribine and others.³¹ In addition to drugs that target purine metabolic enzymes directly, antifolates have also been employed to disrupt purine biosynthesis by interfering with the biosynthesis of folate cofactors. These molecules, which include methotrexate and pemetrexed, are anticancer agents and are also employed as a treatment for autoimmune disorders including rheumatoid arthritis.^{32–34}

2. The *de novo* biosynthesis of purines

The *de novo* biosynthesis of purines is a highly conserved pathway that forms inosine monophosphate (IMP) from the



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Stephen J. Benkovic

*Dr Stephen J. Benkovic, an Evan Pugh Professor of Chemistry and Eberly Chair in Chemistry at the Pennsylvania State University, has focused on the assembly and kinetic characteristics of the protein machinery that is responsible for DNA replication; the importance of dynamic coupling of proximal and distal residues in the catalytic cycle of dihydrofolate reductase; the intracellular observation of *de novo* purine biosynthesis, and the collaborative development of novel sound based technologies for studying cellular communication. Benkovic was awarded the 2009 National Medal of Science and the National Academy of Science Award in Chemical Sciences in 2011.*

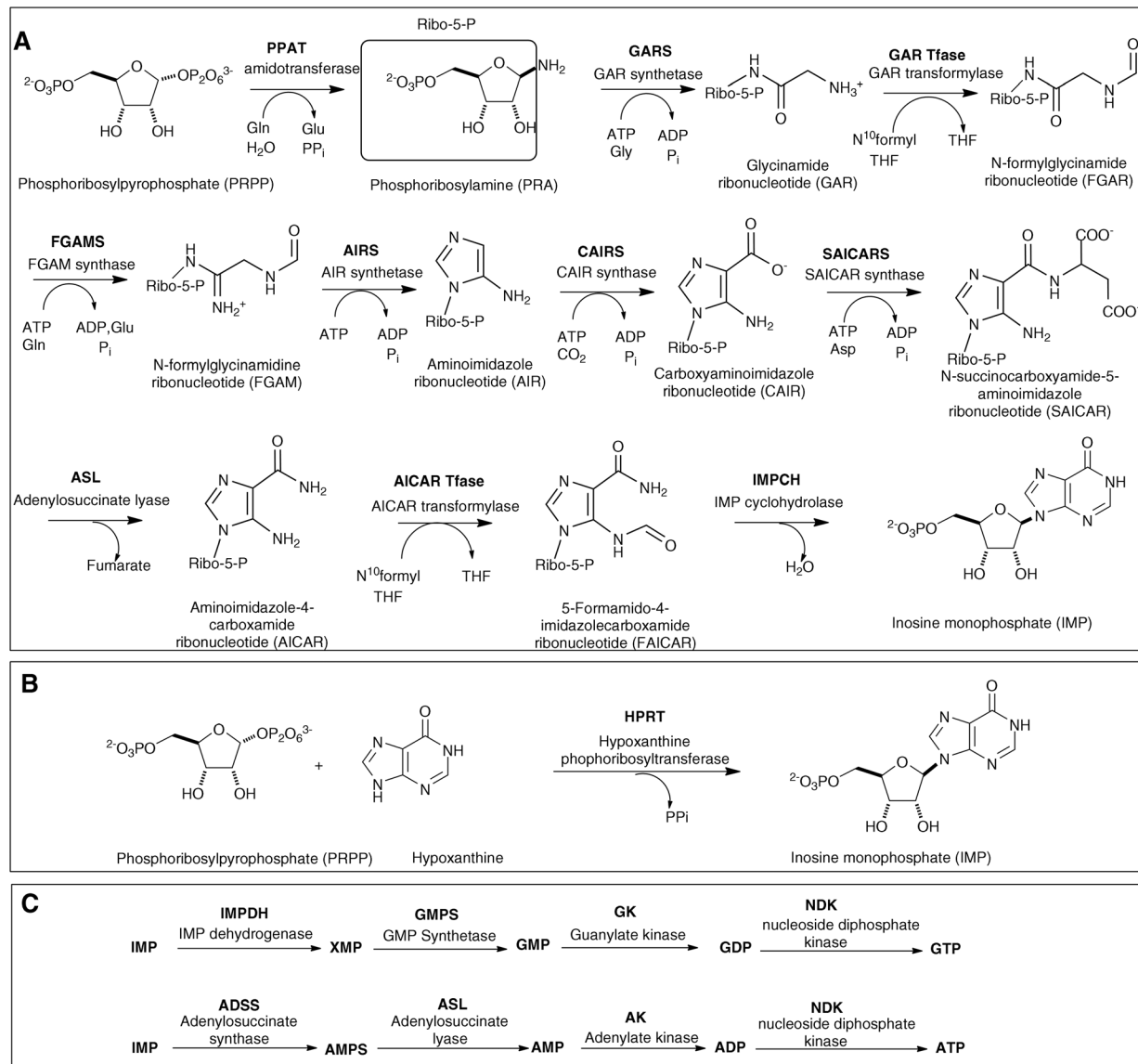


Fig. 1 Purine biosynthesis pathways. (A) The *de novo* purine biosynthetic pathway in human. The pathway contains 10 chemical steps and 6 human enzymes are involved. The trifunctional enzyme: TGART including GARS, GAR Tfase and AIRS, catalyzes the 2, 3 and 5 steps. The bifunctional enzyme PAICS contains CAIRS (6th step) and SAICARS (7th step). The last 2 steps are catalyzed by another bifunctional enzyme: ATIC, which is composed of AICAR transformylase and IMP cyclohydrolase. (B) The purine salvage pathway. HPRT transfers the hypoxanthine onto PRPP to form IMP. GMP or AMP can be synthesized by the same pathway from guanine or adenine using GPRT or APRT, respectively. (C) A simplified representation of the biosynthesis of GTP and ATP downstream of the *de novo* biosynthesis of IMP. GTP and ATP are synthesized from IMP via four steps in two separate pathways.

5-phosphoribosyl- α -1-pyrophosphate (PRPP) and glutamine with the help of several cofactors and four to six molecules of ATP. Seminal discoveries by Buchanan and coworkers starting in the 1950s^{35,36} established that ten enzymatic activities are required to carry out this conversion in chicken liver extracts. More recent studies, in particular the availability of complete genomes for a variety of organisms have allowed for the further characterization of this pathway in both prokaryotes and eukaryotes.

In bacteria, the ten chemical transformations of the pathway employ as many as twelve enzymatic activities and require additional ATP to complete. Humans, however, make use of several multifunctional proteins to carry out the complete transformation from PRPP to IMP.

The ten, highly conserved, chemical steps of *de novo* purine biosynthesis in humans are catalyzed by six enzymes (Fig. 1A). These enzymes include phosphoribosylpyrophosphate amidotransferase (PPAT), a trifunctional enzyme (TGART) which is composed of glycinamide ribonucleotide synthetase (GARS), GAR formyltransferase (GART) and aminoimidazole ribonucleotide synthetase (AIRS), formylglycinamide ribonucleotide synthetase (FGAMS), a bifunctional enzyme (PAICS) which is composed of

[‡] Buchanan *et al.* published a series of 35 reports from 1952 through 1971 in the *J. Biol. Chem.* entitled 'The Biosynthesis of Purines'. These seminal contributions laid the foundation for our current understanding of the enzymes and chemistry involved in purine biosynthesis.

carboxyaminoimidazole ribonucleotide synthase (CAIRS) and succinoaminoimidazolecarboxamide ribonucleotide synthetase (SAICARS), adenylosuccinate lyase (ASL) and a bifunctional enzyme (ATIC) which is composed of aminoimidazolecarboxamide ribonucleotide transformylase (AICART) and inosine monophosphate cyclohydrolase (IMPCH). The first step in this pathway, catalyzed by PPAT, is rate-limiting and is known to be under allosteric control.^{13,37} Two distinct nucleotide binding sites, both overlapping the PPAT active site, bind AMP, ADP, GMP and GDP and provide a mechanism of feedback control for the pathway.^{37,38}

Most of what we know about the structures and mechanisms of the enzymes of the *de novo* purine biosynthetic pathway come from studies carried out on the prokaryotic proteins.² The human enzymes range in size from approximately 57 kDa for PPAT to nearly 150 kDa for FGAMS and assume varied quaternary structures. While the structures of PPAT from both *E. coli* and *B. subtilis*^{38,39} reveal a tetrameric structure for this enzyme, work on the human enzyme suggests that there may be an equilibrium between the dimeric and tetrameric states.⁴⁰ X-ray crystal structures have been solved for each of the three domains of human TGART; however an intact, full-length structure still eludes researchers.^{41–43} Small angle X-ray scattering analysis has suggested that in solution this trifunctional enzyme assumes a relatively disordered, extended conformation.⁴² This observation may indicate that an additional protein, such as FGAMS or PPAT, may bind to TGART and interact with two or all three of the domains. The octameric crystal structure of human PAICS⁴⁴ and dimeric human ATIC^{45,46} have also been reported. While the structure of ATIC reveals an extensively intertwined obligate dimer, there is little structural evidence that channeling of the intermediate could traverse the approximately 50 Å between the active sites of this bifunctional enzyme. Conversely, the structure of PAICS reveals a potential tunnel system within the enzyme to accommodate channeling of the intermediate, CAIR.

3. The purinosome

For several decades there has been a great deal of speculation that the enzymes of the *de novo* purine biosynthetic pathway come together to form a functional complex. Several lines of evidence, accumulated over the years, support this hypothesis. One of the first indications of this phenomenon was the co-purification of enzymes from native sources.⁴⁷ Sequence information on the genes in the pathway provided additional support for complex formation. Not only does the mammalian pathway utilize several multifunctional enzymes, the trifunctional protein TGART catalyzes three non-sequential steps in the pathway (Fig. 1A). The most logical explanation for the retention of such a fusion protein in the genome is an evolutionary driving force that favors an important interaction between this protein and the protein that catalyzes the intermediate step, FGAMS.

Additional evidence that these proteins assemble into a complex comes in the form of kinetic arguments. An investigation of the degradation of the first intermediate in the pathway, PRA (Fig. 1A), suggests that this molecule is unstable and

possesses a short half life in solution.⁴⁸ In-depth kinetic analysis of the reaction catalyzed by the first two enzymes in the bacterial pathway argues in favor of complexation and presents a model that is not consistent with free diffusion of the reaction intermediate.⁴⁹ More recently, several proteomics initiatives have identified interactions between members of the *de novo* purine biosynthetic pathway. Havugimana *et al.* identified nearly 14 000 physical interactions in human cells using an integrative global proteomic profiling approach based upon chromatographic separation.⁵⁰ One of the interactions identified in this study was between PAICS and TGART. Zhang *et al.* employed a structure based prediction of protein–protein interactions to identify more than 300 000 interactions in humans with high confidence.⁵¹ Among these results were the prediction that FGAMS interacts with both TGART and PAICS.

3.1 Discovery of the purinosome

Despite the mounting evidence that the proteins in this pathway assemble in some functional manner, it wasn't until recently that direct evidence for such a complex was presented. Using fluorescence microscopy, the reversible compartmentalization of all six enzymes of the purine biosynthetic pathway was observed in the cytoplasm of several cancer cell lines.⁵ The phenomenon was demonstrated to be dependent on the level of purines in the media and was shown to be reversibly dynamic (Fig. 2). Immunofluorescence imaging of this protein assembly also demonstrated that the complex was observed in non-transfected cells with only endogenous levels of proteins present. A control protein that is involved in a related pathway, C1-Tetrahydrofolate

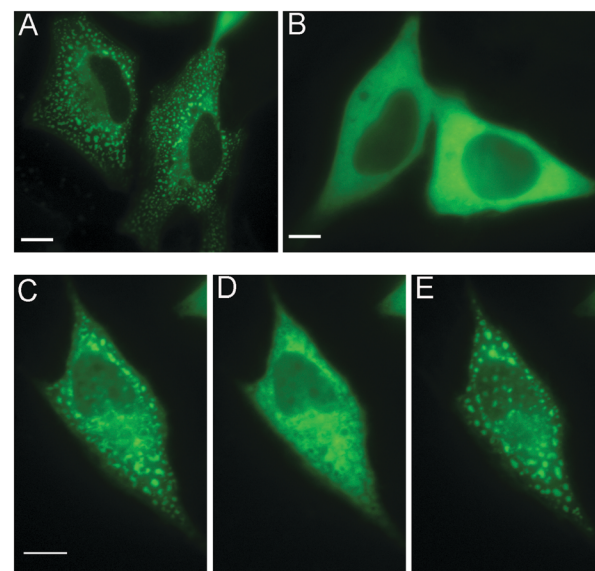


Fig. 2 Images of the purinosome. The pathway 4th enzyme FGAMS was fused with green fluorescent protein (GFP): FGAMS-GFP. (A) Purinosomes formed in HeLa cells transiently transfected with FGAMS-GFP in purine-depleted medium. (B) Diffuse fluorescence signal of FGAMS-GFP in HeLa cells in purine-rich medium. Reversible formation of clusters by FGAMS-GFP is shown in (C)–(E). (C) Purinosome formed when HeLa cells are cultured in purine-depleted medium. (D) Purinosomes disperse within 2 hours upon incubation with purine-rich medium. (E) Purinosomes reformed after returning to purine-depleted medium for 1 hour (Scale bar, 10 μm).

Synthase (C1-THF), did not cluster with the purine biosynthetic proteins. These observations provided strong evidence that a multi-enzyme complex, termed the purinosome, forms in cells under conditions of purine depletion.

While providing the first direct evidence for protein complexation in the *de novo* purine biosynthetic pathway, the purinosome is not a unique form of intracellular structure. In *E. coli*, proteinaceous microcompartments that sequester metabolic pathways involving unstable or toxic metabolites have recently been characterized.^{52,53} A screen of yeast strains yielded a total of 180 proteins that formed discrete physical structures upon nutrient starvation.⁵⁴ Similarly, 4 different filament-forming proteins were identified in screens of *S. cerevisiae*, *D. melanogaster*, and *E. coli*.^{55,56} In all of the above studies, the protein complexes are believed to provide temporal and spatial regulation as well as to provide protection from degradation of the protein components and pathway intermediates.

3.2 Purinosome control, interactions and assembly

A more detailed investigation of the purinosome led to several interesting findings. Several effectors of human protein kinases were found to trigger the association or dissociation of purinosomes.⁵⁷ The complex formation was promoted by the addition of one of several small molecule casein kinase II (CK2) specific inhibitors (Fig. 3A). The inhibitors 4,5,6,7-tetrabromo-1*H*-benzimidazole (TBI), 2-dimethylamino-4,5,6,7-tetrabromo-1*H*-benzimidazole (DMAT), tetrabromocinammic acid (TBCA), and 4,4',5,5',6,6'-hexahydroxydiphenic acid 2,2',6,6'-dilactone (ellagic acid) all stimulated the formation of purinosomes. This result was confirmed by silencing endogenous CK2 using siRNA. Strangely, treatment with the CK2 inhibitor 4,5,6,7-tetrabromobenzotriazole (TBB) resulted in the converse effect and caused the dissociation of purinosomes. While it is not known whether the pathway proteins themselves are phosphorylated by this kinase, it is clear that at least CK2 or a CK2-mediated signaling cascade participates in the reversible regulation of purinosome assembly.

A more recent study also implicated the Heat Shock Protein (Hsp) 90 and Hsp70 chaperone machinery in the formation of the purinosome.⁵⁸ Mass spectrometry analysis of an immunoprecipitation of FGAMS and associated proteins after treatment with a chemical cross-linking reagent yielded several chaperones and co-chaperones. Investigation of these proteins using fluorescence microscopy indicated that Hsp90 and Hsp70, as well as a set of co-chaperone proteins, co-localized dynamically with purinosomes. The abatement of Hsp70 or Hsp90 function through mutation or the decrease of co-chaperone expression using siRNA both led to decreases in purinosome content in the cell. These results suggested a possible means to disrupt purinosomes *via* inhibition of chaperone function. As expected, treatment of cells with several different Hsp90 or Hsp70 inhibitors: 17-*N*-allylamino-17-demethoxygeldamycin (17-AAG), 5-(2,4-dihydroxy-5-isopropyl-phenyl)-*N*-ethyl-4-[4-(morpholino-methyl)phenyl]isoxazole-3-carboxamide (NVP-AUY922), 2-phenylethylenesulfonamide (pifithrin- μ), 1-ethyl-2-[[3-ethyl-5-(3-methylbenzothiazolin-2-yliden)]-4-oxothiazolidin-2-ylidene)methyl]-pyridinium chloride (MKT-077) (Fig. 3B) led to disruption

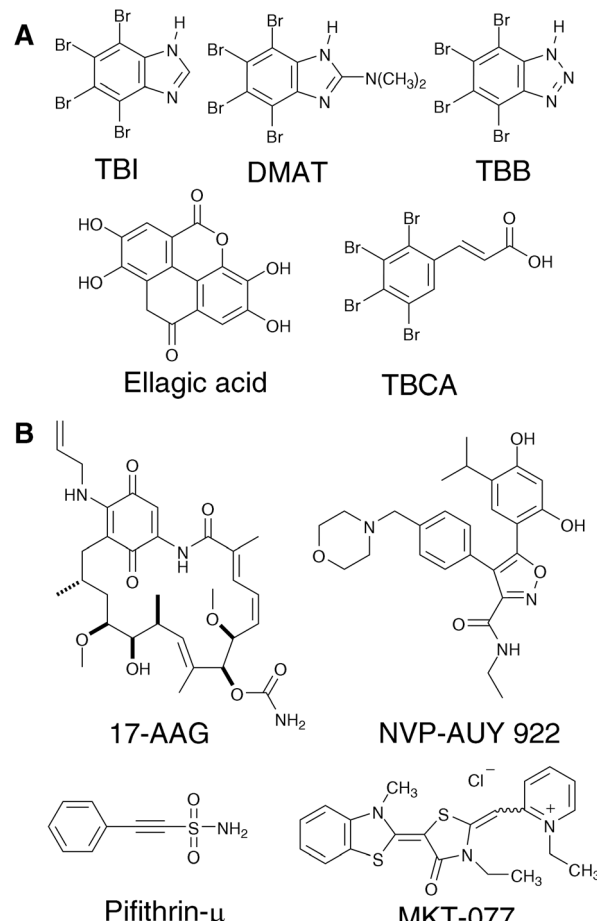


Fig. 3 Chemical structures of small molecules that affect the purinosome assembly inside cell. (A) Inhibitors of casein kinase II (CK2). (B) Inhibitors of Hsp90 and Hsp70 that have demonstrated to disrupt purinosomes.

of purinosomes. In fact, co-treatment of cancer cells with a combination of a known inhibitor of purine biosynthesis, methotrexate,³² and an Hsp90 inhibitor gave a synergistic cytotoxic effect. This work validates the purinosome as a viable target for novel cancer therapeutics and opens up the possibility for the development of new combination therapies that block purine biosynthesis by targeting both individual enzymes within the pathway and the formation of the purinosome complex.

The organization of purine biosynthetic proteins into purinosomes in cells suggests some form of intracellular spatial control of these proteins. Indeed, a microtubule-assisted mechanism for functional purinosome formation was proposed based upon results of fluorescent live-cell imaging studies.⁵⁹ This work determined that cytosolic purinosome clusters associated with microtubule filaments in the cell, but did not colocalize with the actin network. In addition, treatment with nocodazole, a known inhibitor of microtubule formation, not only caused the disruption of purinosomes but also led to a decrease in flux through the *de novo* purine biosynthetic pathway.

A further investigation into the organization of the purinosome employed a novel variation of the Tango assay⁶⁰ in order to examine potential protein–protein interactions. In this assay a transcription factor is tethered to the target protein with a

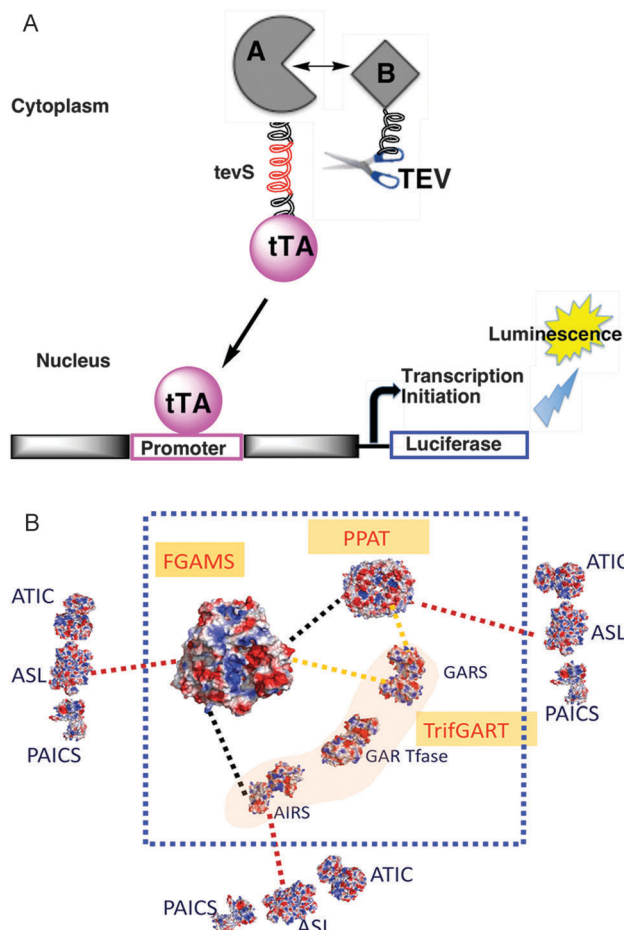


Fig. 4 A modified Tango assay was employed to detect and quantify protein–protein interactions among purinosome proteins. (A) Bait protein A is fused with the transcription factor, tTA linked by a peptide containing the recognition site (tevS) for a modified TEV protease. The prey protein, B, is linked to the modified TEV protease. When a protein–protein interaction occurs between proteins A and B, the TEV protease cleaves at tevS and releases tTA. This transcription factor enters the nucleus and turns on transcription of a luciferase gene. A representation of the interactions measured using this technique is provided in (B). The color of the lines represents the relative strength of the interaction detected between the two protein partners (from strongest to weakest – black, yellow, red). This work demonstrated that FGAMS, PPAT and TGART form a strong core complex that the other enzymes interact with more transiently.

linker that contains the cleavage site for a protease (Fig. 4). The protease, which is linked to a second protein, will cleave off the transcription factor only when the two proteins are in close proximity. Using this assay interactions between all six enzymes of the *de novo* purine biosynthetic pathway were detected and evidence for a core of the proteins that catalyze the first three steps was uncovered.⁶¹ PPAT, TGART and FGAMS exhibited the strongest degree of interaction and presumably nucleate formation of the overall complex. The remaining three enzymes, PAICS, ASL and ATIC interact individually with the core enzymes, but also were determined to interact with one another.

The modified Tango assay employed to investigate the purinosome provided details about the interactions between proteins, but was unable to answer any questions about the signals that drive the assembly or disassembly of this complex.

In addition to what has been reported about the involvement of kinases,⁵⁷ Verrier *et al.* reported that GPCRs can also regulate the assembly of the purinosome.⁶² The regulation of purinosome by small molecules targeting intracellular signaling proteins such as casein kinases suggests that the purinosome dynamics is under control of cell signaling.⁵⁷ To discover the upstream receptor signaling pathways, a methodology combining a label-free biosensor-enabled dynamic mass redistribution (DMR) assay with immunofluorescence imaging and molecular biology techniques was developed.⁶² The DMR assay is capable of translating an agonist-activated signal transduction process into a real-time whole cell phenotypic response, leading to a characteristic DMR signal.^{63,64} Using this label-free technique in conjunction with fluorescent live-cell imaging, Verrier *et al.* determined that GPCR signaling was involved in the regulation of purinosomes.⁶² Changes in DMR responses arising from the purinosome modulators DMAT and TBB were observed for several known adrenergic receptor agonists. Investigation with agents specific for α_2 -AR or β_2 -AR confirmed that purinosome formation was specifically associated with α_2 -AR activation (G_{zi} pathway, Fig. 5). The ability to promote purinosome formation was found to be a general mechanism for several endogenous G_{zi}-coupled receptors including purinergic P2Y receptors, LPA receptors, and prostaglandin receptors in HeLa cells. These findings suggest that, in addition to direct regulation of growth promoting genes in the nucleus, GPCR signaling also regulates purinosome assembly, an essential step in cell mitotic proliferation.⁶⁵

In addition to being a potential target for cancer therapeutics, the purinosome has also been implicated in AICAribosiduria and ASL deficiency.^{66,67} Cultured skin fibroblasts from patients with these diseases harbored mutations that destabilized purinosomes. These mutations cause structural instability in the enzymes which are known to be correlated with the disease phenotype. The results demonstrate that formation of purinosomes are dependent upon structurally intact ATIC and ASL, presumably in addition to the other members of the pathway.

3.3 Linking purine biosynthesis and the purinosome to other metabolic pathways

At two separate steps within the *de novo* purine biosynthetic pathway a formyl group is added to the intermediate using a tetrahydrofolate-derived cofactor (Fig. 1A). The necessity for these cofactors in purine biosynthesis is evident in the widespread use of anti-folates employed as anti-cancer agents to disrupt purine biosynthesis.⁶⁸ The supply of these one-carbon donors is maintained by a folate-dependent one-carbon metabolic network that facilitates the interconversion of tetrahydrofolates.^{69,70} Considering the importance of this pathway to purine biosynthesis, it was not surprising that one of its members, methenyltetrahydrofolate synthetase (MTHFS), was found to colocalize with purinosomes.⁷¹ Functional MTHFS was shown to be necessary for flux through the purine biosynthetic pathway, as mice deficient in MTHFS activity showed reduced levels of purine biosynthesis. In addition, the MTHFS protein appeared to require sumoylation in order to be trafficked to

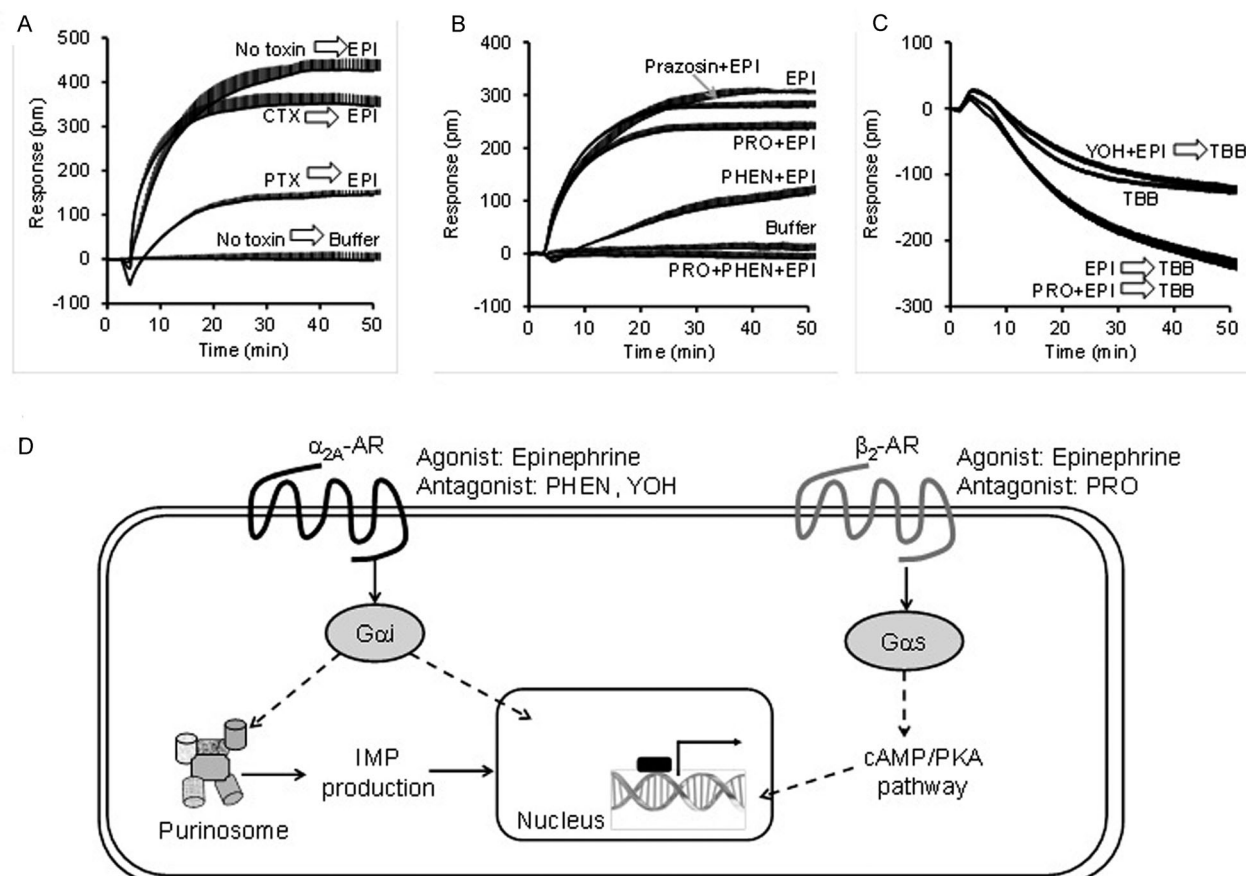


Fig. 5 GPCR signaling regulates purinosome formation. (A) The epinephrine (EPI) DMR contains contributions from both G αi and G αs pathways, as evidenced by the pathway deconvolution data using G protein-specific toxins. (B) Pharmacological profiling using co-stimulation of EPI with known AR antagonists showing that prazosin had little effect, propranolol slightly suppressed, phentolamine or yohimbine markedly suppressed, but the propranolol and phentolamine combination completely inhibited the EPI signal. Prazosin is a potent $\alpha 1$ -selective blocker, while propranolol (PRO) is a β -blocker, phentolamine (PHEN) and yohimbine (YOH) are two $\alpha 2$ -selective blockers. Together with quantitative real time-PCR, it was demonstrated that HeLa cells endogenously express functional G αs -coupled $\beta 2$ -AR and G αi -coupled $\alpha 2A$ -AR. (C) 4,5,6,7-Tetrabromobenzotriazole (TBB), which is known to dissociate the preassembled purinosome via a non-casein kinase II pathway, triggered a negative DMR. Pretreatment with EPI alone potentiated the TBB DMR, suggesting that EPI increases the number of purinosome complexes for TBB to disassemble. The co-pretreatment of EPI with YOH inhibited the EPI-induced potentiation, but propranolol had no impact, suggesting that EPI induces the formation of purinosome via the $\alpha 2A$ -AR. (D) Schematic representation of the expression and signaling of $\alpha 2A$ -AR and $\beta 2$ -AR in HeLa cells. Using both DMR and imaging, formation of the purinosome was determined to be promoted by the G αi -coupled $\alpha 2A$ -AR, but not G αs -coupled $\beta 2$ -AR in HeLa cells.

the purinosome. As well as linking the one-carbon metabolic network to the purinosome, these results provide additional evidence that signaling events, such as post-translational modifications, may be necessary for purinosome formation.

The growth of cancer cells is dependent upon metabolic reprogramming to increase glucose uptake and lactic acid fermentation, a phenomenon called the Warburg effect.^{72,73} One of the enzymes involved in glycolysis, pyruvate kinase, may serve as a bridge between this pathway and purine biosynthesis. This kinase, which plays a vital role in the growth and metabolic reprogramming that occurs in cancer cells, was shown to be allosterically controlled by the purine biosynthetic pathway intermediate SAICAR.⁸ These studies suggest that the accumulation of SAICAR under conditions of glucose depletion is related to an increase in *de novo* purine biosynthesis. This hypothesis is consistent with the observation that in cells where the PAICS activity was knocked-down, levels of SAICAR were not inducible, whereas cells that were deficient in the downstream enzyme ASL had constitutively high levels of SAICAR.

These results provide evidence for a possible link between purine biosynthesis and glycolysis. The starting material for purine biosynthesis, ribose-5-phosphate, is produced by the pentose phosphate pathway. The control of PKM2 by SAICAR-mediated allosteric stimulation may thus allow cells to adjust the generation of energy and metabolic flux in response to nutritional and energetic demands.

The *de novo* purine biosynthetic intermediate AICAR also plays a role in metabolic control, primarily through its effect on AMP kinase (AMPK). The AMPK system acts as a sensor of energy status *in vivo* and is activated by changes in AMP:ATP ratio.⁷⁴ AICAR, which acts as an AMP analog, is a known agonist for AMPK. AMPK acts as a regulatory switch for several intracellular processes including glucose uptake, glycolysis, insulin signaling, lipogenesis, cell cycle control and others.^{74–76} Activators of AMPK, such as metformin, are widely used to treat diabetes and related metabolic disorders.⁷⁴ Considering the cellular implications of AMPK activation it is clear that tight control over levels of activators, such as AICAR, be maintained

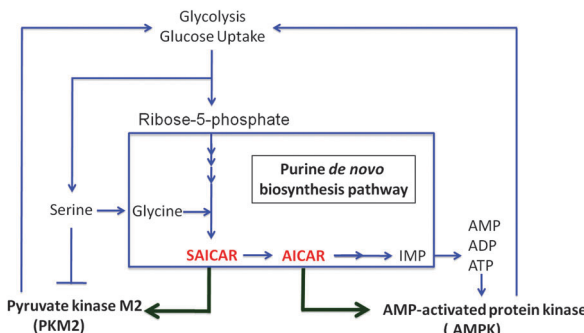


Fig. 6 Summary of some kinase-mediated interactions between the purine biosynthetic pathway, its intermediates and energy metabolism in the cell.

within the cell. Formation of the purinosome represents a potential means to minimize AICAR-mediated AMPK activation during *de novo* nucleotide biosynthesis. Coordination of the enzymes of purine biosynthesis into such an assembly could limit the accumulation of AICAR in order to prevent undesired metabolic consequences. In this fashion, the purinosome may provide a further level of metabolic control and could represent a novel means to control AMPK signaling *via* modulation of AICAR levels.

In a systematic characterization of cancer cell metabolite consumption and release, glycine was shown to be highly correlated with rates of cancer cell proliferation.⁷⁷ The more rapidly dividing cell lines showed a much higher dependence on the availability of glycine. This high degree of dependence of these cancer cells on glycine metabolism was demonstrated to be due, at least in part, to its role in supporting *de novo* purine biosynthesis. In addition, serine levels may also influence purine metabolism in cancer cells, both *via* its role as an allosteric activator of PKM2 and as a precursor to glycine.⁷⁸ The regulation of PKM2 by serine levels provides a feedback mechanism for control of various metabolic pathways within the cell. Fig. 6 summarizes some of the important relationships between intermediates in the purine biosynthetic pathway and several of the kinases responsible for metabolic control. While the significance of these metabolites in metabolic reprogramming and cancer progression are only beginning to be understood, it is clear that the regulation of purine levels in rapidly dividing cells is intimately tied to these processes and under very tight control. The fact that purinosomes are sensitive to purine levels as well as modulation by various kinases suggests that these protein assemblies may also play an important role in metabolic reprogramming.

4. Conclusions

The discovery of the purinosome has heralded a paradigm change in our understanding of nucleotide metabolism. The modulation of the assembly, dynamics and trafficking of this structure represent novel entry points for drug discovery efforts. With the validation of the purinosome as a means to improve the cytotoxicity of currently employed cancer therapeutics, the potential has arisen for an entirely new class of purine antimetabolites that specifically target the purinosome or those factors that govern its assembly. Future challenges involve

elucidating the role and extent of post-translational modifications on the *de novo* purine biosynthetic proteins and how these changes impact purinosome assembly. The interplay and impact of the purinosome with other metabolic pathways is only just beginning to be understood, and its role in metabolic reprogramming is an area that is bound to yield many exciting new insights. Similarly, the relationship between purinosome formation and the control of various important cellular processes such as the cell cycle, DNA replication, or glycolysis are yet to be fully elucidated. In addition, the level of local control over metabolite concentrations that is afforded by the purinosome is a topic that is still not well understood. These investigations will likely require new technologies that enable the measurement of local metabolite concentrations within cells at a very high degree of spatial and temporal resolution. Some imaging technologies that are likely to have a profound effect on how we study the purinosome and similar structures are already beginning to emerge. Super resolution imaging techniques such as Stochastic Optical Reconstruction Microscopy (STORM) and Stimulated Emission Depletion (STED) are examples of tools that will find great utility for the study of such phenomena. Finally, the characterization of the molecular details of the structure of the purinosome and its components is necessary to fully understand its function and to drive drug discovery efforts. Structural elucidation will certainly require a multidisciplinary approach that combines a myriad of biophysical techniques such as X-ray crystallography, NMR, mass spectrometry, super-resolution microscopy, label-free methods and computational approaches. In addition, as we begin to appreciate how important such dynamic multi-protein complexes are to myriad cellular processes, new tools will no doubt be needed to probe the functions, interactions and dynamics of such transient protein complexes *in vivo*.

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