## Research article **[Open Access](http://www.biomedcentral.com/info/about/charter/)**

# **Genome-wide screening for genes whose deletions confer sensitivity to mutagenic purine base analogs in yeast** Elena I Stepchenkova<sup>1</sup>, Stanislav G Kozmin<sup>1,2</sup>, Vladimir V Alenin<sup>1</sup> and Youri I Pavlov\*1,3

Address: 1Department of Genetics, Sankt-Petersburg State University, Sankt-Petersburg, 199034, Russia, 2Laboratory of Molecular Genetics, National Institute of Environmental Health Sciences, RTP, NC 27709, USA and 3Eppley Institute for Research in Cancer and Allied Diseases, the Department of Biochemistry and Molecular Biology, and the Department of Pathology and Microbiology, University of Nebraska Medical Center, Omaha, NE 68198, USA

Email: Elena I Stepchenkova - stepchenkova@yahoo.com; Stanislav G Kozmin - kozmin@niehs.nih.gov;

Vladimir V Alenin - alenin@alenin.usr.pu.ru; Youri I Pavlov\* - ypavlov@unmc.edu

\* Corresponding author

Published: 02 June 2005

*BMC Genetics* 2005, **6**:31 doi:10.1186/1471-2156-6-31

[This article is available from: http://www.biomedcentral.com/1471-2156/6/31](http://www.biomedcentral.com/1471-2156/6/31)

© 2005 Stepchenkov et al; licensee BioMed Central Ltd.

This is an Open Access article distributed under the terms of the Creative Commons Attribution License [\(http://creativecommons.org/licenses/by/2.0\)](http://creativecommons.org/licenses/by/2.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Received: 26 January 2005 Accepted: 02 June 2005

### **Abstract**

**Background:** *N*-hydroxylated base analogs, such as 6-hydroxylaminopurine (HAP) and 2-amino-6-hydroxylaminopurine (AHA), are strong mutagens in various organisms due to their ambiguous base-pairing properties. The systems protecting cells from HAP and related noncanonical purines in *Escherichia coli* include specialized deoxyribonucleoside triphosphatase RdgB, DNA repair endonuclease V, and a molybdenum cofactor-dependent system. Fewer HAP-detoxification systems have been identified in yeast *Saccharomyces cerevisiae* and other eukaryotes. Cellular systems protecting from AHA are unknown. In the present study, we performed a genome-wide search for genes whose deletions confer sensitivity to HAP and AHA in yeast.

**Results:** We screened the library of yeast deletion mutants for sensitivity to the toxic and mutagenic action of HAP and AHA. We identified novel genes involved in the genetic control of base analogs sensitivity, including genes controlling purine metabolism, cytoskeleton organization, and amino acid metabolism.

**Conclusion:** We developed a method for screening the yeast deletion library for sensitivity to the mutagenic and toxic action of base analogs and identified 16 novel genes controlling pathways of protection from HAP. Three of them also protect from AHA.

### **Background**

The accurate replication and repair of genetic material, which is a prerequisite for normal functioning of the eukaryotic genome and the prevention of cancer, relies on coordinated and faithful DNA synthesis. One important mechanism that ensures a high fidelity of DNA replication is a "cleansing" of the DNA precursor pool from deoxyribonucleoside triphosphates containing a modified base [1-4]. Such modified bases may have ambiguous basepairing properties that will result in a high mutagenic activity after their incorporation into DNA during replication. A classic example of the detoxification mechanism is the elimination of dUTP and 8-oxo-dGTP from the dNTP pool by the *E. coli* dUTPase and MutT proteins, respectively [1,5].



**Figure 1** Chemical structures of HAP and AHA and natural purine bases.

Purine analogs 6-hydroxyaminopurine (HAP) and 2 amino-HAP (AHA) are powerful mutagens in bacteria, yeast, and higher eukaryotes [6,7]. It has been suggested that HAP-deoxyriboside-triphosphate (dHAPTP) is a possible endogenous contaminant of nucleotide pools under peroxyl radical stress [8]. HAP and AHA closely resemble the natural purines, hypoxanthine and xanthine (Fig. 1), and therefore, could be exploited to investigate the mechanism preventing mutations that are caused by noncanonical purine nucleotides [9-11].

It was proposed that purine salvage enzymes convert base analogs to the corresponding deoxyribonucleoside triphosphates, which are misincorporated or misreplicated during DNA synthesis, resulting in induction of mutations [12,13]. HAP-induced mutagenesis in yeast is elevated in strains with defects in proofreading activity of replicative DNA polymerases [[14,](#page-9-0)15] and does not depend on excision, mutagenic recombination, and mismatch repair systems [\[14](#page-9-0)-16]. We have described several systems protecting cells from the mutagenic and inhibitory effects of HAP (see review [16]). One is the novel molybdenum cofactor-dependent system in *E. coli* [17]. It has yet to be determined if a similar system exists in higher eukaryotes. Another, versatile HAP-detoxification pathway relies upon the action of triphosphatase, Ham1p, which hydrolyze HAP-containing ribo- and deoxyribo-nucleotides to nucleoside monophosphates, and which prevent incorporation of base analog into DNA and RNA. We initially described the elevated sensitivity to HAP in yeast due to mutations in the *HAM1* gene [[18](#page-9-1)]. When we cloned and sequenced the *HAM1* gene, we found that it has homologs in many organisms, from bacteria to humans [13], and proposed that the gene might code for new triphosphatase [16]. Then, the crystal structure of the Ham1p homologue from a thermostable bacterium (protein Mj0226) was determined [19]. It was found that the Ham1p ternary structure has common features with MutT. Homologs of the yeast Ham1p from other organisms possessed triphosphatase activity on dITP, ITP, XTP, and dHAPTP substrates (Kozmin and Pavlov, unpublished; Burgis and Cunnigham, personal communication; and [19-21]).

There are additional, less thoroughly studied, factors modulating purine base analogues mutagenesis in yeast (see [16] for review). For example, *aah1* mutants are sensitive to HAP, suggesting that adenine deaminase Aah1p may deaminate HAP base to hypoxanthine [16].

In the present study, we carried out a genome-wide search for HAP and AHA sensitive mutants. The release of several complete sets of deletion mutants by the Yeast Deletion Project provides a powerful approach for different types of genome screens in yeast [22]. Haploid and diploid strains have already been used to detect new genes controlling sensitivity to different agents such as UV, ionising radiation, iron, and methyl methane sulfonate (MMS) [23-27], as well as spontaneous mutability [28]. This approach, when combined with other genomics approaches, helps to establish the biological functions of uncharacterized ORFs in yeast, many of which have human orthologs. This approach also allows us to decipher the network responses to endogenous and environmental stress [29]. The present study is the first systematic, genome-wide search for the mutations conferring sensitivity to mutagenic purine base analogs.

### **Results**

#### *Development of the screening method*

To develop a useful method for searching the yeast mutants sensitive to base analogs, we calibrated the experimental conditions using the wild-type strain, BY4742, and two previously described HAP-sensitive mutants, *ham1* and *aah1* (see [6,16]), created on BY4742 background. As shown in Fig. [2](#page-2-0) and described in Materials and Methods, yeast were grown in a 96-well microtiter plate and then transferred, using a multiprong replicator device, to YPD plates containing base analogs. An induction of the Canr mutants was monitored by replica-plating to the minimal-medium plates containing L-canavanine.

The results are presented in Fig. 3. The left panel of Fig. 3 shows the induction of canavanine-resistant mutants by HAP and AHA; and the right panel represents the survival

<span id="page-2-0"></span>

Scheme of the protocol for screening the yeast deletions library for base analog sensitivity and induced mutagenesis.

of the tested strains on YPD plates in the presence of base analogs. In the wild-type strain, as in the *ham1* and *aah1* mutants, 1–3 spontaneous canavanine-resistant colonies per spot arise in the absence of mutagen (Fig. 3A and 3B, left panel). In our experimental conditions, 1 µg/ml of HAP did not induce Canr mutants in the wild-type strain. A moderate induction of Canr clones (fewer than ten per spot) was observed at 10  $\mu$ g/ml of HAP and a very strong HAP-induced mutagenesis was observed in the wild-type strain at 100 µg/ml of HAP (Fig. 3A, left panel). For comparison, 100 µg/ml of AHA were only moderately mutagenic (compare Fig. 3A and Fig. 3E, left panel). Furthermore, both HAP and AHA did not affect the viability of the wild-type strain, even at the maximal concentration of 100 µg/ml (Fig. 3A and 3E, right panel).

In the *ham1* strain, 1 µg/ml of HAP induced Can<sup>r</sup> mutants with the similar frequency that was observed in the wildtype strain at 2 orders in magnitude higher concentration of analog, 100 µg/ml (compare Fig. 3A and 3B, left panel, phenotype of hypermutability, HM). The hypersensitivity of the *ham1* mutant to the toxic action of HAP was clearly detectable at 100 µg/ml of HAP (Fig. 3B, right panel). We will refer later to this phenotype as hypersensitivity, HS. Note that a reduction of the number of canavanine-resistant clones at 10 and 100 µg/ml of HAP, in comparison with 1 µg/ml of HAP observed in the *ham1* mutant (Fig. 3B, left panel, another manifestation of hypersensitivity phenotype, HS), is also due to a dramatic decrease of survival. When the *aah1* mutant was tested (Fig. 3B, second row, for HAP and Fig. 3F, first row, for AHA), the drop of viability was less severe than that for the *ham1* mutant (phenotype of elevated sensitivity, S). HAP-induced mutagenesis was detectable at a low dose of 1 µg/ml, but was much less compared to the *ham1* mutant (phenotype of elevated mutability, M). Mutagenesis was somewhat stronger at the dose 10  $\mu$ g/ml and was not seen at 100  $\mu$ g/ ml (another manifestation of elevated sensitivity, S). For the *aah1* mutant, AHA-induced mutagenesis was moderate at the dose of 10 µg/ml and very strong at a concentration of 100 µg/ml. The *aah1* mutant exemplified what we expect to observe for mutants moderately sensitive to both HAP and AHA. These data suggested that the procedure we devised for micro-titre plate format is effective for the detection of mutants with altered parameters of HAP and AHA sensitivity and mutability.

### *Screening of the deletion strains library*

We screened the yeast deletion strains library as described above. After an initial screening of 4,823 deletion strains,



### Results of the screening of the yeast de **Figure 3** letion library for elevated mutagenesis and sensitivity in micro-titer plates

**Results of the screening of the yeast deletion library for elevated mutagenesis and sensitivity in micro-titer plates**. Left panel – Mutagenesis on selective plates with canavanine. Right panel – The estimation of the number of colonyforming units on YPD medium.



#### <span id="page-4-0"></span>**Table 1: HAP-sensitivity of the mutants selected in our screening.**

†- HM – hypermutable, M – more mutable than wild-type, HS – hypersensitive, S – more sensitive than wild-type, see first paragraph od the Results Section for the explanation.

 $# - 25$  ug/ml

1 – 'DNA' – includes genes involved in the control of DNA precursor metabolism, purine salvage, DNA repair.

2 – 'Cell' – includes genes involved in cytoskeleton organization, cell walls and organelles.

 $3 -$  'Metabolic' – includes genes involved in general metabolic pathways.

 $\rm{*}$  – These mutants were also sensitive to the mutagenic or toxic effect of AHA (see Fig. 3).

° – small difference from the wild-type strain was reproducible and statistically significant by Wilcoxon-Mann-Whitney test.

43 mutants strains appeared to affect base analog-induced killing or mutagenesis. However, in subsequent testing of candidate strains in same type of plate tests, we have confirmed HAP-sensitivity of 16 mutants (Fig. 3, Table [1,](#page-4-0) column 4, where we refer to phenotypes of mutants according to abbreviations described in previous section). Next, we examined the mutability and survival of these 16 mutants in quantitative tests with HAP (see Materials and Methods). Based on the results of these two types of tests, we categorize HAP-sensitive strains in three groups, as shown in Table [1.](#page-4-0) Group I comprises, in addition to *ham1* and *aah1* strains, mutants *ade12* and *ade2*. These strains were hypersensitive to the mutagenic and lethal action of HAP in both types of tests (Fig. 3 and Table [1](#page-4-0), columns 4– 6). Strains of this group were sensitive to the low doses (1

and/or 10  $\mu$ g/ml) of HAP, in contrast to the wild-type strain. The *ade12* mutant was almost as mutable as the *ham1* strain, but the hypermutability could only be demonstrated in a quantitative test, due to poor plating efficiency (compare Fig 3B, row 3 and Table [1,](#page-4-0) column 5, row 3). The *ade2* mutant was as mutable as the *aah1* mutant (Table [1\)](#page-4-0). Deletion mutants of the first group showed variable degree of sensitivity to HAP-induced killing Table [1,](#page-4-0) column 6).

Eleven mutants fall into Group II. Mutants of this class were sensitive to the mutagenic effect of HAP, but their growth was not severely inhibited on HAP-containing medium. As a result, there is a smaller difference, in comparison to mutants of the group I, or no difference in the

number of Can<sup>r</sup> mutants induced by 10 and 100 µg/ml of HAP in spot-tests (Fig. 3C, Table [1,](#page-4-0) column 4). These strains produced some HAP-induced Canr mutants at 10 µg/ml of HAP, whereas the parental strain was non-mutable at this HAP concentration. We also did not detect any substantial drops of viability after growth in liquid media containing 25 µg/ml of HAP (Table [1](#page-4-0), column 6). Group II was not homogeneous in respect to HAP mutability and sensitivity. The six most sensitive mutants in the group are *vip1, vid27, ade1, ade5,7, ade6, ade8, ipk1* and *rim101* (see Table [1\)](#page-4-0). These mutants showed a decrease in the number of Canr mutants in a qualitative test when concentration of HAP increased ten-fold (up to  $100 \mu g/ml$ ), which is an indication of some cell death at very high doses of HAP (Fig. 3C). Mutants *yjl055w*, *ygr035c*, and *ade3* were more resistant to HAP, since the number of Canr colonies was similar at 10 and 100 µg/ml of HAP. As could be seen from the results of quantitative assay, mutant classification is quite conditional and there is substantial variation in responses between mutants of the Group II, but all of them were more mutable that the wild-type strain.

Finally, Group III includes mutants *she4, trp2*, and *yml013c-a;* which were sensitive to HAP-induced killing, but not to HAP-induced mutagenesis (Fig. 3D and Table [1,](#page-4-0) column 6). In quantitative tests the *yml013c-a* and *trp2* mutants were even less sensitive to the mutagenic action of HAP than the wild-type strain (Table [1\)](#page-4-0). The *she2* mutant was slightly more mutable that the wild-type strain only in quantitative test. The existence of such a type of mutants suggests that the toxic effect of HAP in yeast may be not only due to the induction of lethal mutations, but also due to inhibition of certain metabolic pathways.

In the present study, we have also characterized three AHA-sensitive mutants, *aah1*, *ade12*, and *yjl055w* (Fig. 3E and 3F). Remarkably, all of these mutants were also sensitive to HAP (Fig. 3C and Table [1](#page-4-0)). Two of those mutants, *aah1* and *yjl055w*, were AHA-hypermutable; whereas a*de12* strain did not mutate in the presence of AHA, but was sensitive to AHA-induced killing. In a quantitative test, 50 µg/ml of AHA did not inhibit survival of the wildtype strain, *aah1* and *yjl055w* strains; whereas survival of the *ade12* mutant was reduced to 50%. In the *aah1* and *yjl055w* strains, 50 µg/ml of AHA induced canavanineresistant mutants with the frequencies  $120 \times 10^{-7}$  and  $270$  $\times$  10<sup>-7</sup>, respectively, that was 4-9-folds higher than observed in the wild-type and *ade12* strains (30 × 10-7, in both strains).

### **Discussion**

In this study, we elaborated the technology for testing base analog-induced mutability and killing of thousands of yeast strains in microtiter format (see Fig. [2](#page-2-0)). We found

that the method is quite sensitive and reliable. Next, we screened the library of haploid yeast strains carrying deletions in all nonessential ORFs for the sensitivity to mutagenic base analogs HAP and AHA. We have found 16 novel HAP-sensitive mutants that fall into several groups, based on the sensitivity profiles (Fig. 3 and Table [1\)](#page-4-0). One group comprises the mutants that are HAP-hypermutable and grow poorly in the presence of HAP. Another class comprises strains with elevated HAP-mutability that grow normally on medium containing HAP. Finally, the third group includes the mutants sensitive to HAP-induced killing, but not to HAP-induced mutagenesis. We have also isolated three AHA-sensitive mutants. All of them were HAP sensitive as well. We summarized the properties of the genes found in our screening in Additional file 1.

One interesting result from our study is that none of the genes involved in the control of base analogs sensitivity except two genes, *YML013c-*a and *ade12*, were found in screenings for genes controlling sensitivity to the other types of mutagens, MMS, UV, and ionizing radiation [23- 27] or for elevated spontaneous mutagenesis [28]. It was previously reported that deletion of the *YML013c-a* and *ADE12* open reading frame specifically enhanced sensitivity to killing (as in case of HAP) induced by γ-radiation and bleomycin, but did not affect sensitivity to MMS, UV, and hydroxyurea [23,30]. It is known that there are overlaps of the sets of genes detected in genome screenings for the MMS-, UV- or ionizing radiation-sensitive strains [29]. Usually, those spectra include genes controlling DNA replication, recombination, and DNA repair. In our study, we did not find any of those genes. This is consistent with our earlier observation that the mutagenic action of HAP in *S. cerevisiae* does not depend on nucleotide excision repair, mutagenic repair, and mismatch repair [16]. A system to repair non-canonical purines and, probably HAP and AHA, in DNA has been recently characterized in *E. coli* [9,10]. It is possible that yeast *S. cerevisiae* lacks such a repair system.

Based on our data, we propose that, in yeast, the major base analogs protective mechanism is a control of the quality of DNA precursor pool that prevents incorporation of base analogs into DNA. This mechanism may work at several levels: transport of analogs into cells, detoxification of analogs by metabolic enzymes, maintenance of nucleotide pools, and fidelity control of DNA replication (Fig. [4](#page-8-0)). HAP and AHA are likely transported into the yeast cell by the same permeases, which are involved in transport of natural purines. One candidate is purine-cytosine permease, Fcy2p, a major purine (adenine, guanine, and hypoxanthine) and cytosine transporter in yeast [\[31](#page-9-2)]. According to our unpublished data, *fcy2* mutants are resistant to HAP. Thus, the active transport of HAP is the first critical step in the HAP mutagenic pathway. The next

step is a conversion of the base analog to the corresponding ribonucleoside monophosphate by enzymes of the purine salvage pathway. Previously we observed that the inactivation of the *APT1* gene, encoding adenine phosphoribosyl transferase, led to a severe decrease of the mutagenic effect of HAP [16], suggesting that this enzyme plays a key role in the biosynthesis of HAP-riboside-5' monophosphate (HAPMP). HAPMP then may be converted to the corresponding nucleoside triphosphate, which could be ambiguously incorporated into DNA by DNA polymerases and provoke replication errors in the subsequent replication cycles [13].

The mechanism preventing HAP- and AHA-induced toxicity most likely involves the conversion of base analogs to non-mutagenic metabolites by purine salvage enzymes. The main argument for this hypothesis is that HAP could be utilized by yeast cells as a sole purine source [13,[18](#page-9-1)[,32](#page-9-3)]. The first enzyme in this HAP and AHA detoxification pathway is adenine aminohydrolase, encoded by the *AAH1* gene. Aah1p from several microorganisms has been biochemically characterized (see [33]). Is has a broad substrate specificity and is capable of converting both adenine and its six-substituted analogs into hypoxanthine *in vitro*. According to our data, yeast Aah1p may convert HAP to hypoxanthine [12] and AHA to guanine *in vivo*, since the inactivation of the *AAH1* leads to a defect in this conversion that is readily detected by UV spectroscopy of yeast medium (unpublished observation). In the *aah1* mutant, the base analog intracellular concentration increases. We propose that this causes the elevation of base analog-induced mutagenesis (Fig. 3B and 3E, Table [1\)](#page-4-0).

Interestingly, in our screening we did not detect the *amd1* mutant, deficient in AMP aminohydrolase, that catalyze deamination of AMP to IMP [34]. We disrupted this gene by the *kanMX* cassette in several yeast strains and did not see any effect on HAP-induced mutagenesis anywhere. Thus, deamination of HAP at the mononucleotide level does not play an important role in HAP detoxification. This can be due to several reasons: the inability of the Amd1p to hydrolyse HAPMP, the minor role of the *AMD1* gene in yeast, or the short life-time of the HAPMP in yeast cells.

We have found that inactivation of adenilosuccinate synthase (ASS or Ade12p) encoded by the *ADE12* gene strongly enhanced HAP-induced mutagenesis and AHAinduced killing (see Fig. 3B and 3F, and Table [1](#page-4-0)). The primary function of this enzyme is the conversion of IMP to SAMP in the pathway of AMP biosynthesis *de novo* (Fig. [4\)](#page-8-0). We propose that the reason for this hypersensitivity is the dysregulation of purine biosynthesis, as follows. First, the blocking of this step of purine biosynthesis causes accumulation of IMP in the cell. The excess of IMP is probably toxic for the cell, since *ade12* mutants have a slow-growth phenotype that can be rescued by blocks of the earlier steps of the *de novo* purine biosynthesis pathway (Dr. A. M. Zekhnov {St-Petersburg State University}, personal communication). Accumulated IMP can be phosphorylated to ITP by nucleotide phosporylases. Thus, we propose that, in the *ade12* mutant, an excess of ITP may saturate Ham1p triphosphatase, an essential enzyme for the destruction of HAPTP and dHAPTP, which leads to increased HAP-sensitivity (see Fig. 3). The data obtained recently in bacteria are consistent with this hypothesis. It was shown that, *in vitro*, *E. coli* Ham1p homologues protein, RdgB, is a triphosphatase that acts to hydrolize noncanonical DNA precursors, dIPT and dXTP. The Ham1p protein was shown to possess a similar activity on dITP, dXTP, and dHAPTP substrates (Kozmin and Pavlov, unpublished; Burgis and Cunnigham, personal communication; and see refs. [19-21]). In *E. coli*, the *rdgB* mutation is synthetically lethal with *recA*. As proposed, absence of RdgB leads to a dramatic increase of hypoxanthine and xanthine in DNA. Accordingly, base excision repair of such modified bases occurring in opposite strands may generate double strand breaks that require the RecA function to be repaired. However, over-expression of adenilosuccinate synthase PurA (homolog of yeast Ade12p) suppresses this lethality [9]. This suggests a critical role of ASS in the regulation of intracellular concentration of genotoxic hypoxanthine-containing nucleotides.

We found that certain mutations affecting IMP biosynthesis *de novo* enhance HAP-induced mutagenesis (Table [1\)](#page-4-0). Seven of sixteen newly identified genes controlling HAP and AHA sensitivity are involved in AMP biosynthesis (see Additional file 1 and Fig. 3). It is known that the regulation of the AMP biosynthesis pathway by adenine is mediated by SAICAR, one of the precursors in adenine biosynthesis *de novo* [35]. The accumulation of certain purine biosynthesis by-products may play a role in the regulation of the nucleotide pool. A defect in endogenous purine biosynthesis probably alters nucleotide pools to favour dHAPTP mis-incorporation into DNA or leads to HAP toxicity. In respect to this hypothesis, it is important that there is a difference in the level of HAP-induced mutagenesis among the strains carrying mutations in the genes of AMP biosynthesis. The less sensitive mutant is *ade3* (Fig. 3 and Table [1](#page-4-0)). The *ADE3* does not directly control any steps of purine biosynthesis. It encodes the C1 tetrahydrofolate synthase that provides C1-tetrahydrofolate, an indispensable precursor for AMP, histidine, thymidylate, and methionine biosynthesis (see Fig. [4](#page-8-0)). In this respect it is of interest that *ade4, ade16* and *ade17* mutants, also defective in IMP biosynthesis, were not found in our screen and were not sensitive to HAP or AHA when constructed *de novo* and tested directly (data not

shown). The *ADE4* stands apart because PRPP, a substrate of the product of the *ADE4* gene, serves as a precursor for additional biosynthetic pathways. This prevents by-product accumulation and might be one of the explanations of lack of HAP sensitivity of the *ade4* mutants. The single *ade16* and *ade17* mutants also do not lead to byproduct accumulation because *ADE16* and *ADE17* are isozymes and the inactivation of one gene does not block the pathway.

We have found that 6 genes detected in our screening, *VIP1, VID27, IPK1, YGR035c, YML013c-a* and *SHE4*, are putatively involved in cell organization or genetically interact with cell-cycle control genes (see Additional file 1). This observation provides new perspectives on the mechanisms of base analogs-induced mutagenesis. It is possible that there is a specific structural route, mediated by cell cycle and cytoskeleton components, initiated by penetration of the analog inside the cell to its final target. We have also identified several hypothetical genes critical for the HAP and/or AHA resistance. This may be an initial clue to their functional significance.

Finally, we would like to mention that out of the 18 genes we found to be involved in HAP and AHA sensitivity control, 11 (60%) have orthologs in all groups of organisms, including mammals. Therefore, the results have relevance to higher eukaryotes and humans as well (see Additional file 1).

### **Conclusion**

We identified novel mutants sensitive to mutagenic and toxic effects of purine base analogs. AHA sensitivity was not previously described for three of the mutants identified in this study. The results reveal a complex control of base analogue mutagenesis by genes encoding the components of metabolic pathways and cytoskeleton.

### **Methods**

### *Yeast strains and media*

We have used a set of 4,823 *S. cerevisiae* mutants carrying deletions of all non-essential ORFs created in the haploid strain BY4742 (*MAT*α *his3*∆*leu2*∆*lys2*∆*ura3*∆). The information about the deletion strains set is available from the Yeast Deletion Project site: [ftp://ftp.resgen.com/pub/dele](ftp://ftp.resgen.com/pub/deletions/mat_alpha_041902.txt) [tions/mat\\_alpha\\_041902.txt](ftp://ftp.resgen.com/pub/deletions/mat_alpha_041902.txt). Deletion strains were constructed by replacement of the ORF's with the *kanMX4* cassette, which confers resistance to geneticin [36].

The standard yeast complete media (YPD) and minimal synthetic media (SD) were used [37]. Deletion strains were cultivated on YPD medium, supplemented with 200 µg/ml of geneticin. Sensitivity to HAP and AHA was examined on YPD media containing analogs in concentrations 100  $\mu$ g/ml, 10  $\mu$ g/ml and 1  $\mu$ g/ml. SD medium containing 40 µg/ml of L-canavanine was used for the selection of Canr mutants.

### *Base analogs sensitivity tests*

HAP was purchased from MP Biomedicals (Irvine, California, USA). AHA was custom- synthesized by Dr. I. Kuchuk at the Department of Chemistry of Indiana University (Bloomington, Indiana, USA) by the method of Janion [38]. Both chemicals were dissolved in DMSO (Fisher, USA) with mild heating. A single colony of each tested strain was inoculated into a well of 96-well microtitre plate containing 200 µl of liquid YPD medium (see Fig. [2](#page-2-0)). Microtiter plates were incubated for 2 days at 30°C with agitation, to reach a stationary phase (approximately 108 cells/ml). For mutagenesis assay, cells were then plated by a multiprong replicator device (approximately 5 µl of cell suspension per prong) to the YPD plates containing various concentrations of HAP or AHA, as shown on Fig. [2.](#page-2-0) After one day of incubation, the plates were replicaplated on SD minimal-medium plates containing L-canavanine. The plates were incubated 5 days and inspected for induction of Canr mutants.

For the survival test, cell cultures from the microplates were diluted in water in series of 96-well microplates, using a multichanel pipette (see Fig. [2](#page-2-0)). Diluted cells suspensions were plated to YPD plates containing HAP or AHA by a multiprong replicator device. After 2–3 days of incubation, the number of colonies was recorded. Strains that produced smaller colonies or a smaller number of colonies on the HAP-containing medium, relatively to wild-type strain, were classified as sensitive to killing.

### *Quantitative assay of the base analog-induced mutagenesis*

For each strain and each concentration of base analogs to be tested, we prepared six independent cultures by inoculating a single colony into 1 ml of liquid YPD medium with or without mutagens. After two days incubation in the roller drum, the mutant frequencies were determined by plating of appropriately diluted cell suspensions on minimal-medium SD plates supplemented with L-canavanine (to determine the number of canavanine-resistant cells per culture), and on YPD plates (to obtain the total number of cells per culture). Then the frequency of mutants was calculated as described [\[14](#page-9-0)]. Each experiment was repeated at least three times. We have used several doses of HAP and found that in this type of test the most reproducible results are obtained at dose 25 µg/ml. The statistical significance of differences between variants was estimated by Wilcoxon-Mann-Whitney nonparametric criterion.

<span id="page-8-0"></span>

### **Figure 4**

**Purine salvage and purine biosynthesis** *de novo* **in yeast** *Saccharomyces cerevisiae*. Intermediates of the purine biosynthesis *de novo* are designated in blue. The salvage pathway is presented in black. Genes, whose deletions lead to HAP and/or AHA sensitivity are highlighted by red boxes. The proposed conversions of HAP and AHA are represented in brackets below the adenine and guanine metabolites, respectively. Dashed arrows represent hypothetical pathways that were not demonstrated experimentally for a given substrate. Abbreviations:Purine biosynthesis *de novo*: PRPP – 5-phospho-ribosyl-1α-pyrophosphate, PRA – 5-phospho-β-D-ribosylamine, GAR – 5-phosphoribosylglycinamide, FGAR – 5'-phosphoribosyl-*N*-formyl glycinamide, FGAM – 5'-phosphoribosyl-*N*-formylglycinamidine, AIR – 5'-phosphoribosyl-5-aminoimidazole, CAIR – 5'-phosphoribosyl-5-aminoimidazole-4-carboxylate, SAICAR – 5'-phosphoribosyl-4-(*N*-succinocarboxamide)-5-aminoimidazole, AICAR – 5'-phosphoribosyl-4-carboxamide-5-aminoimidazole, FAICAR – 5'-phosphoribosyl-4-carboxamide-5-formamidoimidazole, SAMP – adenylosuccinate, 5,10-methylene-THF – 5,10-methylenetetrahydrofolate, 10-formyl-THF – 10-formyltetrahydrofolate. Salvage: Ade – adenine, AdeRib – adenosine, Hyp – hypoxanthine, Gua – guanine, Xan – xanthine.

### **Authors' contributions**

Youri Pavlov and Elena Stepchenkova designed this study. Elena Stepchenkova performed the experimental work and wrote the initial draft of the manuscript. Vladimir Alenin, Youri Pavlov, Stanislav Kozmin, and Elena Stepchenkova analyzed the data set and wrote the final version of the paper.

### **Additional material**

### **Additional File 1**

*Annotation of the genes whose deletion results in HAP and AHA sensitivity* Click here for file [\[http://www.biomedcentral.com/content/supplementary/1471-](http://www.biomedcentral.com/content/supplementary/1471-2156-6-31-S1.pdf) 2156-6-31-S1.pdf]

### **Acknowledgements**

We are grateful to Polina Shcherbakova for comments during this study and for critical reading of the manuscript. This work was supported in part, by NE DHHS LB506 grant #9934 for YIP, and by the CRDF grant provided by The Russian Ministry of Education # ST-012-0.

### **References**

- 1. Hochhauser SJ, Weiss B: *Escherichia coli* **[mutants deficient in](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=148458) [deoxyuridine triphosphatase.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=148458)** *J Bacteriol* 1978, **134(1):**157-166.
- 2. Michaels ML, Miller JH: **[The GO system protects organisms](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=1328155) [from the mutagenic effect of the spontaneous lesion 8](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=1328155) [hydroxyguanine \(7,8-dihydro-8-oxoguanine\).](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=1328155)** *J Bacteriol* 1992, **174(20):**6321-6325.
- 3. Grollman AP, Moriya M: **[Mutagenesis by 8-oxoguanine: an](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=8379000) [enemy within.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=8379000)** *Trends Genet* 1993, **9(7):**246-249.
- 4. Kamiya H: **[Mutagenic potentials of damaged nucleic acids pro](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=12527759)duced by reactive oxygen/nitrogen species: approaches using [synthetic oligonucleotides and nucleotides: survey and](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=12527759) [summary.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=12527759)** *Nucleic Acids Res* 2003, **31(2):**517-531.
- 5. Maki H, Sekiguchi M: **[MutT protein specifically hydrolyses a](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=1309939) [potent mutagenic substrate for DNA synthesis.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=1309939)** *Nature* 1992, **355(6357):**273-275.
- Pavlov YI, Noskov VN, Lange EK, Moiseeva EV, Pshenichnov MR, Khromov-Borisov NN: **The genetic activity of N6-hydroxyadenine and 2-amino-N6-hydroxyadenine in** *Escherichia coli, Salmonella typhimurium and Saccharomyces cerevisiae***[.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=1870608)** *Mutat Res* 1991, **253(1):**33-46.
- 7. de Serres FJ: **[The genetic toxicology of 2-amino-N6-hydroxya](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=1870610)[denine in eukaryotic organisms: significance for genetic risk](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=1870610) [assessment.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=1870610)** *Mutat Res* 1991, **253(1):**5-15.
- 8. Simandan T, Sun J, Dix TA: **[Oxidation of DNA bases, deoxyribo](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=9761719)[nucleosides and homopolymers by peroxyl radicals.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=9761719)** *Biochem J* 1998, **335(Pt 2):**233-240.
- 9. Burgis NE, Brucker JJ, Cunningham RP: **Repair system for noncanonical purines in** *Escherichia coli***[.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=12730170)** *J Bacteriol* 2003, **185(10):**3101-3110.
- 10. Bradshaw JS, Kuzminov A: **[RdgB acts to avoid chromosome](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=12791149)** [fragmentation in Escherichia coli.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=12791149) **48(6):**1711-1725.
- Kouzminova EA, Rotman E, Macomber L, Zhang J, Kuzminov A: **RecA-dependent mutants in** *Escherichia coli* **[reveal strategies](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=15531636) [to avoid chromosomal fragmentation.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=15531636)** *Proc Natl Acad Sci U S A* 2004, **101(46):**16262-16267.
- 12. Noskov V, Negishi K, Ono A, Matsuda A, Ono B, Hayatsu H: **[Muta](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=7516485)[genicity of 5-bromouracil and N6-hydroxyadenine studied by](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=7516485) [yeast oligonucleotide transformation assay.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=7516485)** *Mutat Res* 1994, **308(1):**43-51.
- 13. Noskov VN, Staak K, Shcherbakova PV, Kozmin SG, Negishi K, Ono BC, Hayatsu H, Pavlov YI: *HAM1***, the gene controlling 6-Nhydroxylaminopurine sensitivity and mutagenesis in the yeast** *Saccharomyces cerevisiae***[.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=8789257)** *Yeast* 1996, **12(1):**17-29.
- <span id="page-9-0"></span>14. Shcherbakova PV, Noskov VN, Pshenichnov MR, Pavlov YI: **Base analog 6-N-hydroxylaminopurine mutagenesis in the yeast** *Saccharomyces cerevisiae* **[is controlled by replicative DNA](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=8700180) [polymerases.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=8700180)** *Mutat Res* 1996, **369(1–2):**33-44.
- 15. Shcherbakova PV, Pavlov YI: **3'–>5' exonucleases of DNA polymerases epsilon and delta correct base analog induced DNA replication errors on opposite DNA strands in** *Saccharomyces cerevisiae***[.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=8849882)** *Genetics* 1996, **142(3):**717-726.
- 16. Kozmin SG, Schaaper RM, Shcherbakova PV, Kulikov VN, Noskov VN, Guetsova ML, Alenin VV, Rogozin IB, Makarova KS, Pavlov YI: **[Multiple antimutagenesis mechanisms affect mutagenic](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=9675240) activity and specificity of the base analog 6-N-hydroxylami[nopurine in bacteria and yeast.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=9675240)** *Mutat Res* 1998, **402(1–2):**41-50.
- 17. Kozmin SG, Pavlov YI, Dunn RL, Schaaper RM: **Hypersensitivity of** *Escherichia coli* ∆*(uvrB-bio)* **[mutants to 6-hydroxylaminopu](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=10852865)[rine and other base analogs is due to a defect in molybdenum](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=10852865) [cofactor biosynthesis.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=10852865)** *J Bacteriol* 2000, **182(12):**3361-3367.
- <span id="page-9-1"></span>18. Pavlov Iu I: **[Mutants of** *Saccharomyces cerevisiae* **[supersensi](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=3533720)[tive to the mutagenic effect of 6-N-hydroxylaminopurine\].](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=3533720)** *Genetika* 1986, **22(9):**2235-2243.
- 19. Hwang KY, Chung JH, Kim SH, Han YS, Cho Y: **Structure-based identification of a novel NTPase from** *Methanococcus jannaschii***[.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=10404228)** *Nat Struct Biol* 1999, **6(7):**691-696.
- 20. Chung JH, Park HY, Lee JH, Jang Y: **Identification of the dITP- and XTP-hydrolyzing protein from** *Escherichia coli***[.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=12297000)** *J Biochem Mol Biol* 2002, **35(4):**403-408.
- 21. Lin S, McLennan AG, Ying K, Wang Z, Gu S, Jin H, Wu C, Liu W, Yuan Y, Tang R, *et al.*: **Cloning, expression, and characterization of a human inosine triphosphate pyrophosphatase encoded by the** *ITPA* **[gene.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=11278832)** *J Biol Chem* 2001, **276(22):**18695-18701.
- 22. Giaever G, Chu AM, Ni L, Connelly C, Riles L, Veronneau S, Dow S, Lucau-Danila A, Anderson K, Andre B, *et al.*: **Functional profiling of the Saccharomyces cerevisiae [genome.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=12140549) 418(6896):**387-391.
- 23. Bennett CB, Lewis LK, Karthikeyan G, Lobachev KS, Jin YH, Sterling JF, Snipe JR, Resnick MA: **[Genes required for ionizing radiation](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=11726929) [resistance in yeast.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=11726929)** *Nat Genet* 2001, **29(4):**426-434.
- 24. Birrell GW, Giaever G, Chu AM, Davis RW, Brown JM: **A genomewide screen in** *Saccharomyces cerevisiae* **[for genes affecting](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=11606770) [UV radiation sensitivity.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=11606770)** *Proc Natl Acad Sci U S A* 2001, **98(22):**12608-12613. Epub 12001 Oct 12616.
- 25. Chang M, Bellaoui M, Boone C, Brown GW: **[A genome-wide](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=12482937) [screen for methyl methanesulfonate-sensitive mutants](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=12482937) reveals genes required for S phase progression in the pres[ence of DNA damage.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=12482937)** *Proc Natl Acad Sci U S A* 2002, **99(26):**16934-16939. Epub 12002 Dec 16913.
- 26. Game JC, Birrell GW, Brown JA, Shibata T, Baccari C, Chu AM, Williamson MS, Brown JM: **Use of a genome-wide approach to identify new genes that control resistance of** *Saccharomyces cerevisiae* **[to ionizing radiation.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=12816519)** *Radiat Res* 2003, **160(1):**14-24.
- 27. Davis-Kaplan SR, Ward DM, Shiflett SL, Kaplan J: **[Genome-wide](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=14594803) [analysis of iron-dependent growth reveals a novel yeast gene](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=14594803) [required for vacuolar acidification.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=14594803)** *J Biol Chem* 2004, **279(6):**4322-4329.
- 28. Huang ME, Rio AG, Nicolas A, Kolodner RD: **A genomewide screen in** *Saccharomyces cerevisiae* **[for genes that suppress](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=12972632) [the accumulation of mutations.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=12972632)** *Proc Natl Acad Sci U S A* 2003, **100(20):**11529-11534.
- 29. Begley TJ, Samson LD: **[Network responses to DNA damaging](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=15279801) [agents.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=15279801)** *DNA Repair (Amst)* 2004, **3(8–9):**1123-1132.
- 30. Westmoreland TJ, Marks JR, Olson JA Jr, Thompson EM, Resnick MA, Bennett CB: **Cell cycle progression in G1 and S phases is CCR4 dependent following ionizing radiation or replication stress in** *Saccharomyces cerevisiae***[.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=15075273)** *Eukaryot Cell* 2004, **3(2):**430-446.
- <span id="page-9-2"></span>31. Weber E, Rodriguez C, Chevallier MR, Jund R: **The purine-cytosine permease gene of** *Saccharomyces cerevisiae* **: primary structure and deduced protein sequence of the** *FCY2* **[gene](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=2191181) [product.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=2191181)** *Mol Microbiol* 1990, **4(4):**585-596.
- <span id="page-9-3"></span>Koz'min SG, Domkin VD, Zekhnov AM, Pavlov Iu I: **[Genetic control of metabolism of a mutagenic analog of 6-N-hydroxylaminopurine bases in** *Saccharomyces cerevisiae* **[yeasts\].](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=9273315)** *Genetika* 1997, **33(5):**591-598.
- 33. Hartenstein RC, Fridovich I: **[Adenine aminohydrolase. An inves](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=6017742)[tigation of specificity.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=6017742)** *J Biol Chem* 1967, **242(4):**740-746.
- 34. Sollitti P, Merkler DJ, Estupinan B, Schramm VL: **Yeast AMP deaminase. Catalytic activity in** *Schizosaccharomyces pombe* **and chromosomal location in** *Saccharomyces cerevisiae***[.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=8440738)** *J Biol Chem* 1993, **268(6):**4549-4555.
- 35. Rebora K, Desmoucelles C, Borne F, Pinson B, Daignan-Fornier B: **[Yeast AMP pathway genes respond to adenine through reg](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=11689683)[ulated synthesis of a metabolic intermediate.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=11689683)** *Mol Cell Biol* 2001, **21(23):**7901-7912.
- 36. Wach A, Brachat A, Pohlmann R, Philippsen P: **New heterologous modules for classical or PCR-based gene disruptions in** *Saccharomyces cerevisiae***[.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=7747518)** *Yeast* 1994, **10(13):**1793-1808.
- 37. Rose M, Winston F, Hieter P: **Methods in Yeast Genetics. A Laboratory Course Manual.** Cold Spring Harbor, NY: Cold Spring Harbor Laboratory Press; 1990.
- 38. Janion C: **[The synthesis and properties of N6-substituted 2](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=7092) [amino-purine derivatives.](http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?cmd=Retrieve&db=PubMed&dopt=Abstract&list_uids=7092)** *Acta Biochim Pol* 1976, **23(1):**57-68.

