In *Saccharomyces cerevisiae*, Expression of Arginine Catabolic Genes *CAR1* and *CAR2* in Response to Exogenous Nitrogen Availability Is Mediated by the Ume6 (CargRI)-Sin3 (CargRII)-Rpd3 (CargRIII) Complex

FRANCINE MESSENGUY,* FABIENNE VIERENDEELS, BART SCHERENS, AND EVELYNE DUBOIS

Institut de Recherches Microbiologiques J. M. Wiame and Laboratoire de Microbiologie de l'Universite´ Libre de Bruxelles, 1070 Brussels, Belgium

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The products of three genes named *CARGRI***,** *CARGRII***, and** *CARGRIII* **were shown to repress the expression of** *CAR1* **and** *CAR2* **genes, involved in arginine catabolism.** *CARGRI* **is identical to** *UME6* **and encodes a regulator of early meiotic genes. In this work we identify** *CARGRII* **as** *SIN3* **and** *CARGRIII* **as** *RPD3***. The associated gene products are components of a high-molecular-weight complex with histone deacetylase activity and are recruited by Ume6 to promoters containing a URS1 sequence. Sap30, another component of this complex, is also required to repress** *CAR1* **expression. This histone deacetylase complex prevents the synthesis of the two arginine catabolic enzymes, arginase (***CAR1***) and ornithine transaminase (***CAR2***), as long as exogenous nitrogen is available. Upon nitrogen depletion, repression at URS1 is released and Ume6 interacts with ArgRI and ArgRII, two proteins involved in arginine-dependent activation of** *CAR1* **and** *CAR2***, leading to high levels of the two catabolic enzymes despite a low cytosolic arginine pool. Our data also show that the deletion of the** *UME6* **gene impairs cell growth more strongly than the deletion of the** *SIN3* **or** *RPD3* **gene, especially in the** S**1278b background.**

The first mutations affecting the expression of the arginine catabolic genes *CAR1* and *CAR2*, encoding arginase and ornithine transaminase, respectively, were located in three unlinked genes, *ARGRI* (*ARG80*), *ARGRII* (*ARG81*), and *ARGRIII* (*ARG82*). The products of these genes were required for the induction of arginase and ornithine transaminase synthesis, and their loss impaired cell ability to use arginine and ornithine as nitrogen sources. The same proteins repressed the synthesis of five arginine anabolic enzymes when exogenous arginine was present in the growth medium (1, 42). Later, it was shown that the pleiotropic factor Mcm1 also participated, with the ArgR proteins, in the arginine-specific regulation (10, 28). The growth defect of an *argR* mutant allowed the selection of suppressor mutations falling into three complementation groups containing the *CARGRI* (*CAR80*), *CARGRII* (*CAR81*) and *CARGRIII* (*CAR82*) genes. Mutations in any of these genes led to overproduction of arginase and ornithine transaminase, even in an *argR* background (7, 9). The *CARGRI* gene was identical to *UME6*, a gene whose product is involved in controlling the expression of early meiotic genes (34, 41). Kadosh and Struhl (20) showed that repression by Ume6 at URS1, a sequence present in a wide variety of yeast promoters (24), involved recruitment of a Sin3-Rpd3 complex and targeted histone deacetylation.

Although *CAR1* and *CAR2* genes are coordinately induced by arginine and the ArgR-Mcm1 complex (29) and repressed by the three CargR proteins, only *CAR2* is induced by Dal82 and allophanate, the last intermediate of the allantoin-degrading pathway (17, 35), whereas *CAR1* is activated by Gln3 and shown that arginase derepression upon severe nitrogen starvation required the presence of arginine or a nonmetabolizable inducer, such as homoarginine (46). This work aimed at characterizing the *CARGRII* and *CARGRIII* genes and at determining their role in the response of *CAR1* and *CAR2* genes to exogenous nitrogen availability. In this paper, we identify *CARGRII* as *SIN3* and *CARGRIII* as *RPD3* and show that Ume6, Sin3, and Rpd3 proteins repress the expression of *CAR1* and *CAR2* genes, as long as nitrogen is available in the growth medium. We also identify a physiological interaction between Ume6 and ArgRI or ArgRII upon nitrogen depletion.

Nil1 through multiple GATAA sequences in the absence of optimal nitrogen sources (ammonia, glutamine) (11, 40). In addition *CAR1* expression is controlled not only by the quality of the nitrogen source (NCR) but also by the quantity of nitrogen available to the cell (8). Derepression of *CAR1* upon nitrogen deficiency requires the integrity of both the ArgR and Ume6 proteins and their target sequences (arginine boxes and URS1) but does not require the integrity of the GATAA sequences, the targets of Gln3 and Nil1 (11, 41). It was also

(A preliminary report of this work has been published as an abstract [12].)

MATERIALS AND METHODS

Strains and media. *Saccharomyces cerevisiae* strains 12T7c (*ura3*), 27061b (*ura3 trp1*), and 27029c (*ura3 leu2*) were derived from the wild-type strain S1278b. Strain BY4709 (*ura3*) was derived from the wild-type strain S288c (3). The in vivo-selected *cargRII* (11S52a) and *cargRIII* (02451c) mutants (7) were *ura3* recombinants from crosses between derivatives of wild-type strain S1278b and a *ura3* mutant strain isogenic to FL100. Strain HY (22) was used as the recipient strain for two-hybrid experiments.

Escherichia coli strain XL1-Blue (Stratagene) was used for plasmid amplification, and HB101 (Life Technologies) was used for plasmid recovery from yeast.

All yeast strains were grown on minimal medium containing 3% glucose, vitamins, mineral traces, and 0.02 M (NH₄)₂SO₄ (M.ammonia medium) (27).

^{*} Corresponding author. Mailing address: Institut de Recherches Microbiologiques J. M. Wiame, 1, Ave. E. Gryzon, 1070 Brussels, Belgium. Phone: 32.2.526.72.77. Fax: 32.2.526.72.73. E-mail: Fanarg @resulb.ulb.ac.be.

TABLE 1. Identification of *CARGRII* gene as *SIN3*

Expt	Strain	Plasmid	Genotype	Arginase speci- fic activity on M .ammonia ^a
	12T7c		ura3	$8*$
2	11S52a		ura3 cargRII	$40*$
\mathcal{R}	12T7cII		$ura3 \sin3$:: $kanMX4$	$49*$
4	12T7cHI		ura3 rpd3::kanMX4	$33*$
5	$11S52a \times 12T7cII$		ura3 cargRII \times ura3 $sin3$:: $kanMX4$	$46*$
6	$11S52a \times 12T7cIII$		ura3 cargRII \times ura3 rpd3::kanMX4	Q^*
	11S52a	$pFV33^b$	ura3 cargRII	16
8	11S52a	pFV36c	ura3 cargRII	40

^a Arginase specific activity was measured at 30°C after growth of the different strains on M.ammonia–25 μ g of uracil per ml where indicated (*). The specific activity is expressed in micromoles of urea produced per hour and per milligram of protein. The values are the means of two independent assays, and the standard

error was 10 to 15%. *^b* Corresponds to pRS416 *CEN6 ARS4 URA3 SIN3*. *^c* Corresponds to pRS416 *CEN6 ARS4 URA3 RPD3*.

Nitrogen starvation was achieved by filtering the cells grown on M.ammonia medium and cultivating them on fresh minimal medium without nitrogen for 2 h. The lithium acetate procedure (19) was used to transform the recipient yeast strains.

Construction of yeast deletant strains. The long flanking homology strategy was used to perform deletion of the following genes: *SIN3*, *RPD3*, *SAP30*, *UME6*, *GCN5*, *HDA1*, and *HOS2* (45). Long flanking homology replacement cassettes were synthesized using a two-step PCR, leading to the *kanMX4* cassette flanked by about 500 bp, corresponding to the promoter and terminator regions, respectively, of the target genes. The DNA fragments containing the different cassettes were used to transform yeast strains 12T7c (*ura3*) and BY4709 (*ura3*) on richmedium plates containing $200 \mu g$ of Geneticin/ml. The correct targeting of the deletions in G418^r transformants was verified by PCR, using whole cells as a source of DNA and appropriate primers. In the $\Sigma1278b$ background, the deletion of *UME6* had to be performed in the diploid strain obtained by crossing strain 27061b with 27029c. To construct strains with multiple deletions, we have used the gene disruption cassette *loxP-kanMX-loxP* (16). To eliminate the *kanMX* marker from the disrupted gene, the mutated strain was transformed with the *cre* expression plasmid pSH47, which carries the *URA3* marker gene and the *cre* gene under the control of the inducible *GAL1* promoter. Expression of the cre recombinase was induced by shifting cells from YPD (glucose) to YPG (galactose) medium for 2 h. The loss of the $kanMX$ cassette was detected by plating cells on YPD and replica plating the colonies onto YPD-G418. The *cre* expression plasmid was removed from the strains by streaking cells on plates containing 5-fluoroorotic acid to counterselect for the loss of the plasmid.

Construction of wild-type *UME6***,** *RPD3***, and** *SIN3* **wild-type cognate clones.** To clone these wild-type genes, the gap repair procedure was used (36). The *UME6*, *RPD3*, and *SIN3* open reading frame replacement cassettes were cloned into the vector pRS416 (39). The *kanMX4* modules were excised by restriction with *Eco*RI and *Bam*HI. The linearized plasmids were used to transform the standard FY1679 strain. Plasmids bearing the wild-type alleles pFV111 (*CEN6 ARS4 URA3 UME6*), pFV33 (*CEN6 ARS4 URA3 SIN3*) and pFV36 (*CEN6 ARS4 URA3 RPD3*) from URA⁺ transformants were isolated and amplified in *E. coli*. The presence of the genes of interest was verified by restriction analysis.

Fusion of *CAR2* **promoter to the** *lacZ* **gene.** A fragment of about 1,000 bp containing the *CAR2* promoter and its first two codons was produced by PCR using appropriate oligonucleotides based on *S. cerevisiae* genomic data and was extended with *Bam*HI restriction sites. This fragment was fused in frame to the *lacZ* coding sequence by insertion in the *Bam*HI site of plasmid pMC310 (pFL1 containing the *E. coli lacZ* gene inserted at the *Bam*HI site [5]); gift from M. Crabeel), yielding plasmid pFV118 (*CAR2-lacZ*). We determined the nucleotide sequence of the junction between the promoter and the *lacZ* gene to make sure that the fusion was in frame.

Construction of GBD and GAD fusions. The DNA-binding domain of the Gal4 activator, Gal4(1-147), is referred to as GBD, and its activation domain, Gal4(768-881), is referred to as GAD. GBD-ARGRI and GBD-ARGRII fusions were described previously (14). To produce the GAD-UME6 fusion, we used PCR to synthesize a *Bam*HI-*Bam*HI DNA fragment containing the *UME6* coding sequence (from nucleotide $+4$ to 2505). The primers used were based on the published *UME6* sequence (41), each flanked by a *Bam*HI restriction site. The *Bam*HI-*Bam*HI fragment was inserted into the *Bam*HI site of plasmid pACTII (13), yielding plasmid pFV124 (GAD-Ume6). In this GAD gene fusion, we determined the nucleotide sequence of the junction between the GAD-encoding region and the *UME6* gene to make sure that the fusions were in frame.

Enzyme assay. β-Galactosidase activity was assayed as described by Miller (31). Protein contents were determined by the Folin method (23).

Activity of arginase was assayed as described previously (30).

Measurement of amino acid pools. Differential extractions of cytosolic and vacuolar amino acid pools were performed as described by Ohsumi et al. (33).

The intracellular concentrations of glutamate, glutamine, arginine, ornithine, and lysine were determined by the PICOTag method. This system employs phenylisothiocyanate to rapidly and quantitavely derivatize both primary and secondary amino acids in a simple, one-step reaction. This derivatization is the first step of the well known Edman degradation. The stable phenylthiocarbamyl derivatives were easily separated by reverse-phase high-pressure liquid chromatography (2).

After peak identification, the amount of each amino acid was calculated by integration of the peak surface on the chromatogram and comparison to calibration curves established with standard amino acid solutions. These amounts, taking into account the different dilution factors and the dry weight of each sample, were used to establish pool concentrations. The dry weights were estimated by precisely measuring absorbance of the cells just before filtration using a conversion curve establishing the relation between the absorbance and the dry weight of the cells.

RESULTS

Identification of *CARGRII* **as** *SIN3* **and** *CARGRIII* **as** *RPD3.* Mutations in *CARGRII* and *CARGRIII* genes led to the derepression of two arginine catabolic genes, which was one of the phenotypes observed in *cargRI* (*ume6*) mutants. One role of Ume6 is to recruit to target promoters the Rpd3 histone deacetylase, through its interaction with Sin3. We have therefore tested the effect on *CAR1* and *CAR2* expression of deletions in the *SIN3* and *RPD3* genes. These deletions were created in strain 12T7c (*ura3*) by replacing the complete coding sequence of each gene by the *kanMX4* cassette, confering resistance to Geneticin. The levels of arginase, the product of the *CAR1* gene, from strains 12T7cII (*ura3 sin3*::*kanMX4*; Table 1, experiment 3) and 12T7cIII (*ura3 rpd3*::*kanMX4*; Table 1 experiment 4) were comparable to those of *cargRII* (11S52a; Table 1, experiment 2) and *cargRIII* (02451c; Table 2, experiment 2) mutants. The similar phenotypes produced by the *cargRII* and *cargRIII* mutations and the *sin3* and *rpd3* deletions prompted the question of their allelism. We crossed *cargRII* or *cargRIII* point mutant haploid strains to *sin3* or *rpd3* deletion mutants and sporulated the resulting diploids. The arginase levels presented in Table 1 showed that the *cargRII* mutant allele was incapable of complementing the *sin3* deletion allele (Table 1 experiment 5) but fully complemented the *rpd3* deletion allele (Table 1, experiment 6). In contrast, the *cargRIII* mutant allele complemented the *sin3* deletion allele (Table 2, experiment 6) and not the *rpd3* deletion allele (Table 2, experiment 5). These data suggest that *CARGRII* is identical to *SIN3*

TABLE 2. Identification of *CARGRIII* gene as *RPD3*

Expt	Strain	Plasmid	Genotype	Arginase speci- fic activity on M .ammonia ^a
	12T7c		ura3	$8*$
2	02451c		ura3 cargRIII	$50*$
3	12T7cIII		ura3 rpd3::kanMX4	$33*$
4	12T7cII		$ura3 \sin3$:: $kanMX4$	49*
5	$02451c \times 12T7cIII$		ura3 cargRIII \times ura3 rpd3::kanMX4	$30*$
6	$02451c \times 12T7cH$		ura3 cargRIII \times ura3 sin3::kanMX4	$12*$
7	02451c	pFV36c	ura3 cargRIII	13
8	02451c	$pFV33^b$	ura3 cargRIII	38

^a Arginase specific activity was measured at 30°C after growth of the different strains on M.ammonia–25 μ g of uracil per ml where indicated (*). The specific activity is expressed in micromoles of urea produced per hour per milligram of protein. The values are the means of two independent assays, and the standard error was 10 to 15%.

error was 10 to 15%. *^b* Corresponds to pRS416 *CEN6 ARS4 URA3 SIN3*. *^c* Corresponds to pRS416 *CEN6 ARS4 URA3 RPD3*.

TABLE 3. Effect of mutations impairing histone acetylation and deacetylation activities on *CAR1* expression

Strain	Genotype	Arginase specific activity on M.ammonia ^a
12T7c	ura3	8
12T7cII	$ura3 \, sin3$:: $kanMX4$	49
12T7cIII	ura3 rpd3::kanMX4	33
12T7cIV	ura3 sap30::kanMX4	27
12T7cV	ura3hda1::kanMX4	5
12T7cVI	$ura3$ hos2:: $kanMX4$	4
12T7cVII	ura3 gcn5::kanMX4	9

^a Arginase specific activity was measured at 30°C after growth of the different strains on M.ammonia-25 µg of uracil per ml. The specific activity is expressed in micromoles of urea produced per hour per milligram of protein. The values are the means of two independent assays, and the standard error was 10 to 15%.

and that *CARGRIII* is identical to *RPD3*. We confirmed these results by analysis of the tetrads issued from the sporulation of the *cargRII-* $\Delta sin3$ diploid. All the spores from 10 tetrads had derepressed arginase levels on M.ammonia medium. This analysis could not be performed with the *cargRIII-* Δr *pd3* diploid, which did not sporulate. It was reported that homozygous *sin3* or *rpd3* diploid strains were sporulation defective (43, 44). Our *cargRII* point mutant is thus not impaired in sporulation, in contrast to our *cargRIII* point mutant. The *SIN3* and *RPD3* genes were cloned by gap repair (see Materials and Methods) and introduced into *cargRII* (11S52a) and *cargRIII* (02451c) mutant strains. The arginase-specific activities in the *cargRII* strain transformed with plasmid pFV33 (pRS416 *ARS CEN URA3 SIN3*) (Table 1, experiment 7) and the *cargRIII* strain transformed with pFV36 (pRS416 *ARS CEN URA3 RPD3*) (Table 2, experiment 7) were comparable to that of the wildtype strain, as expected.

Sap30, but not Hda1, Hos2, or Gcn5, controls *CAR1* **expression.** In *S. cerevisiae* as in humans, histone acetylation and deacetylation are catalyzed by structurally distinct multisubunit complexes. In yeast, Gcn5, present in Ada and SAGA (Spt/ Ada) complexes (4, 15, 26), and TAF145/130 (the yeast equivalent of mammalian TAF250), present in TFIID (32), were shown to have histone acetylation activity. Hda1, a component of histone deacetylase A, has identity to Rpd3, Hos1, Hos2, and Hos3, and it was shown that Hda1 and Rpd3 are members of distinct histone deacetylase complexes (38). Sap30 was recently shown to be part of the Rpd3-Sin3 complex in *S. cerevisiae* (47). To test the role of these different proteins involved in histone acetylation and deacetylation in *CAR1* expression, we have created strains with deletions of the *HDA1*, *HOS2*, *SAP30*, and *GCN5* genes by insertion of the *kanMX4* cassette (see Materials and Methods). As shown in Table 3, the arginase levels on minimal medium in strains with the *HDA1* (12T7cV), *HOS2* (12T7cVI), or *GCN5* (12T7cVII) gene deleted were comparable to the level in the wild-type strain, whereas deletion of the *SAP30* gene (strain 12T7cIV) led to a derepression of *CAR1* expression, as in *SIN3*- and *RPD3*-deleted strains. It is noteworthy that the activation of *CAR1* in a *gcn5*::*kanMX4* strain in response to the presence of arginine or a poor nitrogen source was not impaired (data not shown). Only the Rpd3-Sin3-Sap30 complex seems to regulate the expression of the *CAR1* gene, probably by deacetylation of histones. In contrast, the histone acetyltransferase activity of Gcn5 or TAF145/130 (according to global analysis by Holstege et al. [18]) does not seem to be required for *CAR1* expression.

Effect of deletions in *UME6***,** *SIN3***, and** *RPD3* **genes on cellular growth in the** S**1278b background.** Deletions of *SIN3* and *RPD3* genes from strain 12T7c, which is a *ura3* derivative of Σ 1278b, were performed. These two strains resulting from the deletions (12T7cII and 12T7cIII) were affected in their growth on minimal medium (Fig. 1A). In the same background, we could not obtain a viable *ume6*::*kanMX4* strain. We thus performed the *ume6* deletion in the diploid strain 27061b $(ura3^{trp1}) \times 27029c (ura3 *leu2*)$ by insertion of the *kanMX4* cassette and obtained viable diploids resistant to Geneticin (see Materials and Methods). After sporulation of these diploids, tetrad analysis revealed that only two spores were viable (Fig. 1C) and that the viable spores were Geneticin sensitive. The *UME6* gene product is thus essential in the Σ 1278b background, whereas it only reduces the growth rate in other backgrounds. We have indeed created *UME6*, *SIN3*, and *RPD3* deletions in strain BY4709 (*ura3*; gift from J. Boeke), and, as shown in Fig. 1B, deletion of the three genes also affects the cellular growth rate in this background, especially for the strain with the *UME6* gene deleted.

CAR1 **expression in strains bearing different combinations of mutations impairing histone deacetylation.** To test whether simultaneous deletion of *RPD3*, *SIN3*, and *UME6* genes had a cumulative effect on *CAR1* expression, we have created in strain BY4709 single, double, and triple deletions. Each gene was deleted by the *kanMX4* cassette flanked by *lox* sequences, which allowed the removal of the *kanMX4* cassette by recombination using the cre recombinase (16) (see Materials and Methods). As shown in Table 4, the arginase level was higher in the *ume6*::*kanMX4* strain (BY4709I) than in the *rpd3*::*kanMX4* strain (BY4709III), in the *sin3*::*kanMX4* strain (BY4709II), or in the *rpd3*::*kanMX4 sin3*::*kanMX4* strain (BY4709II-III). The derepression of arginase synthesis resulting from the *UME6* deletion was not increased by additional deletion of *SIN3* (BY4709I-II), *RPD3* (BY4709I-III), or *SIN3* plus *RPD3* (BY4709I-II-III). These results confirm the interdependence of these three proteins to repress a gene containing a URS1 sequence. The deletion of Ume6, which is the DNA binding protein, abolishes the repression of *CAR1* expression, whereas the deletion of Sin3 and/or Rpd3, two components of the histone deacetylase complex, relieves only partially the repression. This suggests that other deacetylases and deacetylaseassociated proteins could fulfil the Rpd3 or Sin3 functions.

The Ume6-Sin3-Rpd3 complex represses arginine catabolic genes in response to exogenous nitrogen availability. We have previously shown that the response of the *CAR1* gene to nitrogen starvation is impaired in a *ume6* mutant strain and in a strain in which the Ume6 target sequence, namely, URS1, was deleted from the *CAR1* promoter (11, 41). Since Ume6 works in conjunction with Sin3 and Rpd3, we tested the role of the last two proteins in the control of *CAR1* by nitrogen availability. The results showed that, upon nitrogen starvation, the arginase levels in strains with *sin3*::*kanMX4* (BY4709II) or *rpd3*::*kanMX4* (BY4709III) deleted were comparable to that in the wild-type strain (Table 5), indicating that, besides Ume6, Sin3 and Rpd3 were required for mediating repression of *CAR1* expression as long as nitrogen was available. Since *cargRI* (*ume6*), *cargRII* (*sin3*), and *cargRIII* (*rpd3*) mutations also led to derepression of the *CAR2* gene (7, 9), we analyzed the response of the *CAR2* promoter to nitrogen starvation in these strains with deletions. Since the basal level of the *CAR2* gene product (ornithine transaminase) was very low, we measured *CAR2* expression using a *CAR2-lacZ* fusion (pFV118). The deletion of the *UME6*, *SIN3*, or *RPD3* gene led to a strong increase of β -galactosidase synthesis, and, similar to what was found for *CAR1*, *CAR2* derepression was significantly higher in the *ume6*-deleted strain. When the three strains with deletions were starved completely for exogenous nitrogen, fur-

FIG. 1. Effect of deletions in *SIN3*, *RPD3*, and *UME6* genes on cell growth. (A and B) Tenfold serial dilutions of cells were plated and incubated at 30°C for 3 days on M.ammonia–25 mg of uracil. Strains 12T7c (*ura3*), 12T7cII (*ura3 sin3*::*kanMX4*), and 12T7cIII (*ura3 rpd3*::*kanMX4*) are isogenic to strain S1278b (A). Strains BY4709 (*ura3*), BY4709II (*ura3 sin3*::*kanMX4*), BY4709III (*ura3 rpd3*::*kanMX4*), and BY4709I (*ura3 ume6*::*kanMX4*) are isogenic to S288c (B). (C) Tetrad analysis of the S1278b isogenic diploid strain 27061b (*ura3 trp1*) 3 27029c (*ura3 leu2*) in which one copy of the *UME6* gene was replaced by the *kanMX4* cassette. Spores were isolated on YPD medium. WT, wild type.

ther derepression of *CAR2* was abolished in the *ume6* strain and strongly reduced in the *sin3* and *rpd3* strains.

Induction of arginase upon nitrogen starvation does not result from a burst of arginine in the cytoplasm. It was previously reported that starvation of an arginine auxotroph for both arginine and nitrogen did not result in the synthesis of arginase unless homoarginine, a nonmetabolizable inducer, was added, suggesting that production of arginase under conditions of nitrogen starvation was the result of induction and was contingent upon the presence of the inducer arginine in the amino acid pools of the cells (46). Later it was proposed that the internal induction of arginase in cells deprived of any

TABLE 4. *CAR1* expression in strains bearing single, double, and triple deletions of *UME6*, *SIN3*, and *RPD3*

Strain	Genotype	Arginase speci- fic activity on M .ammonia a	
BY4709	ura3	7	
BY4709I	ura3 ume6::kanMX4	66	
BY4709H	ura3 sin3::kanMX4	38	
BY4709HI	ura3 rpd3::kanMX4	34	
BY4709I-II	$ura3$ ume6:: $kanMX4$ sin3::lox	63	
BY4709I-III	ura3 ume6::kanMX4 rpd3::lox	60	
BY4709II-III	ura3 sin3::kanMX4 rpd3::lox	41	
BY4709I-II-III	ura3 ume6::kanMX4 sin3::lox rpd3::lox	60	

^a Arginase specific activity was measured at 30°C after growth of the different strains on M.ammonia– 25μ g of uracil per ml. The specific activity is expressed in micromoles of urea produced per hour per milligram of protein. The values are the means of two independent assays, and the standard error was 10 to 15%.

nitrogen source resulted from the release of arginine from the vacuole (6). The data shown in Fig. 2 indicate that maximal arginase derepression was reached after 40 min of nitrogen starvation (Fig. 2A). To test whether this rapid increase of arginase level was a consequence of arginine accumulation in the cytosol, we have measured the evolution of amino acid pools in the vacuole and the cytoplasm after a shift from M.ammonia medium to a medium devoid of nitrogen. Using the standard $Cu2^{2+}$ method (33), we have determined the

TABLE 5. Effect of mutations in *UME6*, *SIN3*, and *RPD3* on the starvation response of *CAR1* and *CAR2* genes

			Expression of:	
Strain	Genotype	Median ^a	$CARI^c$	$CAR2$ -lac Z^b
BY4709	ura3	M.ammonia		176
		$-N^d$	60	1,184
BY4709I	ura3 ume6::kanMX4	M.ammonia	66	1,644
		$-N$	76	1,743
BY4709H	$ura3 \, sin3$:: $kanMX4$	M.ammonia	38	900
		$-N$	63	1,165
BY4709III	ura3 rpd3::kanMX4	M.ammonia	34	653
		$-N$	65	1,045

a When no plasmid was present in the strains, 25 μ g of uracil was added. *b* β -Galactosidase activity was measured in extracts from different strains

transformed with plasmid pFV118 ($CAR2\text{-}lacZ$). The β -galactosidase specific activity is expressed in nanomoles of *o*-nitrophenyl-ß-D-galactopyranoside hydro-lyzed per minute per milligram of protein.

^c Arginase-specific activity is expressed in micromoles of urea produced per hour per milligram of protein. The standard error was 10 to 15%.

^d -N, medium with no nitrogen source (see Materials and Methods).

FIG. 2. Determination of arginase-specific activity and cytosolic and vacuolar glutamate, glutamine, arginine, ornithine, and lysine pools upon nitrogen starvation.
Arginase-specific activity was measured in extracts from nitrogen source (times 20, 40, 60, and 120 min) (A). The specific activity is expressed in micromoles of urea produced per hour per milligram of protein. (B to F) Solid diamonds, vacuolar amino acid pool; open circles, cytosolic amino acid pool. Amino acid intracellular concentrations are expressed in nanomoles per milligram of dry weight.

TABLE 6. Interaction between Ume6 and ArgRI or ArgRII under nitrogen starvation conditions*^a*

Protein	Medium	β -Galactosidase specific activity ^{<i>b</i>} for:	
		GAD	GAD-Ume ₆
GBD	M.ammonia	$<$ 1	$<$ 1
	$-N^c$	$<$ 1	$<$ 1
GBD-ArgRI	M.ammonia		\overline{c}
	$-N$		21
GBD-ArgRII	M.ammonia	33	37
	$-N$	35	162

^a The full-length Ume6 was fused in frame with the activation domain of Gal4 (GAD-Ume6), and full-length ArgRI and ArgRII were fused in frame with the

^{*b*} Transcription activation of the *lacZ* reporter gene was determined by β-galactosidase activity assays performed at 30°C on extracts of at least two independent transformants containing both plasmids. The standard error was 15% of the mean. The β-galactosidase specific activity is expressed in nanomoles of *o*-nitrophenyl-b-D-galactopyranoside hydrolyzed per minute per milligram of protein. *^c* ²N, medium with no nitrogen source.

differential pools of glutamate, glutamine, arginine, ornithine, and lysine. As shown in Fig. 2B, the glutamate pool was equally distributed between the cytoplasm and the vacuole. Interestingly, nitrogen deprivation led to a rapid consumption of cytosolic glutamate without release from the vacuole. Glutamine was 90% vacuolar when the cells were grown on M.ammonia but was quickly released in the cytoplasm and immediately utilized as a nitrogen source upon nitrogen starvation (Fig. 2C). In contrast, the vacuolar arginine was released more slowly. After 40 min of starvation, only 15% of the vacuolar arginine was consumed without accumulation in the cytosol (Fig. 2D), while the production of arginase was maximal. These results did not contradict those of Kitamoto et al. (21). In their experiments, nitrogen starvation was performed after growth on arginine as the sole nitrogen source. In that condition at time zero, the arginase level was about 50-fold higher than that on M.ammonia, which explained why the vacuolar arginine pool decreased more rapidly but no burst of arginine was detected in the cytosol. During the nitrogen starvation, the ornithine pool remained constant (Fig. 2E), probably because the ornithine transaminase level was not sufficient to degrade it, and the lysine vacuolar pool accumulated (Fig. 2F), since lysine was not used as a nitrogen source and was less incorporated into proteins. These results indicate that the production of arginase after transfer to a nitrogen-free medium does not result from an induction in response to a higher cytosolic arginine concentration.

Arginase induction upon nitrogen starvation could result from the interaction between Ume6 and components of the ArgR-Mcm1 complex. We have previously shown that the control of *CAR1* expression by nitrogen availability required URS1, the arginine boxes (arginine upstream activation sequence [UASarg]), and the integrity of Ume6 and the ArgR-Mcm1 complex (11). We had proposed that when cells were starved for nitrogen, the release of URS1 facilitated the accessibility of the ArgR-Mcm1 complex to UASarg in spite of the weak arginine internal pool. Using the two-hybrid system we identified an interaction between Ume6 and ArgRI or ArgRII, but only under nitrogen starvation conditions. As shown in Table 6, in strain HY transformed with plasmid pME46 (GBD-ArgRI) or plasmid pNA33 (GBD-ArgRII) and plasmid pFV124 $(GAD-Ume6)$, the level of β -galactosidase was increased in the absence of nitrogen. No interaction between Ume6 and the pleiotropic regulators ArgRIII and Mcm1, which are also regulators of the arginine genes (data not shown), was detected. The interaction between Ume6 and ArgRI or ArgRII seems thus specific and physiological.

DISCUSSION

CARGRI, *CARGRII*, and *CARGRIII* gene products were identified as repressors of *CAR1* and *CAR2* genes. CargRI turned out to be Ume6, and we show here that CargRII is identical to Sin3 and that CargRIII is identical to Rpd3. These three proteins are part of a high-molecular-weight complex regulating the expression of a large set of genes, which explains the wide variety of names attributed to these three genes. Ume6 is able to interact with DNA in vitro at a sequence called URS1 and recruits Sin3, which then binds to Rpd3 causing repression of transcription through core histone deacetylation (20). Sap30, which is part of this protein complex, is also required for *CAR1* repression, whereas two other histone deacetylases, Hda1 and Hos2, belonging to other histone deacetylase complexes, do not control *CAR1* expression. In contrast, neither Gcn5 nor TAF145/130, both of which show histone acetyltransferase activity $(4, 32)$, is involved in the derepression of *CAR1*. Formation of a more active chromatin state at the *CAR1* promoter should thus depend on another histone acetyltransferase. As for other genes controlled by Ume6, the repression at *CAR1* results mainly from the action of the Sin3-Rpd3 complex, since deletion of *SIN3* and/or *RPD3* does not increase the derepression of arginase observed in a *ume6*-deleted strain. However, this derepression is always higher in a *ume6*-deleted strain than in a *sin3*- or *rpd3*-deleted strain. This could result from a repressing activity of Ume6 independent of histone deacetylase recruitment or from a partial effect of other histone deacetylases in the absence of Sin3 or Rpd3. The fact that Ume6 could play a role independent of Sin3 and Rpd3 is sustained by our observation that the growth of a *ume6* deletant is more impaired than the growth of *sin3* or *rpd3* deletants. This difference in behavior is increased in the S1278b background, in which deletion of *UME6* leads to lethality. Such a phenotype suggests that Ume6 could recruit positive transcription factors involved in the control of essential genes. It has already been reported that one of the positive effects of Ume6 is to recruit the transcriptional activator Ime1 to activate the expression of early meiotic genes upon nitrogen starvation (37). This Ime1-Ume6 complex formation requires an interaction between Rim11 and Ume6, resulting in a carbon source-dependent phosphorylation of Ume6 (25).

In the control of arginine catabolic genes, the role of the Ume6-Sin3-Rpd3 complex is to block the expression of *CAR1* and *CAR2* promoters as long as exogenous nitrogen is available. Indeed, a mutation in *UME6* abolishes completely the response of these two promoters to nitrogen depletion. Arginase and ornithine transaminase production under nitrogen starvation conditions also requires the integrity of the ArgR-Mcm1 complex (11). However, this enzyme synthesis does not result from a burst of arginine stored in the vacuole toward the cytosol, as shown by our differential arginine pool measurements. Under these growth conditions, arginine leaks slowly out of the vacuole to the cytosol, where it is used as a nitrogen source, without sufficient accumulation of arginine to allow the interaction between the ArgR-Mcm1 complex and the arginine boxes (11). This induction could rather result from an interaction between Ume6 and the components of the ArgR-Mcm1 complex, leading to a more efficient binding at the arginine boxes at low arginine concentration. Such a hypothesis is supported by our two-hybrid experiments showing an interaction between ArgRI or ArgRII and Ume6 only under nitrogen starvation conditions.

To confirm the importance of chromatin structure in the regulation of *CAR1* and *CAR2* expression, mapping nucleosomes along the promoters of these two genes under different growth conditions will be our next goal.

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