

# DNA SEQUENCE POLYMORPHISMS IN THE GENUS SACCHAROMYCES. I. COMPARISON OF THE *HIS4* AND RIBOSOMAL RNA GENES IN LAGER STRAINS, ALE STRAINS AND VARIOUS SPECIES

by

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The region of chromosome XII containing the *RDNI* gene which encodes the cytosolic ribosomal RNA molecules and the region of chromosome III containing the *HIS4* (histidine 4) gene were analysed in 30 lager yeast strains, 11 ale strains and 20 strains from a number of different species in the genus *Saccharomyces*.

With the aid of restriction endonuclease fragment patterns and cloned probes to the *RDNI* gene of *S. cerevisiae* three forms of this gene were identified, two of them corresponding to the previously known forms I and II and a third one characterized by an additional HindIII site located in the 3' spacer region. A more distantly related form of the *RDNI* gene containing a single HindIII restriction site was found in *Saccharomyces fermentati* and one form without any HindIII site in a wild yeast contaminant.

With the help of the restriction endonuclease fragments derived from the *HIS4* region seven genotypes can be recognized. They result from various combinations of three restriction endonuclease fragment patterns designated I, II and III, each pattern represents a chromosome.

All lager strains are homozygous for form II of the ribosomal RNA gene and heterozygous for patterns I and II of the *HIS4* gene. An exception is one German brewing strain which is homozygous for pattern II.

With one exception the ale strains were homozygous for form II of the *RDNI* gene and for pattern I of the *HIS4* gene. One British strain contains form I of the *RDNI* gene. Bakers yeast, *S. diastaticus* and *S. italicus* are homozygous for form I of the *RDNI* gene and for pattern I of the *HIS4* gene. In *S. bayanus* and *S. pastorianus* homozygosity for form III of the *RDNI* gene was combined with heterozygosity for patterns II and III of the *HIS4* gene. *S. uvarum* is homozygous for both form III of the *RDNI* gene and pattern III of the *HIS4* gene. Form III of the *RDNI* gene and patterns I and II of the *HIS4* gene were combined in a Chinese brewing strain and a strain designated as a type strain of *S. carlsbergensis*.

The nucleotide sequence polymorphisms are useful markers for strain characterization in addition to the generally used fermentation properties.

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Abbreviations: bp = basepairs; kb = kilobase; SSC = 0.15 M-NaCl, 15 mM-Na citrate; Tris = tris-(hydroxymethyl)-amino methane.

## 1. INTRODUCTION

In yeast taxonomic studies a genus or a species is usually delimited on the basis of morphology, fermentation characteristics (33), capacity of spore formation and ability to hybridize (15, 36). Closely related strains are distinguished from one another by individual fermentative abilities, enzyme polymorphisms (8, 37) and most recently nucleic acid sequence variations (4, 12, 17, 19, 20, 35). The analysis of dissociation of DNA and its reassociation with DNA or RNA has added unlimited possibilities for characterization of strains and species within as well as among genera. Low sequence homology has been established among the genera *Schwanniomyces*, *Saccharomyces*, *Debaryomyces* and *Pichia* (1, 25). Within the genus *Saccharomyces* some species show close homology (28).

In the present study two specific chromosomal regions of the *Saccharomyces* genomes in 30 lager strains, 11 ale strains and 20 strains from a number of different species have been characterized for polymorphisms using restriction endonuclease fragment patterns and molecular hybridization to specific probes. The first region is in chromosome XII and contains the *RDN1* gene encoding the cytosolic 25S, 5.8S, 18S and 5S ribosomal RNA molecules (20, 21, 22). The 5S sequence is transcribed separately from the opposite strand of the gene. In *Saccharomyces cerevisiae* this gene is present in over 100 tandemly repeated copies (22, 29). The gene is generally highly conserved in sequence and overall organization throughout the pro- and eukaryotic kingdoms (6, 9, 32, 34). As a probe was used the plasmid constructed by PETES, HERFORD and SKRYABIN (20) which contains form II of the *RDN1* gene. The second region analysed contains the *HIS4* gene on chromosome III which encodes three enzymatic functions in the biosynthesis of histidine. The plasmid containing this gene has been constructed by HOLMBERG et al. (11) and has previously disclosed the existence of at least two homeologous chromosomes III in the Carlsberg lager strain (12, 17). The nucleotide sequence of the *HIS4* region of one of them is so different from the corresponding region in *S. cerevisiae* that genetic recombination is not possible between them. Extensive nucleotide sequence differences

between the Carlsberg lager strain and *S. cerevisiae* have also been shown to exist in other portions of chromosome III (12) and in other chromosomes (35). Various *Saccharomyces* species have extensively diverged nucleotide sequences in the members of the *SUC* gene family encoding invertase (4).

## 2. MATERIALS AND METHODS

### 2.1. Strains and media

The strains used in this study are listed in Tables I, II and III. Agar plates with complete medium (YPD) and plates containing sporulation medium (SPOR) were prepared according to SHERMAN, FINK and HICKS (30). Growth on melibiose was tested on plates of the following composition: 0.2% melibiose (Merck 12240), 0.67% yeast nitrogen base without amino acids (Bacto 919-15) and 2% agar (Bacto 0140-01). All strains were propagated on YPD plates for DNA isolation and replicated to sporulation plates and melibiose containing plates. The plates were incubated at 22 °C, the frequency of asci determined in each strain after seven days (10) and the capacity to grow on melibiose recorded. The strains were further tested for excretion of  $\alpha$ -galactosidase (melibiase) by growing them on plates containing 0.2% galactose (Merck 4062), 0.67% yeast nitrogen base without amino acids (Bacto 919-15) and 2% agar for one or two days. Thereafter the colonies were covered with a layer of agar (5 ml) containing 1.5% agar, 31 mM-citric acid, 39 mM-KH<sub>2</sub>PO<sub>4</sub>, pH 4.0 according to the procedure of BUCKHOLZ and ADAMS (2). Instead of using *p*-nitrophenyl- $\alpha$ -D-galactoside as the indicator in the overlayer, 40  $\mu$ mole ml<sup>-1</sup> 4-methyl-umbelliferyl- $\alpha$ -D-galactoside (Sigma M-7633 USA) was included. Melibiase will produce free 4-methyl-umbelliferone which can be detected by its fluorescence (B.S. ENEVOLDSEN, pers. comm., 27). Detection of the fluorescing compound is possible after 5-40 min incubation of the plates at room temperature.

### 2.2. Preparation of DNA

DNA from the yeast cells was isolated by a modification of the method of CAMERON, PHILIPPSEN and DAVIS (3). Yeast cells were scraped from a YPD plate which previously had been incubated for 48 hours, suspended in 5 ml of 20

Table I.

Characteristics of lager yeast strains (*Saccharomyces carlsbergensis*). *RDNI*: Restriction endonuclease fragment pattern (cf. Fig. 9) of the *RDNI* gene encoding the cytosolic ribosomal RNA molecules. *HIS4*: Restriction endonuclease pattern (cf. Fig. 9) of the *HIS4* region in chromosome III. Spor: sporulation, Mel: capacity to grow on melibiose minimal medium, M.ase: Excretion of melibiase. -/+ : Indicates colonies of both types. n.d.: not determined. (-): Very poor growth on melibiose minimal medium.

Strain number	<i>RDNI</i>	<i>HIS4</i>	Spor%	Mel	M.ase	Source
BK1100	II	I + II	< 1	+	+	Denmark, Tuborg, 1945
BK1101	II	II	< 1	+	+	Germany, Pschorr, 1939
BK1103	II	I + II	1	+	+	Denmark, Carlsberg, 1964
BK1111	II	I + II	< 1	+	+	Germany, Weihenstephan, 1966
BK1112	II	I + II	< 1	+	+	Denmark, Alfr. Jørgensen no. 2190
BK1115	II	I + II	< 1	+	+	USA, Schaefers, New York, Belgian yeast, 1947
BK1116	II	I + II	< 1	+	+	Germany, Dortmunder Union, 1936
BK2224	II	I + II	< 1	+	+	Germany, Kindl Berlin, 1950
BK2229	II	I + II	< 1	+	+	Denmark, Alfr. Jørgensen no. 1522
BK2231	II	I + II	< 1	+	+	Sweden, Lyckholm, Gothenburg, 1943
BK2237	II	I + II	15	(-)	+	Pilsen, Erste Pilsener Akt. Br., 1937
BK2246	II	I + II	1	+	+	Denmark, Carlsberg lager strain
C82-FP2	II	I + II	2	+	+	Malta
C82-FP3	II	I + II	20	+	+	Italy
C82-FP4	II	I + II	5	+	+	Cameroun
C82-FP5	II	I + II	12	+	+	Norway
C82-FP6	II	I + II	3	+	+	France
C82-FP7	II	I + II	24	+	+	Portugal
C82-FP8	II	I + II	11	+	+	Portugal
C82-FP9	II	I + II	20	+	+	Canada
C82-FP11	II	I + II	< 1	-	+	Norway
C82-FP13	II	I + II	n.d.	-	+	Portugal
C82-FP14	II	I + II	6	-	+	Portugal
C82-FP18	II	I + II	7	+	+	Great Britain
C82-FP19	II	I + II	9	+	+	Cameroun
C82-FP20	II	I + II	10	+	+	Cameroun
C82-FP21	II	I + II	38	+	-	Portugal
C82-FP22	II	I + II	20	-/+	-/+	Sweden
C82-FP26	II	I + II	41	+	+	Great Britain
C82-FP32	II	I + II	6	+	+	Finland

mm-dithiothreitol (Sigma D-0632 USA) and incubated at 30 °C for 30-45 min. Then zymolyase-60000 (from *Arthrobacter luteus*, Seikagaku Kogyo Co. Ltd., Japan) was added (1.4 mg). The solution was further incubated at 30 °C for 30 min. Cell lysis was induced by addition of 0.2 ml 10% sodium dodecyl sulfate (SDS), 0.5 ml of 0.5 M-EDTA (pH 8.5) and 0.5 ml 2 M-Tris base (tris-(hydroxymethyl)-amino methane). After stirring the solution was heated to 65 °C for 30 min. The tube was then cooled on ice and one ml of 5 M-potassium acetate was added. After one hour on ice the tube was spun for 10-15 min at 4000 g

and the nucleic acids in the supernatant were precipitated in 70% ethanol. After one hour at -20 °C the precipitate was spun down at 4000 g for 10 min, resuspended in 10 mM-Tris, 1 mM-EDTA buffer (pH 7.5) and 2.5 µg ribonuclease A (Sigma R-5125) as well as 40 units of ribonuclease T<sub>1</sub> (Sigma R-8251 USA) were added. The ribonuclease treatment was performed for 30-45 min at room temperature.

After the ribonuclease treatment one volume of phenol (equilibrated with Tris-EDTA buffer) was added. The phenol treatment was performed twice and the aqueous phase was washed three

Table II.

Characteristics of ale strains (*Saccharomyces cerevisiae*). *RDNI*: Restriction endonuclease fragment pattern (cf. Fig. 9) of the *RDNI* gene encoding the cytosolic ribosomal RNA molecules. *HIS4*: Restriction endonuclease fragment pattern (cf. Fig. 9) of the *HIS4* region in chromosome III. Spor: Sporulation. Mel: Capacity to grow on melibiose minimal medium. M.ase: Excretion of melibiase. -/+ : Colonies of both types.

Strain number	<i>RDNI</i>	<i>HIS4</i>	Spor%	Mel	M.ase	Source
BK3300	II	I	<1	-	-	Denmark, Ceres 1920
BK3302	II	I	1	-	-	Denmark, Horsens 1932
BK3304	II	I	<1	-	-	Great Britain, Whitbread, London 1937
BK3313	II	I	5	-	-	Sweden, Stockholms Bryggerier, Clö 5 1949
BK3331	I	I	50	-	-	Great Britain, St. Austel Brewery
C82-FP10	II	I	<1	-	-	Canada
C82-FP15	II	I	>90	-	-	Great Britain
C82-FP16	II	I	59	-	-	Great Britain
C82-FP17	II	I	8	-	-	Great Britain
C82-FP23	II	I + II	1-20	-/+	-/+	Sweden
C82-FP25	II	I + II	20	-/+	-/+	Great Britain

Table III.

Characteristics of type strains and other strains for various species of *Saccharomyces*. *RDNI*: Restriction endonuclease fragment pattern (cf. Fig. 9) of the *RDNI* gene encoding the cytosolic ribosomal RNA molecules. *HIS4*: Restriction endonuclease pattern (cf. Fig. 9) of the *HIS4* region in chromosome III. Spor: Sporulation. Mel.: Capacity to grow on melibiose minimal medium. M.ase.: Excretion of melibiase. n.d.: not determined. 0: No hybridization.

Strain number	<i>RDNI</i>	<i>HIS4</i>	Spor%	Mel	M.ase	Source
BK4410	I	I	50	-	-	<i>S. odessa</i> ( <i>S. cerevisiae</i> ) Baarn 1950
BK4411	III	II+III	37	-	-	<i>S. bayanus</i> Baarn
BK4428	I	I	39	-	-	<i>S. diastaticus</i> NCYC 447, 1969
BK4433	III	I + IIa	3	+	+	<i>S. (carlsbergensis)</i> NCYC 396
BK4435	I	I	20	-	-	<i>S. cerevisiae</i> var. <i>ellipsoideus</i> NCYC 93
BK4436	different <i>RDNI</i>	0	n.d.	-	-	<i>S. fermentati</i> NCYC 161
BK4439	I	I	6	-	-	<i>S. italicus</i> (syn. <i>S. steineri</i> ) NCYC 406
BK4479	I	I	40	-	-	<i>S. ellipsoideus</i>
BK4505	III	II + III	60	-	-	<i>S. bayanus</i> NCYC 374
BK4515	III	I + II	12	+	+	<i>S. spec.</i> , China, brewing strain
BK4516	III	II + III	80	-	-	<i>S. pastorianus</i> NCYC 392
BK4525	n.d.	I	n.d.	n.d.	n.d.	<i>S. spec.</i> , wine yeast strain
BK4526	n.d.	I	n.d.	n.d.	n.d.	<i>S. spec.</i> , wine yeast strain
C81-1471	I	I	43	-	-	<i>S. spec.</i> , wild yeast, beer contaminant
C81-1483	I	I	>90	-	-	<i>S. cerevisiae</i> , bakers yeast
C81-1510	III	IIIa	<1	+	+	<i>S. uvarum</i> CBS 395., P. PHILIPPSEN
C81-1511	III	III	4	+	+	<i>S. uvarum</i> R.DAVIS, P.PHILIPPSEN
C81-1537	I	I	3	-	-	<i>S. spec.</i> , Belgium, brewing strain
C82-CG625	n.d.	I + II	n.d.	n.d.	n.d.	C. GJERMANSEN
C82-FP33	different <i>RDNI</i>	0	n.d.	n.d.	n.d.	This study

times with one volume of water saturated ether. After removal of the ether 1/10 volume 5 M-sodium acetate and 96% ethanol were added to a final concentration of 70% ethanol. After one hour at -20 °C the precipitated DNA was spun

down and resuspended in Tris-EDTA.

Plasmid DNAs were prepared and isolated in preparative scale according to HOLMES and QUIGLEY (13).

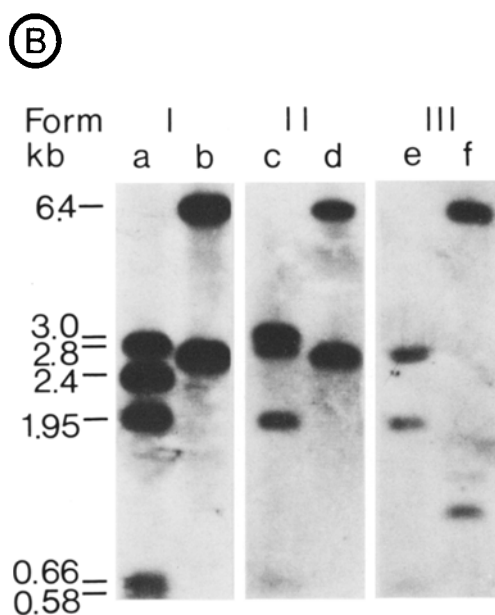
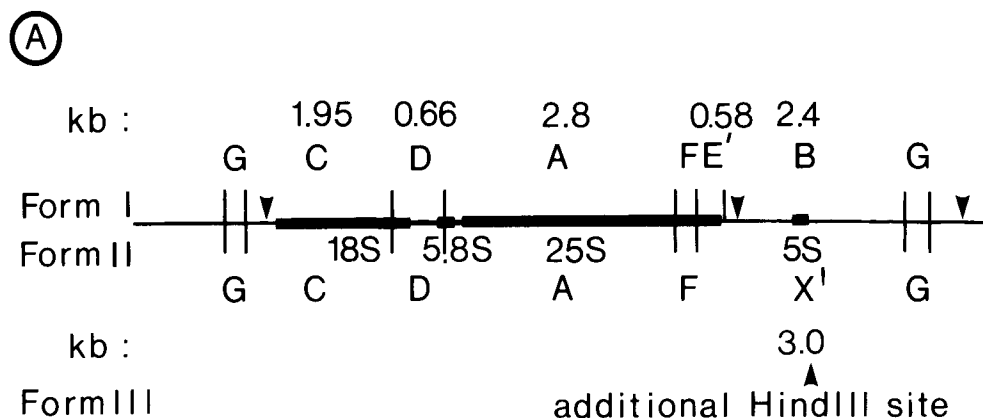


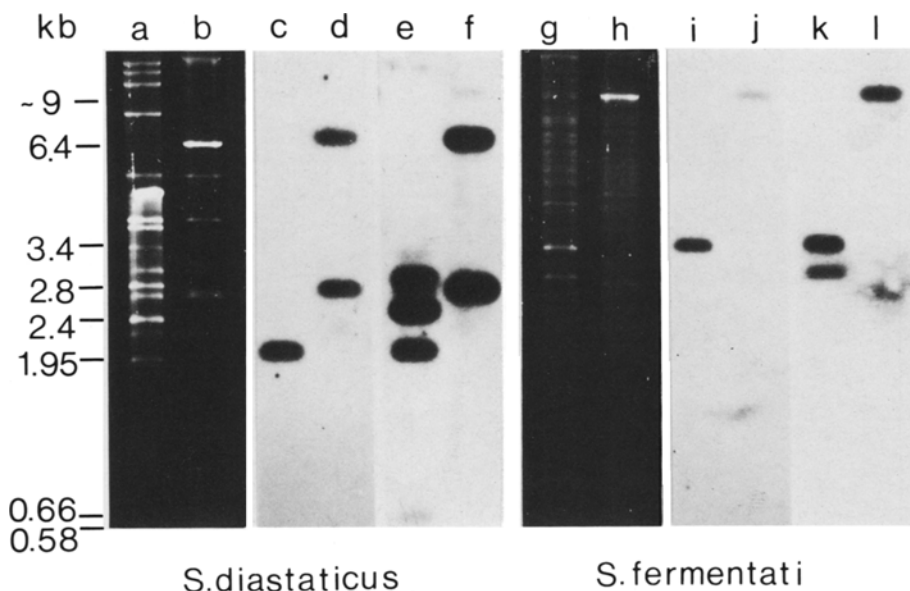
Figure 1. A: EcoRI restriction site maps (vertical bars) of the forms I and II of the *RDN1* gene according to PETES et al. (21). Also indicated are the HindIII sites (arrowheads). B: Autoradiograph of restriction endonuclease fragments separated by gel electrophoresis, transferred to a filter and hybridized to the radioactively labelled *RDN1* gene containing plasmid. The DNA under I was isolated from *S. ellipsoideus* BK4479, that under II from the Carlsberg lager strain BK2246 and that under III from *S. bayanus* BK4411. The DNA of the respective strains has been subjected to either EcoRI digestion (lanes a, c and e) or HindIII digestion (lanes b, d and f). About 2  $\mu\text{g}$  of DNA has been loaded in each slot. Plasmid pY1rA12 was labelled with  $10^8$  cpm  $[\alpha\text{-}^{32}\text{P}]$  dATP  $\cdot \mu\text{g}^{-1}$  of DNA. Molecular hybridization and washing of the filter was performed at 60  $^{\circ}\text{C}$  with  $3\times\text{SSC}$ . Form I is cut into seven EcoRI fragments of which five, namely A, B, C, D and E, are seen in lane a. Form II is cut into six fragments of which four ( $X' = B+E$ , A, C and D) are seen in lane c. Form III contains a double band of 2.8-2.9 kb representing fragment

A and a fragment derived from  $X'$  possibly by a deletion. It also contains bands corresponding to fragments C (1.95 kb) and D (0.66 kb). HindIII cuts the *RDN1* gene of forms I and II (lanes b and d) into two fragments with sizes of 6.4 kb and 2.6 kb. Form III has an additional HindIII site which generates two fragments of 1.45 kb and 1.15 kb from the 2.6 kb fragment of forms I and II (lane f).

### 2.3. Molecular hybridization analysis

Restriction endonucleases EcoRI, HindIII, PstI, PvuII and Sall were obtained from Boehringer Mannheim and used according to MANIATIS, FRITSCH and SAMBROOK (16). DNA fragments were separated electrophoretically in 0.7% agarose (Sigma A-6877 USA) at 1.25 V  $\cdot \text{cm}^{-1}$  for 14-16 hours. The gels were stained with ethidium bromide (Sigma E-8751) (1

$\mu\text{g} \cdot \text{ml}^{-1}$ ) in 1 litre of gelbuffer (Tris 10.8 g, disodium EDTA 0.93 g and boric acid 5.5 g, pH 8.3) (18) and photographed in UV-light. Transfer of the DNA fragments from the agarose gel was done either to a nitrocellulose filter (Millipore HAWP 000 10) or to GENESCREEN (New England Nuclear) by the method of SOUTHERN (31), but with the modification that transfer of DNA to GENESCREEN was performed in low salt

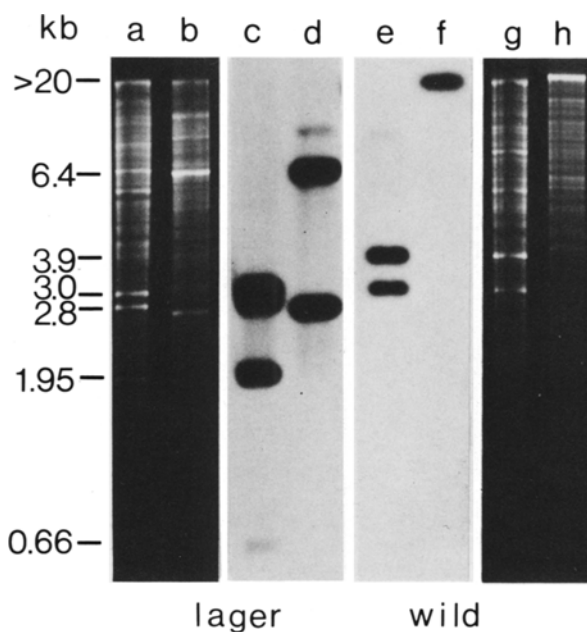


Lane	Restriction endonuclease	<i>RDN1</i> probe	rDNA bands (kb)	Form
a	EcoRI	-	2.8,2.4,1.95	I
b	HindIII	-	6.4,2.6	"
c	EcoRI	"18S"pY1rA12	1.95	"
d	HindIII	"18S"pY1rA12	6.4,2.6	"
e	EcoRI	pY1rA12	2.8,2.4,1.95,0.66,0.58	"
f	HindIII	pY1rA12	6.4,2.6	"
g	EcoRI	-	3.4,2.8	
h	HindIII	-	8.5-9	
i	EcoRI	"18S"pY1rA12	3.4	
j	HindIII	"18S"pY1rA12	8.5-9	
k	EcoRI	pY1rA12	3.4,2.8	
l	HindIII	pY1rA12	8.5-9	

Figure 2. Restriction endonuclease fragment patterns of *S. diastaticus* NCYC 447 (lanes a to f) and *S. fermentati* NCYC 161 (lanes g to l). Lanes a, b, g and h present the ethidium bromide stained fragments separated by gel electrophoresis. The other lanes are hybridizations with *RDN1* probes according to SOUTHERN (31). Hybridization to filter bound DNA was performed at 60 °C with 3×SSC.

buffer (1×SSC). Labelling of the plasmid pY1rA12 (20) carrying Form II of the *RDN1* gene, its derivative ("18S") (this study) and the *HIS4* plasmid pC503 containing a 9.4 kb PstI fragment (11) with [ $\alpha$ -<sup>32</sup>P]dATP (NEG-012A New England Nuclear) was done by nick translation according to RIGBY et al. (26) giving a specific activity in the range of 10<sup>7</sup> to 10<sup>8</sup> cpm · μg<sup>-1</sup> of plasmid DNA. For characterization of the *HIS4*

region and the rDNA genes by molecular hybridization the method of DENHARDT (7) with the modifications introduced by JEFFREYS and FLAVELL was used (14). The molecular hybridizations were performed in the temperature range of 58 °C to 68 °C. Autoradiographs were obtained at -70 °C in Kodak X-Omatic cassettes with regular screens.



Lane	Restriction endonuclease	<i>RDNI</i> probe	rDNA bands (kb)	Form
a	EcoRI	-	3.0,2.8,1.95,0.66	II
b	HindIII	-	6.4,2.6	"
c	EcoRI	pY1rA12	3.0,2.8,1.95,0.66	"
d	HindIII	pY1rA12	6.4,2.6	"
e	EcoRI	pY1rA12	3.9,3.0	
f	HindIII	pY1rA12	>20	
g	EcoRI	-	3.9,3.0	
h	HindIII	-	>20	

Figure 3. Identification of a wild yeast with the aid of the *RDNI* gene. Lanes a to d are from the Carlsberg lager strain while lanes e to h are from the contaminant C82-FP33 in the brewing yeast. Hybridization to filter bound DNA and washing was performed at 60 °C with 3×SSC.

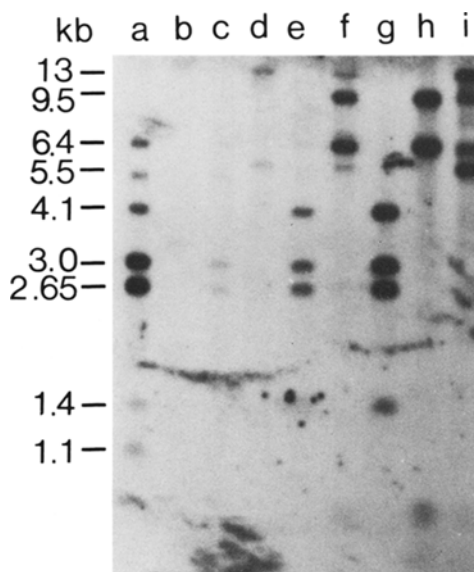
### 3. RESULTS

#### 3.1. Physiological characteristics of the *Saccharomyces* strains analysed

In Tables I, II and III are given for each strain the capacity to sporulate, to grow on melibiose minimal medium and to excrete the enzyme melibiase. In Table I are grouped the lager strains (*S. carlsbergensis*), in Table II the ale strains (*S. cerevisiae*) and in Table III various other yeast strains including *Saccharomyces* type strains obtained from different culture collections. The lager strains (Table I) tend to be poor sporulators. (Here tested at 22 °C.) Twenty-five of the strains could grow on melibiose and excrete melibiase. Four of the lager strains would not grow on the

melibiose minimal plates but displayed detectable amounts of melibiase activity when the strains were grown on galactose for induction of the enzyme and subsequently tested with 4-methyl-umbelliferyl- $\alpha$ -D-galactoside. One strain grew on melibiose but the test for enzyme excretion was negative. One sample contained a mixture of cells with a wide range of melibiose activity and a wide range of sporulation capacity.

Only one of the ale strains analysed (Table II) sporulated with a high frequency at 22 °C. Nine of the strains were unable to grow on melibiose and excrete melibiase. Two of the samples contained a mixture of cells with a wide range of melibiase activity and capacity to grow on



Lane	restriction endonuclease	strain	<i>HIS4</i> bands (kb)	pattern
a	EcoRI	Carlsberg lager BK2246	6.2, 5.0, 4.1, 3.0, 2.65, 1.4, 1.1	I and II
b		bacteriophage $\lambda$ DNA		
c	EcoRI	German lager BK1101	6.2, 5.0, 3.0, 2.65	II
d	HindIII	German lager BK1101	13.0, 5.5	II
e	EcoRI	<i>S. (carlsbergensis)</i> NCYC396	6.2, 4.1, 3.0, 2.65, 1.4	I and II
f	HindIII	<i>S. (carlsbergensis)</i> NCYC396	13.0, 9.5, 6.4, 5.5, 0.69, 0.62	I and II
g	EcoRI	<i>S. italicus</i> NCYC406	4.1, 3.0, 2.65, 1.4	I
h	HindIII	<i>S. italicus</i> NCYC406	9.5, 6.4, 0.69, 0.62	I
i	HindIII	Carlsberg lager BK2246	13.0, 9.5, 6.4, 5.5, 0.69, 0.62	I and II

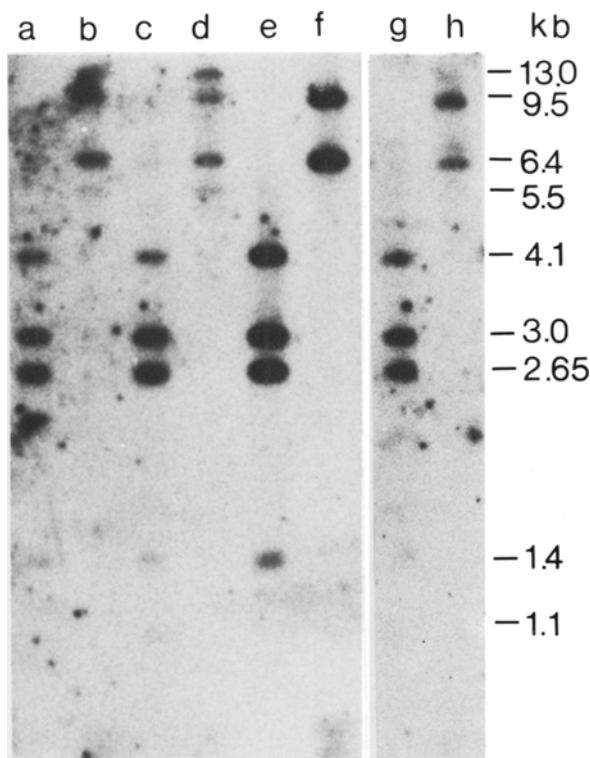
Figure 4. Autoradiograph of restriction endonuclease fragment patterns of alleles of the *HIS4* region in four *Saccharomyces* strains. Lanes a, c, e, and g contain EcoRI digested DNA, lanes d, f, h and i are loaded with HindIII digested DNAs. Molecular hybridization of the radioactively labelled *HIS4* containing plasmid and washing of the filter bound DNA was performed at 60 °C with 3 $\times$ SSC. 1  $\mu$ g of the plasmid DNA was labelled with 10<sup>8</sup> cpm [ $\alpha$ -<sup>32</sup>P] dATP.

melibiose, but only one of these revealed a similar variability in sporulation frequency.

A wide range of sporulation frequencies at 22 °C was observed among the various species and type strains studied (Table III). Four accessions could ferment melibiose and excrete melibiase. These were the two strains of *S. uvarum*, a brew-

ing yeast from China and the strain NCYC 396 considered as a type strain of *S. carlsbergensis*. These strains are not listed under the lager yeast because they clearly differ from them in the organization of the *RDN1* and *HIS4* genes as is detailed below.





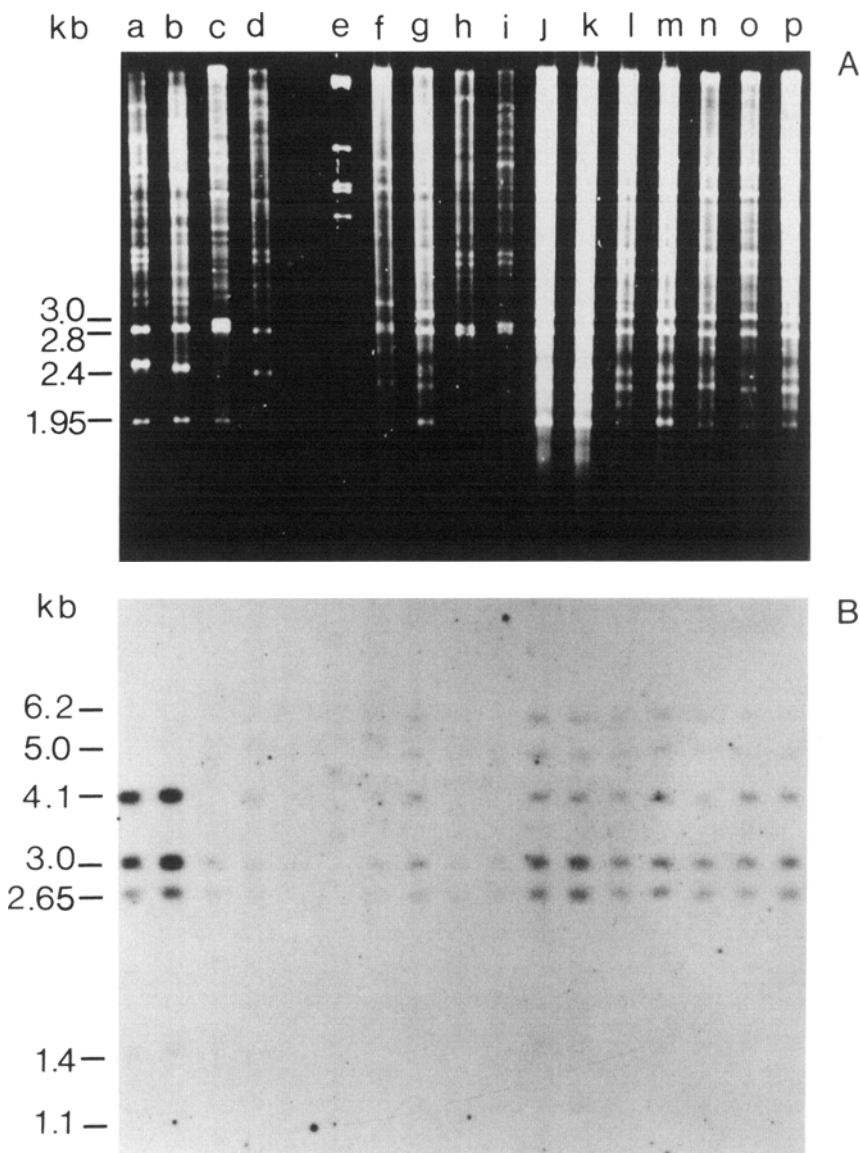
Lane	restriction endonuclease	strain	<i>HIS4</i> bands (kb)	pattern
a	EcoRI	lager strain BK1100	4.1, 3.0, 2.65, 1.4*	I and II
b	HindIII	lager strain BK1100	13.0, 9.5, 6.4, 5.5*	I and II
c	EcoRI	lager strain BK2237	6.2, 4.1, 3.0, 2.65, 1.4, 1.1*	I and II
d	HindIII	lager strain BK2237	13.0, 9.5, 6.4, 5.5*	I and II
e	EcoRI	ale strain BK3300	4.1, 3.0, 2.65, 1.4	I
f	HindIII	ale strain BK3300	9.5, 6.4	I
g	EcoRI	<i>S. (carlsbergensis)</i> NCYC396	4.1, 3.0, 2.65, 1.4	I and II
h	HindIII	<i>S. (carlsbergensis)</i> NCYC396	13.0, 9.5, 6.4*	I and II

Figure 5. Autoradiograph of restriction endonuclease fragment patterns of *S. (carlsbergensis)* NCYC396 compared with two lager strains and one ale strain. Lanes a, c, e and g contain EcoRI digested DNA while lanes b, d, f and h are loaded with HindIII digested DNA. Approximately 1 µg of DNA has been applied in each slot. Molecular hybridization of the radioactively labelled *HIS4* containing plasmid has been performed at high stringency of hybridization (68 °C with 3xSSC). Washing was done at low salt condition at the same temperature. \* = hybridizing with low intensity

### 3.2. Molecular analysis of the rDNA genes corresponding to the *RDNI* gene in *Saccharomyces cerevisiae*

The EcoRI and HindIII restriction site maps

of the two previously identified forms of the *RDNI* gene are given in Figure 1A (5, 20, 23, 24). Also indicated are the regions encoding the different cytosolic ribosomal RNA molecules and



the spacers separating them. As an example for the restriction endonuclease fragment pattern of form I the one for *Saccharomyces ellipsoideus* (BK4479) is presented (Figure 1B). Of the seven EcoRI fragments the five largest are easily identified (lane a). HindIII produces only two fragments as seen in lane b of Figure 1B. Form II of the *RDN1* gene, here represented by the Carlsberg lager strain (BK2246), yields the same HindIII restriction endonuclease fragment pattern as form I but is distinguished from the latter by the absence of the EcoRI restriction site sepa-

rating the E and B fragments. This leads to a fragment X' with a size of about 3 kb (lane c, Figure 1B).

In the course of this investigation a third form of the *RDN1* gene has been discovered. As an example for the gene the restriction endonuclease fragment patterns of the DNA from *Saccharomyces bayanus* (BK4411) are given in Figure 1B. Form III is distinguishable from forms I and II in both the EcoRI and the HindIII patterns. In the EcoRI pattern fragment A (2.8 kb) and a fragment corresponding to X' but

Lane	strain	<i>RDNI</i> Form	<i>HIS4</i> bands	pattern
a	<i>S. spec.</i> C81-1471	I	4.1, 3.0, 2.65, 1.4	I
b	<i>S. cerevisiae</i> C81-1483	I	4.1, 3.0, 2.65, 1.4	I
c	<i>S. uvarum</i> C81-1511	III	5.0, 3.0, 2.65	III
d	brewing strain C81-1537	I	4.1, 3.0, 2.65, 1.4	I
e	bacteriophage $\lambda$ DNA			
f	lager strain BK4515	III	6.2, 4.1, 3.0, 2.65	I and II
g	Carlsberg lager strain BK2246	II	6.2, 5.0, 4.1, 3.0, 2.65, 1.4, 1.1	I and II
h	<i>S. bayanus</i> NCYC374	III	3.0, 2.65	
i	<i>S. pastorianus</i> NCYC392	III	3.0, 2.65	
j	Carlsberg lager strain BK2246	II	6.2, 5.0, 4.1, 3.0, 2.65, 1.4, 1.1	I and II
k	lager strain BK1103	II	"	I and II
l	lager strain BK1112	II	"	I and II
m	lager strain BK1115	II	"	I and II
n	lager strain BK1116	II	"	I and II
o	lager strain BK2229	II	"	I and II
p	lager strain BK2236	II	"	I and II

◀ Figure 6. EcoRI restriction endonuclease fragment patterns of the genes *RDNI* (Figure 6A) and *HIS4* (Figure 6B). 0.5 to 2  $\mu$ g of EcoRI digested DNA has been loaded in each slot. The major *RDNI* gene fragments are visible in Figure 6A. Molecular hybridization of the radioactively labelled *HIS4* probe has been performed at 58 °C with 3 $\times$ SSC. Washing of the filter bound DNA was done at low salt conditions (0.1 $\times$ SSC at 58 °C).

smaller (also about 2.8 kb) appear as a double band. In addition, fragments C (1.95 kb) and fragment D (0.66 kb) are present (Figure 1B, lane c). HindIII cuts this *RDNI* gene into three fragments, with sizes of 6.4, 1.45 and 1.15 kb, respectively, (Figure 1B, lane c).

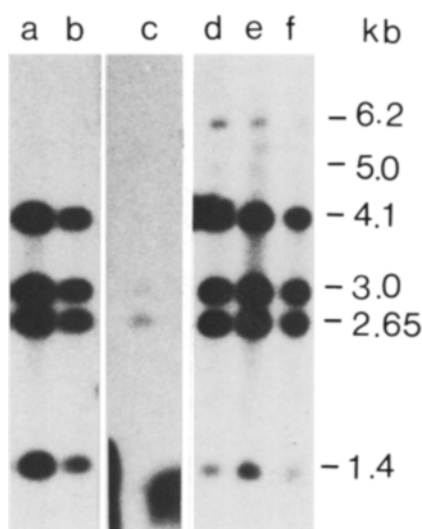
All 30 lager strains (*Saccharomyces carlsbergensis*) had form II of the *RDNI* gene (Table I). Of the 11 ale strains (*Saccharomyces cerevisiae*) all had form II with the exception of strain BK3331, which contained form I (Table II). Form III of the *RDNI* gene was characteristic for *Saccharomyces bayanus*, *S. pastorianus*, *S. uvarum*, a brewing strain from China and strain NCYC 396 considered as a type strain of *S. carlsbergensis* (Table III). All other species studied except one contained form I.

*Saccharomyces fermentati* (NCYC 161) revealed a very different form of the *RDNI* gene. In order to compare this additional form of the *RDNI* gene with form I it was probed with both pY1rA12 plasmid containing the entire *RDNI* gene of form I (*S. cerevisiae*) and a plasmid derivative thereof containing 100 bp of EcoRI fragment G and 900 bp of fragment C thus covering about half of the region transcribed into the 18S

ribosomal RNA (Figure 1A).

This latter plasmid ("18S"pY1rA12) originated as a recombinant product from the original plasmid which contained as insert a fragment bordered on one side by a portion of the G region and on the other by a full G region plus a 900 bp segment of the C region. A recombination event in the duplicated segments has deleted all of the *RDNI* gene except the 1000 bp retained in the "18S"pY1rA12 plasmid.

Figure 2 shows that the plasmid containing the 18S region hybridizes in agreement with the expectation to the EcoRI fragment C (1.95 kb) of the *RDNI* gene (form I) and to both HindIII fragments (lanes c and d). In *S. fermentati* the 18S region probe hybridizes to a 3.4 kb EcoRI fragment instead of the 1.95 kb fragment (Figure 2, lane i). The plasmid containing the entire *RDNI* gene recognizes in addition the EcoRI fragment C of 2.8 kb (Figure 2, lane k). Both probes hybridize to a single HindIII generated fragment about 9 kb in size (Figure 2, lanes j and l), revealing that the entire *RDNI* gene of *S. fermentati* contains a single HindIII restriction site. The restriction endonuclease fragment patterns of *S. fermentati* presented in Figure 2 are the same as those de-



Lane	strain	<i>HIS4</i> bands (kb)	pattern
a	<i>S. spec.</i> BK4525	4.1, 3.0, 2.65, 1.4	I
b	<i>S. spec.</i> BK4526	4.1, 3.0, 2.65, 1.4	I
c	<i>S. uvarum</i> C81-1510	3.0, 2.65	
d	Carlsberg lager strain BK2246	6.2, 4.1, 3.0, 2.65, 1.4	I and II
e	Hybrid strain C82-CG625	6.2, 4.1, 3.0, 2.65, 1.4	I and II
f	lager strain C82-FP32	6.2, 4.1, 3.0, 2.65, 1.4	I and II

Figure 7. Comparison of *Eco*RI restriction endonuclease *HIS4* fragments of six *Saccharomyces* strains. Approximately 2  $\mu$ g of *Eco*RI digested DNA have been loaded in each slot. 1  $\mu$ g of the *HIS4* containing plasmid was labelled with  $6 \times 10^7$  cpm of [ $\alpha$ - $^{32}$ P] dATP. The molecular hybridization of the probe and washing of the filter were performed at 62 °C with 3 $\times$ SSC. The *S. uvarum* strain C81-1510 shows low homology to the probe compared to the other five strains.

scribed for *S. rosei* (34).

During a microbiological analysis of a production sample of the Carlsberg lager strain BK2246 an aberrant colony was found and cultured (C82-FP33). To determine whether the colony was a mutant of the lager strain or a wild yeast colony, the DNAs from the two strains were compared by restriction analysis and molecular hybridization to the *RDNI* gene probe. In Figure 3, lanes a, b and g, h, are shown the fluorescing bands of the ethidium bromide stained fragments from the endonuclease digested DNA of the Carlsberg lager strain (lane a: *Eco*RI, lane b: *Hind*III) and of the aberrant strain (lane g: *Eco*RI and lane h: *Hind*III). The strongly fluorescing bands originate from restriction fragments present in multiple copies and they produce quite dissimilar patterns in the two strains. Such sequence divergence cannot be obtained by a single mutational

event. The autoradiographs in lanes c - f identify the fluorescing bands containing DNA fragments that can hybridize to the radioactively labelled *RDNI* gene probe. The Carlsberg lager strain BK2246 (lanes c and d) gives the band patterns characteristic for the *RDNI* gene form II, while in the wild yeast different DNA fragments are identified by the *RDNI* probe (lanes e and f). The probe pY1rA12 hybridizes to two *Eco*RI fragments with a size of 3.9 kb and 3.0 kb (lane e) and to a single *Hind*III fragment band with a size larger than 20 kb (lane f). Thus the rDNA gene of the contaminant was not cut by *Hind*III. While the *Eco*RI and *Hind*III restriction fragment patterns of this contaminant superficially look similar to those of *S. fermentati* (Figure 2, k and l) measurement of the *RDNI* fragment sizes reveal very different genotypes in the two strains.

### 3.3. Molecular analysis of the *HIS4* region in *Saccharomyces* strains

With regard to the *HIS4* region two variants of chromosome III have so far been identified in the genus *Saccharomyces* (17). Laboratory strains of *Saccharomyces cerevisiae* contain a chromosome III which upon probing with the *HIS4* containing plasmid pC503 (11) is recognized by four EcoRI restriction fragments with sizes of 4.1, 3.0, 2.65, and 1.4 kb. The pattern of this chromosome is designated as I and corresponds to a pattern of six HindIII generated fragments with sizes of 9.5, 6.4, 0.69, 0.62, 0.46 and 0.39 kb. The Carlsberg lager strain contains a chromosome with pattern I and a homeologous chromosome which upon endonuclease digestion generates pattern II consisting of EcoRI fragments with sizes of 6.2, 5.0, 3.0, 2.65, 1.1 kb and two HindIII fragments with sizes of 13.0 and 5.5 kb. All these fragments hybridize only weakly with the *HIS4* containing plasmid indicating extensive sequence differences between the "cerevisiae" *HIS4* region and that in the homeologous chromosome.

Of the strains analysed in Figure 4, *S. italicus* (lanes g and h) has pattern I, the German lager strain BK1101 has pattern II (lanes c and h) while the Carlsberg lager strain BK2246 has a summation of patterns I and II and thus contains both homeologous chromosomes (lanes a and i). The difference in the hybridization intensity of the fragments originating from the two chromosomes III in the lager yeast is clearly illustrated in lane a. A drastic difference in hybridization intensity is also noted between fragments of pattern I (lanes g and h) and those of pattern II in the German lager strain (lanes c and d), the latter hybridizing with low efficiency. Strain NCYC 396 considered as a type strain of *S. carlsbergensis* reveals the combination of patterns I and II when EcoRI or HindIII digested DNA is probed with the radioactive plasmid (lanes e and f). Probably the strain is similar to the tested lager yeasts, but it was consistently observed that the probe hybridized less intensely to the restriction fragments of the NCYC 396 strain than to those of lager strains (cf. lane i with lane f).

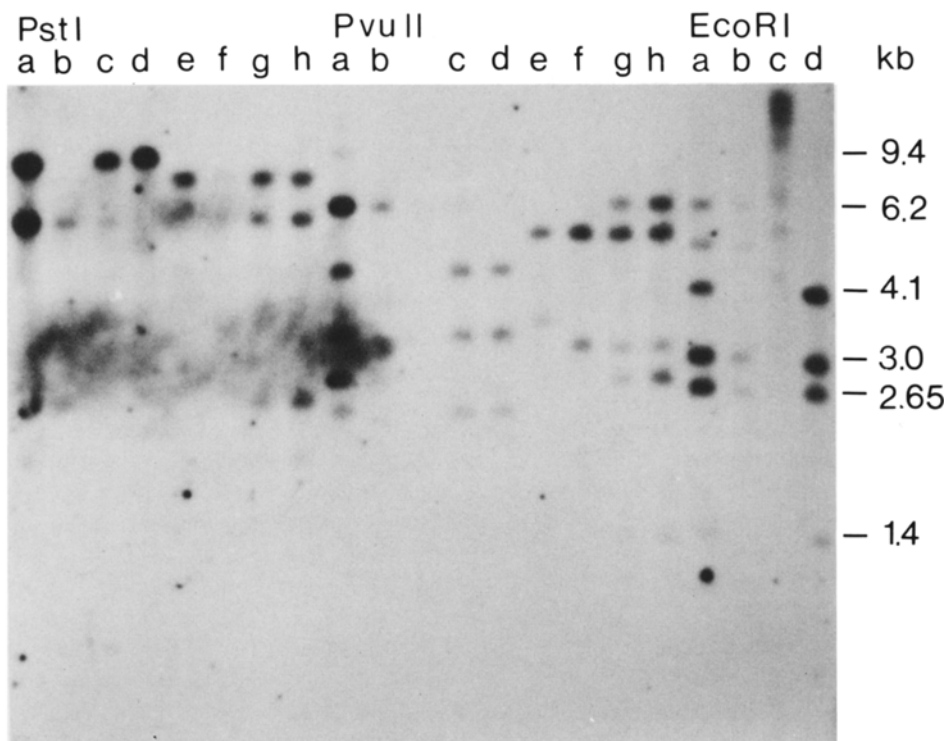
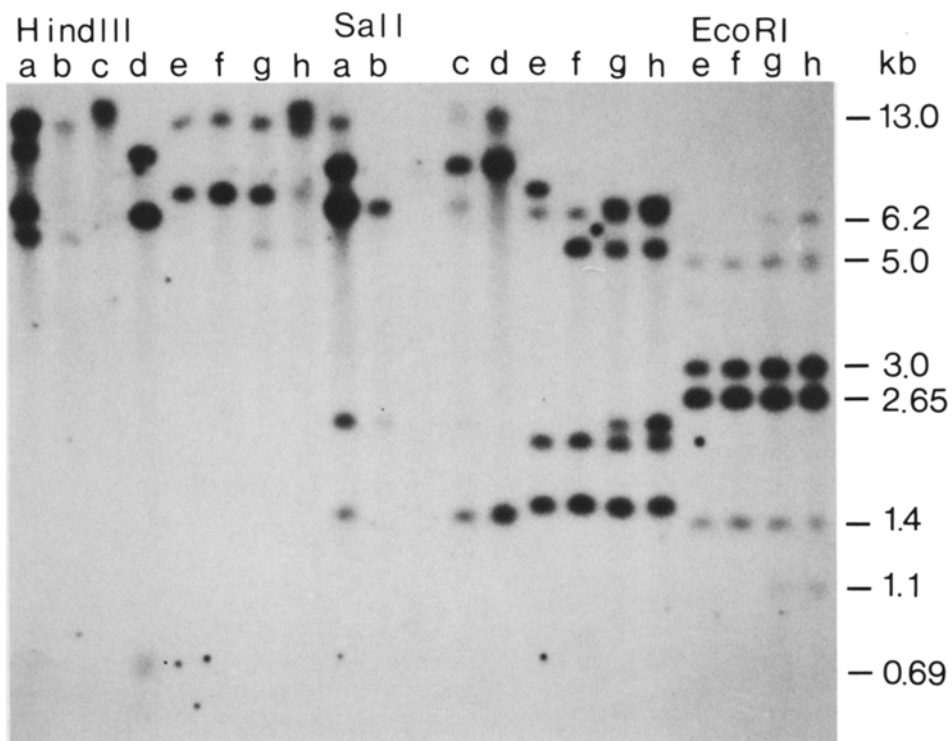
At high stringency of hybridization (68 °C, 3×SSC) the fragments diagnostic for pattern II of the *HIS4* region in strain NCYC 396 *S. carls-*

*bergensis* and the lager strain hybridize with weak intensity (Figure 5). This is visible for the 6.2 and 1.4 kb EcoRI fragment band as well as for the 13.0 and 5.5 kb HindIII fragment bands. The absence of a 4.1 kb EcoRI fragment in pattern II results in a less intense band than that shown by the pattern I allele present in the ale strain BK3300 (Figure 5, lane e).

In Figure 6 are shown the fluorograph of the ethidium bromide stained fragments of the EcoRI digested DNA (Figure 6A) and the autoradiograph after transfer of the fragments to a filter and their hybridization to the radioactively labelled *HIS4* probe (Figure 6B). In this experiment the chromosomes displaying pattern I of *HIS4* in the *Saccharomyces* strains C81-1471 and C81-1483 (lane a and b) show stronger homology to the probe than those in the lager strains (lanes j through p).

Three strains show weaker homology of their *HIS4* regions to the probe than all the other strains (Figure 6 lanes c,h,i). These are *S. uvarum* C81-1511, *S. pastorianus* NCYC 392 (BK4516) and *S. bayanus* NCYC 374 (BK4505) which only give detectable hybridization to the 2.65, 3.0 and 5.0 kb fragments containing the *HIS4* region. At higher stringency (62 °C 3×SSC) a prominent difference in hybridization intensity appears between the 2.65 kb and 3.0 kb fragments of *S. uvarum* strain C81-1510, whereas such a difference is not observed in the other *Saccharomyces* strains (Figure 7). The *Saccharomyces* strains, BK4525 and BK4526, contain pattern I of *HIS4*, while the Carlsberg lager strain BK2246, the hybrid lager strain C82-CG625 and the Finnish lager strain C82-FP32 carry both patterns I and II of *HIS4*.

At low stringency (58 °C) six to seven genotypes can be distinguished with the *HIS4* probe. The DNA of the eight strains has been digested with restriction endonucleases EcoRI, HindIII, PstI, PvuII and Sall (Figure 8). Looking at the patterns obtained with the restriction endonuclease HindIII (lanes a to h) six genotypes can be distinguished: (a) Pattern I + II from the Carlsberg lager strain. (b) The German lager strain with pattern II. (c) This lane contains the DNA of the strain considered a type strain of *S. carlsbergensis*, but was not cut in the particular experiment. Its pattern I + II has been presented in



Strain, pattern	Restriction fragment sizes in kb				
	EcoRI	HindIII	PvuII	PstI	Sall
<b>a</b> Carlsberg lager strain BK 2246 pattern I and II	6.2, 5.0, 4.1, 3.0, 2.65, 1.4, 1.1	13.0, 9.0, 6.4, 5.5, 0.69, 0.62,	9.4, 6.2, 4.9 3.5, 2.75, 1.4	9.4, 6.0, 2.5	13.0, 9.4, 6.7, 2.3, 1.5
<b>b</b> German lager strain BK 1101 pattern II	6.2, 5.0, 3.0, 2.65	13.0, 5.5	6.2, 3.5	6.0, 2.5	6.7, 2.3
<b>c</b> <i>S. carlsbergensis</i> NCYC396 pattern I and IIa	6.2, 5.0, 4.1, 3.0, 2.65	(cf. Fig. 4 and 5)	9.4, 6.2, 4.9 3.5, 2.5	9.4, 6.0	13.0, 9.4, 6.7, 2.3, 1.5
<b>d</b> <i>S. cerevisiae</i> C81-1483 pattern I	4.1, 3.0, 2.65 1.4	9.5, 6.4, 0.69 0.62	9.4, 4.9, 3.5 2.5	9.4	13.0, 9.4, 1.5
<b>e</b> <i>S. uvarum</i> C81-1510 pattern IIIa	5.0, 3.0, 2.65 1.4	13.0, 7.5	5.9, 3.8, 1.4, 1.3	8.0	7.5, 6.4, 2.1, 1.6
<b>f</b> <i>S. uvarum</i> C81-1511 pattern III	5.0, 3.0, 2.65, 1.4	13.0, 7.5	5.9, 3.4, 1.4, 1.3	8.0	6.4, 5.5, 2.1, 1.6
<b>g</b> <i>S. pastorianus</i> NCYC392 pattern II and III	6.2, 5.0, 3.0, 2.65, 1.4, 1.1	13.0, 7.5, 5.5	6.2, 5.9, 3.4 2.75, 1.4, 1.2	8.0, 6.0, 2.5	6.7, 6.4, 5.5 2.3, 2.1, 1.6
<b>h</b> <i>S. bayanus</i> NCYC374 Pattern II and III	6.2, 5.0, 3.0, 2.65, 1.4, 1.1	13.0, 7.5, 5.5	6.2, 5.9, 3.4 2.75, 1.4, 1.2	8.0, 6.0, 2.5	6.7, 6.4, 5.5 2.3, 2.1, 1.6

Figure 8. Structural differences in the *HIS4* region of eight *Saccharomyces* strains revealed by restriction endonuclease analysis and molecular hybridization to the radioactively labelled plasmid pC503 containing the *HIS4* region (11). Each of the isolated DNAs have been digested by five restriction enzymes.

Four of the strains are heterozygous for *HIS4* (a, c, g and h), the others seem to be homozygous for either pattern I, II or III (b, d, e and f).

Figure 4, lane f. (d) The typical pattern I of *S. cerevisiae*. (e) and (f) The patterns of the *S. uvarum* accessions contain a 13 kb and a more intense hybridizing 7.5 kb fragment designated as pattern III. (g) and (h) The patterns are from *S. pastorianus* and *S. bayanus* and represent a summation of pattern II (=b) and III (=f). (The DNA in lane h was only partially cut.)

The Sall patterns (lanes a - h) permit the distinction of seven genotypes. (a) The Carlsberg lager strain pattern I + II reveals 5 fragments, which represent the summation of the fragments present in (b) (= pattern II) and (d) (= pattern I). (c) The strain considered as a type strain for *S. carlsbergensis* contains all the fragments present in the lager yeast (a) but there is a drastic difference between the two strains in hybridization intensity. (e) and (f) The restriction fragment pattern of the *S. uvarum* strain (C81-1510) is distinguished from the *S. uvarum* strain (C81-1511)

by having a 7.5 kb fragment instead of a 5.5 kb fragment. (g) and (h) *S. pastorianus* and *S. bayanus* have patterns consisting of the summation of those in (b) and (f) i.e. pattern II + III.

The PstI and PvuII fragment patterns of the 8 strains confirm the existence of seven genotypes as detailed in the legend to Figure 8. The EcoRI restricted DNA of *S. uvarum* gives the pattern III for this endonuclease (lanes e and f). It consists of 4 fragments with sizes of 5.0, 3.0, 2.65 and 1.4 kb. The EcoRI fragment patterns of *S. pastorianus* and *S. bayanus* (lanes g and h) are in agreement with the summation of patterns II (b) and III (f).

#### 4. DISCUSSION

As summarized in Figure 9 the EcoRI restriction fragments of the *RDNI* gene encoding the cytosolic ribosomal RNA molecules allow to distinguish three forms. Form III is distinct from the previously mapped forms I and II (cf. Figure

1A) by having a fragment of 2.9 kb. Most likely, this fragment has originated from the X' fragment of form II by deletion of a segment in the 3' spacer region. Further work is required to demonstrate the presence of the EcoRI F and G fragments. Form III is also distinguished in the HindIII pattern by the presence of a third HindIII site in the gene. Form III allows to identify *S. bayanus*, *S. pastorianus* and *S. uvarum* which all are homozygous for this form.

By virtue of its ability to ferment glucose, sucrose, maltose and raffinose (33) and its inability to utilize galactose, *S. fermentati* superficially resembles *S. bayanus*. It is considered, however, to be most closely related to *S. rosei* (33). The *RDN1* of *S. fermentati* (3.2 and Figure 2) was found to have an organization which corresponds to that determined by VERBEET et al. (34) for *S. rosei*. This *RDN1* gene is about 500 bp smaller than forms I and III and characterized by having a single HindIII site. The "18S" ribosomal RNA probe used in the present investigation and the similar probe used by VERBEET et al. hybridizes in both strains to a 3.4 kb EcoRI restriction fragment indicating a similar overall organization. The probe with the total form II of *RDN1* hybridizes to *S. fermentati* EcoRI fragments of 3.4 kb and 2.8 kb. Fragments of the same size in *S. rosei* hybridize to the probe pMY60 containing a form I *RDN1* gene (34). In the latter paper a detailed comparison is made between form I and the *S. rosei* rDNA unit by heteroduplex mapping and partial nucleotide sequencing. The large difference between the *S. rosei* and the form I rDNA is entirely due to less sequence homology in the spacer regions. The analysis of the ribosomal RNA genes is in agreement with the observations by YARROW and NAKASE (38) that *S. rosei* and *S. fermentati* have the same sero-type composition and cell wall structures. They thus belong to the same species and are furthermore closely related to *Torulaspora delbrueckii* (38).

The chromosomal region containing the *HIS4* gene occurred in three different allelic forms, designated as patterns I, II and III. With one exception the lager strains (Table I) contained chromosomes with patterns I and II but were homozygous for form II of the *RDN1*. The exception is the strain formerly used by the Bavarian

brewery Pschorr which is homozygous for the *HIS4* pattern II. A hybrid between two spore clones of the Carlsberg lager strain BK2246 with such a homozygous pattern II has been produced and was recently successfully employed in production of beer (10, 25).

With the exception of strain BK3331, ale strains (Table II) contained form II of the *RDN1* gene and were homozygous for pattern I of the *HIS4* region. The ale strain BK3331 contained form I of the *RDN1* gene. Progeny analysis of the strains C82-FP23 and C82-FP25 are required to determine the nature of the populations observed in these two accessions.

Most useful is the combination of the *RDN1* and *HIS4* analysis to distinguish the strains listed in Table III. *S. odessa*, *S. diastaticus*, *S. ellipsoideus*, *S. italicus*, bakers yeast and a beer contaminant were homozygous for form I of *RDN1* and pattern I of *HIS4*. This genotype was also found in a strain isolated from Trappiste beer (C81-1537). *S. bayanus*, *S. pastorianus* and *S. uvarum* were characterized by homozygosity for the *RDN1* gene form III and the presence of pattern II of the *HIS4* gene. *S. pastorianus* and *S. bayanus* were heterozygous for patterns III and II of *HIS4*, while the two accessions of *S. uvarum* were homozygous for pattern III.

Previous studies on molecular cross hybridization of DNA from *S. uvarum* and *S. bayanus* have shown 95% homology between the DNA of the two species (28) while only 40% is found between *S. uvarum* and *S. cerevisiae* (1). *S. uvarum*, *S. bayanus* and *S. pastorianus* are thus closely related and taxonomical evaluation based on the capacity to ferment melibiose is misleading in this case.

The ribosomal RNA gene form III is also present in strain NCYC 396 considered as a type strain of *S. carlsbergensis*, but combined with pattern I and II of the *HIS4* region. It was noted that the *HIS4* region pattern II fragments of this strain hybridize less intensely with the *S. cerevisiae* probe than the corresponding fragments from the lager yeast strains listed in Table I. This type strain differs thus from *S. uvarum* in the *HIS4* region of chromosome III, it is similar to *S. uvarum* in the *RDN1* region of chromosome XII but differs in that respect from the lager yeast strains. All three groups of strains ferment



		HIS4 region				RDN1 gene			
						Form I	II	III	Band
kb	6.2	—	—	—					X'
	5.0	—	—	—	—	—	—	—	A
	4.1	—	—						B
	3.0	—	—	—	—	—	—	—	C
	2.65	—	—	—	—				
	1.4	—	—	—	—	—	—	—	D
	1.1	—	—	—	—	—	—	—	E
		—	—	—	—	—	—	—	F
		—	—	—	—	—	—	—	G
EcoRI	pattern	I	I+II	II	II+III	III			

Figure 9. Diagnostic key to EcoRI restriction endonuclease fragment patterns for the *HIS4* region and the *RDN1* gene. Fragment sizes are in kb (thousands of base pairs).

melibiose. Further studies which include additional genes and properties are necessary to evaluate the taxonomic status of strain NCYC 396 and the strain with the same genotype obtained from China (BK4515).

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