UC Irvine UC Irvine Previously Published Works

Title

Lessons from the Genome Sequence of Neurospora crassa: Tracing the Path from Genomic Blueprint to Multicellular Organism

Permalink https://escholarship.org/uc/item/76h9307w

Journal Microbiology and Molecular Biology Reviews, 68(1)

ISSN 1092-2172

Authors

Borkovich, KA Alex, LA Yarden, O <u>et al.</u>

Publication Date

2004-03-01

DOI

10.1128/MMBR.68.1.1-108.2004

License

<u>CC BY 4.0</u>

Peer reviewed

Lessons from the Genome Sequence of *Neurospora crassa*: Tracing the Path from Genomic Blueprint to Multicellular Organism

Katherine A. Borkovich,^{1*} Lisa A. Alex,² Oded Yarden,³ Michael Freitag,⁴ Gloria E. Turner,⁵ Nick D. Read,⁶ Stephan Seiler,⁷ Deborah Bell-Pedersen,⁸ John Paietta,⁹ Nora Plesofsky,¹⁰ Michael Plamann,¹¹ Marta Goodrich-Tanrikulu,¹² Ulrich Schulte,¹³ Gertrud Mannhaupt,¹⁴
Frank E. Nargang,¹⁵ Alan Radford,¹⁶ Claude Selitrennikoff,¹⁷ James E. Galagan,¹⁸ Jay C. Dunlap,¹⁹ Jennifer J. Loros,²⁰ David Catcheside,²¹ Hirokazu Inoue,²² Rodolfo Aramayo,⁸ Michael Polymenis,²³ Eric U. Selker,⁴ Matthew S. Sachs,²⁴ George A. Marzluf,²⁵ Ian Paulsen,²⁶ Rowland Davis,²⁷ Daniel J. Ebbole,²⁸ Alex Zelter,⁶ Eric R. Kalkman,⁶ Rebecca O'Rourke,²⁹ Frederick Bowring,²¹ Jane Yeadon,²¹ Chizu Ishii,²² Keiichiro Suzuki,²² Wataru Sakai,²² and Robert Pratt⁸

Department of Plant Pathology, University of California, Riverside, California 92521¹; Department of Chemistry, California State Polytechnic University, Pomona, California 91768²; Department of Plant Pathology and Microbiology, Faculty of Agricultural, Food and Environmental Quality Sciences, The Hebrew University of Jerusalem, Rehovot 76100, Israel³; Institute of Molecular Biology, University of Oregon, Eugene, Oregon 97403⁴; Department of Chemistry and Biochemistry, University of California, Los Angeles, California 90095⁵; Institute of Cell and Molecular Biology, University of Edinburgh, Edinburgh EH9 31H,⁶ and School of Biology, Leeds University, Leeds LS2 917,¹⁶ United Kingdom; Department of Molecular Microbiology & Genetics, Institute of Microbiology & Genetics, Georg-August-University, D-37077 Goettingen,⁷ Institute of Biochemistry, Heinrich Heine University, 40225 Dusseldorf,¹³ and Department of Genome Oriented Bioinformatics, Technical University of Munich, 85350 Freising-Weihenstephan,¹⁴ Germany; Department of Biology,⁸ Department of Biochemistry and Biophysics,²³ and Department of Plant Pathology and Microbiology,²⁸ Texas A&M University, College Station, Texas 77843; Department of Biochemistry and Molecular Biology, Wright State University, Dayton, Ohio 45435⁹; Department of Plant Biology, University of Minnesota, St. Paul, Minnesota 55108¹⁰; School of Biological Sciences, University of Missouri—Kansas City, Kansas City, Missouri 64110¹¹; Bio-Rad Laboratories, Inc., Hercules, California 94547¹²; Department of Biological Sciences, University of Alberta, Edmonton, Alberta T6G 2E9, Canada¹⁵; Department of Genetics¹⁹ and Department of Biochemistry,²⁰ Dartmouth Medical School, Hanover, New Hampshire 03755; School of Biological Sciences, Flinders University, Adelaide 5001, Australia²¹; Laboratory of Genetics, Department of Regulation Biology, Saitama University, Saitama City, Saitama 338-8570, Japan²²; Department of Environmental and Biomolecular Systems, School o

INTRODUCTION
Neurospora crassa: a Model Filamentous Fungus
Basic Features of the <i>Neurospora</i> Genome
Sequencing and assembly
Integration of the sequence with the genetic map7
Nuclear and mitochondrial genes7
Codon bias
Comparative multigene family and domain analysis8
CHROMATIN ASSEMBLY AND GENE REGULATION
Centromere Organization and Kinetochore Complexes
Organization of centromeres
Kinetochore complexes and motors that move chromosomes10
Chromosomes move through checkpoints10
Chromatin Structure and Gene Regulation
Nucleosome assembly and histone modification
(i) Nucleosome assembly and nucleosome spacing
(ii) Histone modifications
Chromatin assembly and remodeling14
(i) CAFs
(ii) CRFs
Transcription Factors
Zn(II) ₂ Cys _c fungal binuclear cluster family17

^{*} Corresponding author. Mailing address: Department of Plant Pathology, 2338 Webber Hall, 900 University Ave., University of California, Riverside, CA 92521. Phone: (909) 787-2753. Fax: (909) 787-4294. E-mail: Katherine.Borkovich@ucr.edu.

C2112 ZIIC Hilgers	······································
GATA factors	
bHLH transcription factors	21
B-ZIP transcription factors	21
Miscellaneous factors	
Translation Factors	
GENOME DEFENSE, DNA REPAIR, AND RECOMBINATION	
Genome Defense Mechanisms	
Heterochromatin silencing and DNA methylation	
RIP	
RNA-dependent silencing	
DNA Renair	
Photoreactivation	27
Fxcision	
(i) NFR	2.7
(ii) RFR	28
(II) DER	28
Postronlication repair	20
Charlengint control	30
Maiotia Decombination	30
Defore the DSP	
DED concretion	
DSB generation	
Removal of Spo11 protein from DNA	
Resection of ends	
Strand Invasion	
Synapsis and SC formation	
Regulation of crossover frequency	
Mismatch repair	
Resolution of recombination intermediates	
Nonhomologous end joining	
METABOLIC PROCESSES AND TRANSPORT	
Extracellular Digestion	
Glycosyl hydrolases	
Proteases	
Proteases Nucleases and phosphatases	
Proteases Nucleases and phosphatases Lipases	
Proteases Nucleases and phosphatases Lipases Transporters	
Proteases Nucleases and phosphatases Lipases Transporters Glycolysis, Fermentation, and Gluconeogenesis	
Proteases Nucleases and phosphatases Lipases Transporters Glycolysis, Fermentation, and Gluconeogenesis Glycolysis and the pentose phosphate cycle	
Proteases Nucleases and phosphatases Lipases Transporters Glycolysis, Fermentation, and Gluconeogenesis Glycolysis and the pentose phosphate cycle (i) Hexose phosphorylation	
Proteases Nucleases and phosphatases Lipases Transporters Glycolysis, Fermentation, and Gluconeogenesis Glycolysis and the pentose phosphate cycle (i) Hexose phosphorylation	
Proteases Nucleases and phosphatases Lipases Transporters Glycolysis, Fermentation, and Gluconeogenesis Glycolysis and the pentose phosphate cycle (i) Hexose phosphorylation (ii) EM glycolysis (iii) HM and ED glycolysis and the pentose phosphate cycle	
Proteases Nucleases and phosphatases Lipases Transporters Glycolysis, Fermentation, and Gluconeogenesis Glycolysis and the pentose phosphate cycle (i) Hexose phosphorylation (ii) EM glycolysis (iii) HM and ED glycolysis and the pentose phosphate cycle Alcoholic fermentation	
Proteases Nucleases and phosphatases Lipases Transporters Glycolysis, Fermentation, and Gluconeogenesis Glycolysis and the pentose phosphate cycle (i) Hexose phosphorylation (ii) EM glycolysis (iii) EM glycolysis (iii) HM and ED glycolysis and the pentose phosphate cycle Alcoholic fermentation	
Proteases Nucleases and phosphatases Lipases Transporters Glycolysis, Fermentation, and Gluconeogenesis Glycolysis and the pentose phosphate cycle (i) Hexose phosphorylation (ii) EM glycolysis (iii) EM glycolysis and the pentose phosphate cycle (iii) HM and ED glycolysis and the pentose phosphate cycle Alcoholic fermentation Gluconeogenesis Mitochondrion and Energy Metabolism	
Proteases Nucleases and phosphatases Lipases Transporters Glycolysis, Fermentation, and Gluconeogenesis Glycolysis and the pentose phosphate cycle (i) Hexose phosphorylation (ii) EM glycolysis (iii) EM glycolysis and the pentose phosphate cycle (iii) HM and ED glycolysis and the pentose phosphate cycle Alcoholic fermentation Gluconeogenesis Mitochondrion and Energy Metabolism Sulfur Metabolism	
Proteases Nucleases and phosphatases Lipases Transporters Glycolysis, Fermentation, and Gluconeogenesis Glycolysis and the pentose phosphate cycle (i) Hexose phosphorylation (ii) EM glycolysis	
Proteases Nucleases and phosphatases Lipases Transporters Glycolysis, Fermentation, and Gluconeogenesis Glycolysis and the pentose phosphate cycle (i) Hexose phosphorylation	
Proteases Nucleases and phosphatases Lipases Transporters Glycolysis, Fermentation, and Gluconeogenesis Glycolysis and the pentose phosphate cycle	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
Proteases Nucleases and phosphatases Lipases Transporters Glycolysis, Fermentation, and Gluconeogenesis Glycolysis and the pentose phosphate cycle (i) Hexose phosphorylation	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
Proteases	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
Proteases	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
Proteases	38 38 39 39 40 40 40 40 40 40 40 40 40 40 41 42 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 44 47 49 50 51 51
Proteases	$\begin{array}{c} & & & & & & & & & & & & & & & & & & &$
Proteases	$\begin{array}{c} & & & & & & & & & & & & & & & & & & &$
Proteases	38 38 39 39 40 40 40 40 40 40 40 40 40 40 40 41 43 43 43 43 43 43 43 43 43 43 43 43 43 43 43 44 47 49 50 51 51 52 52 52 52 55
Proteases	38 38 39 39 40 40 40 40 40 40 40 40 40 40 41 42 43 43 43 43 43 44 47 49 50 51 51 52 52 55 55
Proteases	$\begin{array}{c} & & & & & & & & & & & & & & & & & & &$
Proteases	$\begin{array}{c} & & & & & & & & & & & & & & & & & & &$
Proteases	$\begin{array}{c} & & & & & & & & & & & & & & & & & & &$
Proteases	$\begin{array}{c} & & & & & & & & & & & & & & & & & & &$
Proteases	$\begin{array}{c} & & & & & & & & & & & & & & & & & & &$
Proteases	$\begin{array}{c} & & & & & & & & & & & & & & & & & & &$
Proteases	$\begin{array}{c} & & & & & & & & & & & & & & & & & & &$
Proteases	$\begin{array}{c} & & & & & & & & & & & & & & & & & & &$

MAPKs	64
Calcium signaling	65
Protein phosphatases	67
Mammalian signaling proteins not found in <i>Neurospora</i>	67
Photobiology and Circadian Rhythms	67
Heat Shock and Stress Responses	69
GROWTH AND REPRODUCTION	74
Cell Wall	74
Glucan synthases	74
(i) (1,3)β-Glucan synthesis	74
(ii) (1,6)β-Glucan synthesis	75
Chitin substrate synthesis—the Leloir pathway	75
Cell wall precursors	75
Chitin synthases	75
Hyphal Morphogenesis	76
Generation of hyphal polarity	77
(i) Proteins important for cell polarity development	77
(ii) Rho-type GTPases as key regulators of polarity	78
Cytoskeleton and motor proteins	80
(i) Structural components	80
(ii) Kinesins	80
(iii) Myosins	82
(iv) Dynein	82
Cyclin/CDK Machinery	82
Asexual and Sexual Sporulation	83
Macroconidiation	83
Meiosis and the sexual cycle	84
FUNGAL PATHOGENESIS AND HUMAN DISEASE	86
Relationship to Animal and Plant Pathogens	86
Animal pathogens	86
Plant pathogens	87
Human Disease Genes	
PERSPECTIVES AND FUTURE DIRECTIONS	
ACKNOWLEDGMENTS	
REFERENCES	91

INTRODUCTION

Neurospora crassa: a Model Filamentous Fungus

The kingdom Fungi contains an estimated 1.5 million species, the majority of which are filamentous (321). The impact of this group of organisms on human affairs and the ecosystem rivals that of plants and bacteria. Fungi are probably the most biotechnologically useful group of organisms and are used to synthesize a wide range of economically important compounds, enzymes, and secondary metabolites, including antibiotics and other pharmaceuticals (841). Fungi are the primary agents of decay on this planet (225) and play critical ecological roles in nutrient recycling. This biodegradation activity can also be deleterious, leading to decomposition of economically useful products, including building timber, man-made materials, and food (841).

Fungi are often found in association with other organisms. Numerous diseases of humans and other animals are caused by fungi, and the incidence of life-threatening fungal infections is on the rise, in parallel with the increased number of immunocompromised patients (453). Fungi are the most important group of plant pathogens, causing significant and often devastating losses in crop yield worldwide (8). Mycorrhizal fungi form symbiotic associations with the roots of higher plants (744) and effectively determine what type of plant ecosystem develops (see, e.g., reference 389).

The fungi have played a major role in the progress of biochemistry, genetics, and molecular biology. George W. Beadle and Edward L. Tatum (55) defined the role of genes in metabolism, and this led quickly to the mid-century revolution in genetics. Their work took advantage of the filamentous ascomycete Neurospora crassa (hereafter referred to as Neurospora), which was first described in 1843 as the causative agent of an orange mold infestation in French bakeries; (Fig. 1) (607, 610). Neurospora was later domesticated as an experimental organism by Bernard O. Dodge (725) and Carl C. Lindegren (see, e.g., reference 476). Beadle and Tatum sought an organism displaying Mendelian genetics that could be grown on simple media and might display additional nutrient requirements arising by mutation. Their success emboldened others to use bacteria, algae, and other fungi in similar studies. Together with the elucidation of the structure of DNA in 1953, molecular biology as we know it was born.

Neurospora soom became a popular experimental model organism (185). Diverse research programs centered on *Neurospora* have ranged from formal, population, and molecular genetics, biochemistry, physiology, and molecular cell biology to more recent studies of development, photobiology, circadian rhythms, gene silencing, ecology, and evolution. Substantial genetic and molecular information has been obtained about species differences and intraspecific variation, building on the efforts of David Perkins, who has sampled natural iso-



rapidly. c. Branching filament, about 150×. g". Spore treated successively, under the microscope, with a dilute solution of potassium hydroxide, and aqueous alcoholic solution of iodine, then with gradually more concentrated solutions of sulfuric acids. This acid, which separates parts of the cellulose envelope that contains less nitrogenous substance, results in a blue color turning to purple, which is characteristic of the state intermediate between cellulose and dextrin. i. Normal vegetative growth as seen with the naked eye; well-developed, especially under conditions FIG. 1. Plate from the first published scientific study of *Neurospora*. Plate 1 from Payen (607). The following translated legend for the portions of the figure labeled a, a', b', c, g'', i, and k' is taken from reference 610. "a. Colonies of the red-orange fungus Oidium aurantiacum as they appear to the naked eye in the cavities of infected bread. a'. A similar colony cut in two, showing in the red area a thick layer composed of innumerable small spores formed at the end of radiating filaments. The latter are yellowish white. b. Similar colonies that have grown up completely in the dark, with the result that the red color has not developed. b'. One of the colonies in b seen after exposure to light for one hour. Color begins to appear and then pigmentation progresses high humidity. k'. Termini of well developed filaments, showing spores and young cells." zlates from all over the world (809). The legacy of 70 years of intense research with this organism continues to be driven by a large and interactive research community that has also served to draw together a wider group of scientists working with other filamentous fungi.

One of the attractive features of Neurospora as a model organism is its complex yet genetically and biochemically tractable life cycle (Fig. 2). Neurospora is multicellular and produces at least 28 morphologically distinct cell types (82), many of which are derived from hyphae. Neurospora vegetative hyphae are tip-growing cellular elements that undergo regular branching (294, 798, 800, 812) and are multinucleate. These hyphae contain incomplete cross walls (septa) (315) that allow the movement of organelles between compartments. Frequent fusion among hyphal filaments produces a complex hyphal network (the mycelium) (336) and promotes the formation of heterokaryons in which multiple genomes can contribute to the metabolism of a single mycelium. Specialized aerial hyphae are differentiated from vegetative hyphae in response to nutrient deprivation, desiccation, or various stresses, and these form chains of asexual spores (the multinucleate macroconidia) for dispersal (752). The timing of macroconidiation is controlled by a circadian rhythm, which in turn is modulated by exposure to blue light. Another type of asexual spore, the uninucleate microconidium, is differentiated from microconidiophores or directly from the vegetative hypha (82, 495, 752). Limiting nitrogen induces a type of hyphal aggregation that leads to generation of multicellular female sexual organs (protoperithecia) (564, 642). Mating is accomplished by chemotropic growth of a specialized female hypha from the protoperithecium toward the male cell (typically a conidium) in a process involving pheromones (81). Fertilization and meiosis result in development of the female structure into a beaked fruiting body (the perithecium) within which asci, each containing eight ordered sexual spores (ascospores), are formed (638).

The genome sequence of Neurospora was recently reported (269, 498). Here we provide a more detailed analysis, with the annotation of approximately 1,100 genes, or more than 10% of the total predicted in the genome. Several themes emerge from this study. First, the multicellular Neurospora possesses a large number of genes without homologues in Saccharomyces cerevisiae, suggesting that Neurospora will be a better model for higher eukaryotes in many aspects of cell biology, including multicellularity (579). Among the unshared genomic equipment is an expanded group of sugar transporters, transcription factors, and environmental sensing pathways, plus a diversified metabolic machinery. Second, Neurospora displays a number of gene-silencing mechanisms acting in the sexual or the vegetative phase of the life cycle. Best studied is repeat-induced point mutation (RIP), an effective defense against duplicated sequences such as those arising from the multiplication of transposons (see "Genome defense, DNA repair, and recombination" below) (709). However, RIP appears to impose constraints on gene and genome evolution, raising the question of whether Neurospora currently is able to utilize gene duplication as a means of gene diversification. Finally, Neurospora, a generalist species in its life cycle, genetic system, and growth requirements, provides a basis for comparison with the highly diversified plant pathogens and other specialized fungi of narrow habitat. For example, the number of



FIG. 2. Life cycle of *Neurospora*. "Depending on surrounding conditions, the vegetative mycelium can undergo the asexual sporulation processes of macroconidiation and microconidiation. It can enter the sexual cycle by forming protoperithecia; which, upon fertilization, can initiate development leading to the production of meiotically-derived ascospores." Reprinted from reference 751 with permission from the publisher.

homologues it shares with pathogens offers a starting point for study of evolutionary gene recruitment in these specialized organisms. Conversely, genes found in pathogenic fungi but absent from *Neurospora* provide potential targets to exploit for development of antifungal drugs and fungicides.

Basic Features of the Neurospora Genome

Sequencing and assembly. The Neurospora genome was sequenced using a whole-genome shotgun strategy (a summary of its general features is given in Table 1). Paired end reads were acquired from a variety of clone types, including plasmids, fosmids, cosmids, bacterial artificial chromosomes (BACs), and jumping clones. In all, greater than 20-fold sequence coverage and greater than 98-fold physical converage were generated. These data were assembled into a draft sequence by using the Arachne whole-genome assembler (50, 378). The resulting assembly consists of 958 sequence contigs (contiguous stretches of sequence derived from reads that overlap with high sequence similarity) with a total length of 38.6 Mb and an average N50 length of 114 kb (meaning that 50% of all bases are contained in contigs of at least 114 kb). Contigs were assembled into 163 scaffolds (sets of contigs that are ordered and oriented with respect to each other by using paired read information) with an N50 length of 1.6 Mb.

The long-range continuity of the assembly was confirmed by comparisons with previously described BAC physical maps for linkage groups II and V (11), in which only one discrepancy

was noted. Confirmation of long-range continuity was also provided by comparisons with the *Neurospora* genetic map (see below). The assembly also has high accuracy, with 99.5% of the sequence having Arachne quality scores of \geq 30. A comparison with 17 Mb of finished BAC sequence (http://mips.gsf.de/proj /Neurospora/) confirmed the sequence accuracy. Only 12 discrepancies were identified in this comparison. Five lie at the end of contigs and are most probably caused by misaligned or low-quality terminal reads. Four are short insertions or deletions, ranging from 9 to 559 bp. The remaining three discrepancies appear to be instances in which the finished sequence does not correctly represent the genome, owing to chimerism in the BACs and in which the whole-genome assembly is correct.

The total genome size can be approximated from the draft assembly by estimating the size of gaps between contigs and scaffolds. The size of gaps between adjacent contigs in a scaffold can be derived from the size of clones spanning the gap. When these gap sizes are included, the total physical length of all scaffolds is estimated to be 39.9 Mb. The size of gaps between scaffolds is more difficult to estimate since spanning clones are not available. In addition, these gaps include difficult-to-sequence regions of the genome including the ribosomal DNA (rDNA) repeats, centromeres, and telomeres. A total of \sim 1.7 Mb of additional sequence (251) is probably accounted for by these regions. Based on these considerations, the genome size is estimated to be 41 Mb. The most recent

TABLE 1. Features of the Neurospora genome

Feature	Value	No. (%) of genes with BLASTP hits ^a	No. (%) of genes with BBH hits ^b
General			
Size (bp)	38,044,345		
No. of chromosomes %G+C	7 50		
Protein-coding genes	10,082		
Protein-coding genes >100 aa ^c	9,200		
Introns	17,118		
tRNA genes	424		
5S rRNA	74		
Percent coding	44		
Avg gene size (bp)	1,673 (481 aa)		
Avg intron size (bp)	135.4		
Avg intergenic distance (bp)	1,953		
Predicted protein-coding sequences			
Identified by similarity to known sequences	1,336 (13%)		
Conserved hypothetical proteins	4,606 (46%)		
Predicted proteins (no simi- larity to known sequences)	4,140 (41%)		
Organism comparison			
Schizosaccharomyces pombe		3,883 (39)	2,225 (22)
Saccharomyces cerevisiae		3,722 (37)	2,209 (22)
Drosophila melanogaster		3,120 (31)	1,619 (16)
Caenorhabditis elegans		2,986 (30)	1,521 (15)
Arabidopsis thaliana		3,291 (33)	1,756 (17)
Homo sapiens		3,379 (34)	1,350 (13)

^{*a*} At e < 1e-5.

^b BBH, best bidirectional hit.

^c aa, amino acid.

estimate of the genome size based on pulsed-field gel electrophoresis of intact *Neurospora* chromosomes is 42.9 Mb (www .fgsc.net/fgn39/oneline.html). The size predicted from the sequence (41 Mb) is well within the limits of resolution for pulsed-field gel electrophoresis measurements of such large molecules.

Integration of the sequence with the genetic map. Approximately 1,000 genetic markers exist for *Neurospora*. The majority have been ordered on the genetic maps for the seven linkage groups and are described in a recent compendium (612). The *Neurospora* assembly was correlated with the genetic maps by using a subset of 252 markers for which there is sequence in *Neurospora* (or other closely related fungi). The marker sequences were located on the current assembly by using BLASTN and filtering for unique high-quality alignments (http://www.ogi.edu/satacad/ase [269]). The 243 (96% of the total) markers that aligned were then used to place contigs and scaffolds on the physical map, according to the genetic marker order. In all, 95% of the assembly was assigned to a linkage group; 85% of this sequence was further ordered and oriented within a linkage group.

Only a handful of discrepancies were noted between the physical and genetic maps. There were three cases where gene order differs between the two maps. Five markers were located in more than one contig, indicating places where the assembly failed to merge contigs. Twelve markers failed to be located within the physical map, indicating sequence gaps within the current assembly. Finally, nine scaffolds contained markers on different linkage groups, indicating either misplaced markers on the genetic map or contigs incorrectly linked within a scaffold.

Nuclear and mitochondrial genes. An automated annotation of the Neurospora draft genome sequence was performed by the Whitehead Institute Center for Genome Research (WICGR) by using the Calhoun annotation and analysis system. A combination of three gene prediction algorithms (FGenesH, FGenesH+, and Genewise) was combined with available protein homology to predict protein-coding genes. Gene predictions were compared with BLAT alignments of available expressed sequence tags (ESTs) (63, 565, 900) to assess accuracy. A total of 10,082 protein-coding genes were predicted. Eliminating proteins shorter than 100 amino acids that lack protein or EST similarity reduces this number to 9,200. This number of genes is within the range of 9,200 to 13,000 estimated by previous authors (56, 409, 442, 565). An additional 26 protein-coding genes reside in the mitochondrial genome (see "Metabolic processes and transport" below).

Consistent with the greater biological complexity of filamentous fungi compared to both fission and budding yeast, Neurospora possesses nearly twice as many genes as Schizosaccharomyces pombe (\sim 4,800) and S. cerevisiae (\sim 6,300). Neurospora contains almost as many as genes as Drosophila melanogaster (\sim 14,300), despite the relative developmental complexity of the latter. In addition, 41% of the predicted Neurospora proteins do not have significant similarity to known or predicted proteins in other organisms and 57% do not have good matches to proteins in either S. cerevisiae or S. pombe (269). The Neurospora gene complement also displays greater structure complexity than those of the two yeasts. Neurospora genes possess a predicted 17,118 introns (1.7 introns per gene), compared to roughly 286 (0.04 intron per gene) and 4435 (0.95 intron per gene) in S. cerevisiae and S. pombe, respectively. However, as with the yeasts and other simple eukaryotes, Neurospora introns do appear to be biased toward the 5' regions of genes.

A total of 413 tRNA genes were identified using tRNAscan (487), including 234 (57%) with introns. Of this number, 396 are predicted to decode all standard amino acids and one could potentially decode UAG termination codons. Ten tRNA pseudogenes were identified, two of which were inferred to be mutated by RIP. An additional six tRNAs were predicted with undetermined specificity; one of these was inferred to be a relic of RIP.

All annotation data are available at the WICGR *Neurospora crassa* website (http://www-genome.wi.mit.edu/annotation/). In addition, a manually curated annotation of the *Neurospora* gene set is available at the Munich Information Center for Protein Sequences (MIPS) *Neurospora crassa* database (MNCDB; http://mips.gsf.de/proj/neurospora) (498). At present, MNCDB contains 8,500 *Neurospora* proteins; this number is expected to increase as manual gene prediction and annotation progress. MIPS protein codes were chosen according to the cosmid, BAC, and DNA shotgun contigs from which they were derived. Linkage was established with their respective counterparts in the WICGR database that were identified using automated gene prediction tools. The proteins in the WICGR database may differ from those in MNCDB, due to manual

MICROBIOL. MOL. BIOL. REV.

correction, but the proteins in the different databases are linked as long as partial matches are found.

Codon bias. The mRNA expression level is influenced by synonymous codon usage in a number of organisms. In particular, increasing codon bias is correlated with greater expression level in Escherichia coli (362), S. cerevisiae (156, 266, 606 [although see also reference 305]), as well as Caenorhabditis elegans, D. melanogaster, and Arabidopsis thaliana (208). Correspondence between tRNA gene copy number and codon usage has also been demonstrated for highly expressed genes in S. cerevisiae (816) and E. coli (362). It has been proposed, based on these and other data, that codon bias reflects coadaptation between codon usage and tRNA abundance in order to maximize the efficiency of protein translation for highly expressed genes. However, in mammalian genomes, codon bias has been attributed to regional variations in genomic G+C content (i.e., isochores). In support of this, it has been shown that in mammals the G+C content of regions flanking genes (816) and the GC content of introns (207) covary with the G+C content in the third position of codons.

Although Neurospora genes display significant variation in codon bias, the determinants of this bias are not known for filamentous fungi. To determine whether this variation might reflect mutational selection for translational efficiency, EST sequences from a number of previously characterized libraries were used to estimate relative transcript levels (63, 565, 900). In particular, a count of the number of distinct EST clones that align with a given gene (or flanking region) was used as an estimate of the relative transcript level for that gene. Two different measures of codon bias were used: the codon bias index (CBI) (65) and the effective number of codons (Nc). CBI is a measure of the amount of bias toward a particular set of favored codons, with a large CBI indicating greater bias; for this analysis, the set of favored codons from reference 481 was used. Nc is a measure of codon bias away from uniform codon usage, with a smaller Nc indicating greater bias.

A statistically significant correspondence between estimated transcript level and codon bias was detected using both CBI (Spearman rank correlation coefficient R = 0.30, n = 10,082, P < 1e-197) and Nc (Spearman rank correlation coefficient R = -0.25, n = 10,082, P < 1e-138). Furthermore, a significant correspondence between estimated transcript level and the degree of correlation of codon usage with synonymous tRNA copy number was detected. In other words, more highly expressed genes showed a strong tendency to display a codon usage that was more closely aligned with a synonymous tRNA gene copy number. A significant correspondence between codon third-position G+C content in genes and estimated transcript levels was also detected; however, there was no significant relationship between intron G+C content and estimated transcript levels. These data suggest that, similar to the situation in S. cerevisiae, codon usage and tRNA abundance in Neurospora have coevolved to maximize the efficiency of protein translation for highly expressed genes.

Comparative multigene family and domain analysis. Despite the presence of RIP, *Neurospora* possesses 527 multigene families, including 118 families expanded relative to their counterparts in *S. cerevisiae*. In addition, *Neurospora* possesses numerous Interpro protein domains that display expansions in number relative to other sequenced eukaryotes. Particularly

surprising is the abundance of cytochrome P450 domains, which are numerous in plants and in *Neurospora* but very scarce in both *S. cerevisiae* and *S. pombe*. The cytochrome P450 enzyme domain, including the E-class P450 group 1 domain and E-class P450 group IV domain subclasses, are represented by 38 proteins in *Neurospora*. In contrast, *S. cerevisiae* and *S. pombe* contain only two to four proteins with these domains. Accounting for genome size, this represents a six- to eightfold increase in genes with these domains in *Neurospora*. Cytochrome P450s are known for playing roles in both detoxification and secondary metabolism, and the implications of their high representation in *Neurospora* have been discussed previously (269).

Other domains abundant in *Neurospora* include the zinc finger C2H2-type domain, the *S*-adenosylmethionine (SAM) binding motif domain, the short-chain dehydrogenase/reductase (SDR) superfamily domain, and the flavin adenine dinucleotide-dependent pyridine nucleotide-disulfide oxidoreductase domain. Interestingly, a number of domains involved in signaling appear underrepresented compared to other fungi and plants. These include the eukaryotic, serine/threonine, and protein tyrosine kinase domains. Other underrepresented domains include certain helicases, RNA binding protein motifs, and the AAA-ATPase superfamily domain.

In the following sections, different variations of the BLAST program (17) were used to search DNA or protein databases using DNA or protein sequences. The resulting e value is dependent on the database size, and various databases are of different sizes and many are increasing in size over time. Hence, the magnitude of e should be treated as an indication and not as absolute measure of the similarity between two sequences.

CHROMATIN ASSEMBLY AND GENE REGULATION

Centromere Organization and Kinetochore Complexes

Centromeres and centromere-associated proteins are necessary to accomplish the movement of chromosomes on microtubule spindles during cell division (for a recent review, see reference 154). Conventional wisdom holds that a genetic locus and the information it conveys are defined by its DNA sequence. Most eukaryotic centromeres, however, run counter to this concept. In fact, they represent one of the best examples of how cells make use of so-called "junk DNA", because centromeric DNA in most eukaryotes is an assembly of satellite DNA and transposons or transposon relics. While the function of the small centromere of the budding yeast S. cerevisiae can be disrupted by point mutations, the much larger centromeres of other eukaryotes, from the fission yeast S. pombe to metazoans, appear to be functionally redundant (497). The hunt for centromere sequence elements has been supplanted by the realization that redundant, nonconserved DNA segments can act as a "scaffold" for the assembly of a specialized centromere nucleosome complex. The defining unit in this complex is a variant of histone H3, encoded by CENP-A in mammals, cid in Drosophila, CSE4 in S. cerevisiae, and cnp1 in S. pombe (331). Cse4-containing nucleosomes are found predominantly or exclusively at the core of centromeres, surrounded by stretches of transcriptionally inactive heterochromatin (520). In mammals

LG	Scaffold	Size (kb)	Cen contigs	Cen (kb)	Nearest locus/marker
Ι	54	213	522–528	210	Uncertain
II L	93	68	680	28	vma, 41-kDa subunit
II R	20	566	303-304	215	
III L	22	565	323–329	271	<i>acr-2</i> (2 kb)
IV R	29	418	373–374	20	Uncertain, pyr-1 (60 kb)
IV L	17	667	273–277	200	On 3.273: NCU04963.1 (1 kb) On 3.277: NCU04984.1 (1 kb)
VL	82	110	628	11	Uncertain, NCU09669.1; ccg-8
V R	15	769	246-254	300	NCU04676.1
VI L	24	534	335–348	125	NCU06013.1
VI R	12	862	206-208	8	Uncertain, NCU03846.1 (rib-1?)
VII C	6	1,266	119–130	336	CenVII (contig 129); hH2A (contig 119)

TABLE 2. Contigs with centromeric sequences ordered by linkage group^{*a*}

^{*a*} Putative centromeric sequences were detected by BLAST searches with *Tcen*, *Tgl* (121), and *Tad* (122) elements. The edge of the centromere (Cen) is defined as the beginning of uninterrupted AT-rich repeat segments. One large scaffold containing most of the centromere was found for each linkage group (LG). Contigs for LG IIL, IVC, VR, and VI were mapped based on sequencing data available at http://www.mips.biochem.mpg.de/proj/neurospora/ and ordered cosmid libraries at http://gene.genetics.uga.edu/. Additional putative core centromeric pieces are contained on scaffolds 96, 98, 101, 103, 111, and 112, but cannot be mapped using available data. Based on assembly 3, the LG IV centromere may be contained in its entirety on scaffold 17.

and *Drosophila*, centromeric DNA associated with CENP-A and Cid, respectively, is interspersed with histone H3-containing nucleosomes (86). This assembly forms multiple foci for kinetochore subunits to create microtubule attachment points, an arrangement also predicted for *Neurospora*, where centromeres are rich in inactive transposons (121) and where a CENP-A homologue, hH3v, has been identified (322). The centromere binding proteins and kinetochore complexes previously isolated in *S. cerevisiae*, *S. pombe*, and animal systems have few counterparts in *Neurospora* (with the exception of the Ndc80 complex), lending support to the idea that centromeric regions and their associated complexes undergo "accelerated evolution" (330).

Organization of centromeres. S. cerevisiae has the simplest eukaryotic centromere known, only ~125 bp of DNA associated with a single nucleosome. This short region is divided into three centromere DNA elements (CDEs) which are conserved on all 16 chromosomes and serve as binding sites for the sequence-specific DNA binding protein Cbf1 and the essential CBF3 complex (457). The sequence of CDE II is not conserved, but the length (\sim 80 bp) and A+T content (\sim 90%) is similar at all yeast centromeres. CDE III is associated with the histone H3 variant Cse4/CENP-A and the kinetochore chromatin binding protein Mif2/CENP-C. In contrast to S. cerevisiae, S. pombe centromeres are much larger (40 to 100 kb) and are composed of two inverted repeats surrounding a nonconserved core. The inner inverted repeats and the core sequence are associated with Cnp1/CENP-A nucleosomes and the Ctf19 homologues Mis6 and Mis12 (603, 772). The flanking regions are assembled into heterochromatin in part by the histone methyltransferase, Clr4, and the heterochromatin protein Swi6 (see "Genome defense, DNA repair, and recombination" below). Drosophila centromeres are large (400 to 500 kb) and are composed of 5-bp satellite sequences interspersed with transposons or transposon relics, while human centromeres are 0.5 to 5 Mb long and are homogenously composed of 171-bp long α -satellites (for a review, see reference 154). In most animals and in plants, the repeat sequences of the satellite arrays are not conserved but the array repeats usually approximate the length of a nucleosome repeat of DNA sequence (330).

The seven centromeres of Neurospora remain largely uncharacterized, even after the cloning of Cen VII (137) and detailed analysis of a 17-kb segment (121). It is clear, however, that *Neurospora* centromeres are large (~ 200 to 400 kb) and AT rich, like those in *Drosophila* (Table 2). As in flies, they appear to consist of an accumulation of complete or fragmented and rearranged transposon relics, in particular the gypsy-type Tgl1 or copia-type Tcen retrotransposon relics, the LINE-like element Tad, and a homolog of the S. cerevisiae Ty3 transposon, Tgl2 (121). All such transposon relics have been inactivated by the genome defense mechanism of RIP (see "RIP" below), which introduces CG-to-TA transition mutations and thus renders the DNA highly AT rich. Micro- and minisatellites and homopolymeric stretches can be identified in centromeric regions (498). No specific accumulation of tRNA genes has been noted close to the centromeres. Assembly 3 of the Neurospora genome sequence contains the sequence of most of the centromeres of all chromosomes, but some of the putative centromeric regions cannot yet be assigned to supercontigs on specific chromosomes. Similarly, the German sequencing project (498) has not yielded complete sequence information for the centromeres of linkage groups (LG) II and V. Curiously, no significant increase in the ratio of physical to genetic distance between known markers has been observed in regions near the centromeres of LG II and V (498), in contrast to that previously reported for LG III (183).

Comparison of centromeric DNA sequences in eukaryotes suggests that epigenetic factors, rather than simply DNA sequence, determine the activity of the locus. Both centromeres and the specialized histone H3, CENP-A, evolve quickly and show no obvious sequence conservation across species or even at different centromeres of the same species (330). DNA se-

10 BORKOVICH ET AL.

Duratain	I	Match found by BLAST					
Tiotem	Locus (gene)	Best match ^c	S. cerevisiae	S. pombe	Animal ^a	Plant ^b	
HsCENP-B	0996.1	A. thaliana	No match	8e-39	1e-25	1e-40	
ScCbf1	8999.1	P. carinii, 2e-33	2e-20	4e-29	2e-7	3e-4	
HsCENP-C/ScMif2	1635.1	S. pombe	2e-6	2e-7	1e-6	No match	
ScCBF3d (=Skp1)	8991.1 (scon-3)	E. nidulans, 5e-44	4e-35	1e-45	7e-36	5e-34	
ScNdc80/HsHec1	3899.1	S. pombe	1e-52	2e-56	2e-27	3e-9	
HsNuf2	6568.1	S. pombe	0.013	6e-21	3e-5	0.32	
ScSpc24	5312.1	S. pombe	2e-4	6e-6	No match	No match	
ScSpc25	0965.1	S. pombe	2e-6	1e-6	4e-5	1.1	
ScDam1	6878.1	S. pombe	0.11	8e-6	No match	No match	
ScOkp1	0367.1	S. cerevisiae	0.16	No match	No match	No match	
ScMtw1/SpMis12	9120.1	S. cerevisiae	1e-4	0.002	No match	No match	
SpMis6	4131.1	S. pombe	No match	3e-32	3e-27	No match	
ScCh14	3537.1	S. cerevisiae	2e-6	No match	No match	No match	
ScSli15/HsINCE NP	5211.1	S. cerevisiae	0.025	No match	No match	No match	
ScBir1/HsSurvivin	6621.1	S. pombe	No match	3e-10	1e-8	No match	
ScIp11/HsAurora B	0108.1	S. pombe	1e-64	1e-102	1e-87	4e-80	

TABLE 3. Putative Neurospora homologues of proteins involved in centromere binding and kinetochore assembly

^a Caenorhabditis elegans, Drosophila melanogaster, Anopheles gambiae, Mus musculus, or Homo sapiens (Hs).

^b Arabidopsis thaliana or Oryza sativa.

^c Species listed as best match are Saccharomyces cerevisiae (Sc), Schizosaccharomyces pombe (Sp), Pneumocystis carinii, and Emericella (Aspergillus) nidulans.

quences characteristic of centromeres are by themselves unable to direct centromere function, and in rearranged chromosomes where bona fide centromeric DNA sequences have been deleted, novel sequences can aquire centromere function (154).

Kinetochore complexes and motors that move chromosomes. To move chromosomes during cell division, centromeres are attached to spindle microtubules via the kinetochore and various motor complexes. In S. cerevisiae, the Cse4p/ CENP-A nucleosome is bound by a centromere protein clamp, consisting of a homodimer of Cbf1p, the essential CBF3 complex, and Mif2/CENP-C (274, 519). Interestingly, Neurospora has putative homologues to both, presumably functionally equivalent, S. cerevisiae helix-loop-helix Cbf1p and the animaltype CENP-B proteins (Table 3). Curiously, alignments of putative Neurospora CENP-B homologues with S. pombe proteins (Cbh1, Cbh2, and Abp1) involved in centromere binding revealed that the Neurospora CENP-B homologues contain RIPtype mutations, including numerous nonsense mutations. Because CENP-B and Cbh1-like proteins are also related to the Drosophila Pogo and human Tigger transposons, most of the Neurospora Cbh1 homologues have been considered to be transposon relics (717). CBF3 is an octameric complex composed of four subunits (four Ndc10p, two Ctf13p, and one each of Cep3p and Skp1p). Neurospora has one Skp1p homolog, SCON-3, which is part of an SCF complex (E3 ubiquitin ligase see "Sulfur metabolism" below) involved in sulfur regulation (741), but there are no matches to the other three essential CBF3 subunits.

In *S. cerevisiae*, the Okp1/Ctf3 (577) and Ndc80 (379, 853) complexes connect the core centromere to distal spindle components and serve as kinetochore "glue," while the Dam/Duo complex is important for direct microtubule binding (141, 142, 380). *Neurospora* has all four previously identified components of the Ndc80 complex, and all of them are most closely related to the *S. pombe* homologues. *Neurospora* contains only three poor matches to proteins of the *S. cerevisiae* Okp1 complex, however (Table 3). Similarly, there is one Dam1p homologue

but no good match to the other subunits (i.e., Dad1p-4p, Duo1p, Spc19p and Spc34p, Ask1p) of the yeast Dam/Duo complex in *Neurospora*. The activity of Dam1p is itself regulated by the conserved Ipl1p/Sli15p (Aurora B/INCENP) kinase complex (351, 613). It appears that the Ndc80 complex and proteins important for regulation of microtubule capture and proper segregation (Dam1p and its kinase, Ipl1p) are evolutionary conserved while functional homologues of the CBF3, Okp1p and Dam/Duo complexes are possibly subject to the same "accelerated evolution" that has been suggested for the CENP-A protein family and the centromeric DNA substrate (330).

In mammals, the kinesin CENP-E, cytoplasmic dynein, microtubule tracking proteins, and disassemblases are involved in chromosome movement (see "Growth and Reproduction" below); whereas in yeast, dynein positions the spindle but is not involved in actual chromosome movement (154). It remains to be seen whether dynein is involved in chromosome segregation in *Neurospora;* nevertheless, effects similar to those in mammals have been observed in *Tetrahymena* and *Drosophila* (154). It is likely that both active motor movement and microtubule flux contribute to anaphase movements of chromosomes.

Chromosomes move through checkpoints. The mitotic (or spindle assembly) checkpoint blocks the entry into anaphase until the two kinetochores of duplicated chromatid pairs have attached to spindle microtubules. This ensures accurate segregation (655). It appears that a combination of unattached kinetochores coupled to lack of tension acting on both kinetochores of a chromatid pair causes the block (154). Genetic dissection in *S. cerevisiae* has identified seven components of the mitotic checkpoint (552), all of which appear conserved in *Neurospora*. The molecular interactions between kinetochores and all checkpoint proteins are not established, but it appears that Mad2p and Cdc20p play central roles in signaling the unattached kinetochore, either directly or through a signal-amplifying cascade which inactivates or sequesters Cdc20p. Loss or reduced levels of checkpoint proteins in metazoans

Duratain	I ()	Match found by BLAST ^e					
Protein	Locus (gene)	Best match	S. cerevisiae	S. pombe	Animal ^a	Plant ^b	
Core histones							
Histone H2A	2437.1 (hH2A)	P. anserina, 3e-52	1e-48	1e-50	3e-45	4e-41	
Histone H2B	2435.1 (hH2B)	P. anserina, 2e-32	2e-32	1e-32	6e-41	3e-29	
Histone H3	1635.1 (hH3)	A. nidulans, 7e-53	3e-50	1e-51	8e-53	7e-50	
Histone H4-1	1634.1 (hH4-1)	A. nidulans, 1e-31	9e-30	3e-30	7e-32	3e-29	
Histone H4-2	212.1 (hH4-2)	Same as H4-1					
H2Az	5347.1 (hH2Az)	S. pombe	3e-37	1e-42	1e-35	1e-32	
H3v/ScCse4	0145.1 (hH3v)	S. pombe	3e-28	2e-30	3e-24	4e-24	
H4v	4338.1 (hH4v)	P. polycephalum, 3e-18	2e-16	4e-16	1e-16	1e-15	
Linker histones							
Histone H1	6863.1 (hH1)	A. nidulans, 2e-14	1e-9	No match	6e-5	No match	
Histone-like proteins							
HMG1.2/ScNhp6	9995.1	V. faba	1e-17	2e-10	2e-11	1e-21	
HMG2A	2819.1	M. musculus	4e-9	1e-5	2e-13	3e-13	
HMG4	2695.1	P. anserina, 3e-11	0.008	No match	3e-5	2e-6	
MATA-3	1960.1 (matA-3)	S. macrospora, 1e-37	No match	1e-4	No match	No match	
MATa-1	No ID (mata-1)	S. macrospora, 1e-135	No match	2e-14	2e-11	No match	
S. pombe Ste11-like	2326.1	N. crassa, 1e-10, 9387.1	No match	0.011	No match	No match	
S. pombe Ste11-like	9387.1	N. crassa, 9e-11, 2326.1	No match	1e-7	4e-6	No match	
ScRox1/Rfg1/S RY	3481.1	C. albicans, 4e-14	2e-13	6e-12	4e-9	No match	
TBF (TATA-bind.)	2017.1	S. pombe	2e-15	6e-29	1e-18	4e-16	
HAP (CAAT-bind)	9248.1	H. jecorina, 3e-49	2e-26	1e-30	3e-31	4e-26	
CHRAC17/POL IÍ	3073.1	S. pombe	4e-4	1e-11	2e-6	6e-5	
Polyamine oxidase with HMG box	9120.1	H. sapiens	No match	8e-25	4e-69	9e-54	

TABLE 4. Neurospora histones and histone variants involved in nucleosome assembly and nucleosome spacing

^a Caenorhabditis elegans, Drosophila melanogaster, Strongylocentrotus purpuratus, Mus musculus, or Homo sapiens.

^b Arabidopsis thaliana, Vicia faba, or Oryza sativa.

^c Histones and histone-like proteins are short; therefore BLAST scores are comparatively low and e values are high. Species listed as best matches are *Podospora* anserina, Aspergillus nidulans, Physarum polycephalum, Vicia faba, Sordaria macrospora, Candida albicans, and Hypocrea jecorina.

causes chromosome missegregation, tumorigenesis, and apoptosis (154).

Chromatin Structure and Gene Regulation

Nucleosome assembly and histone modification. (i) Nucleosome assembly and nucleosome spacing. (a) Core histones. Nucleosomes are assembled into an octamer from dimers of the core histones H2A, H2AB, H3, and H4. Except for H4, which is encoded by two unlinked genes, hH4-1 and hH4-2, all *Neurospora* core histones are encoded by single genes (Table 4) (322, 860). This is similar to the situation in other fungi but is in contrast to that in plants and metazoans, which have numerous histone genes (322). The distribution and small number of histone genes, as well as the presence of introns within the histone genes of filamentous ascomycetes other than yeasts, may reflect the operation of the genome surveillance systems RIP and MIP (322). Introns are absent from the histone genes of most other eukaryotes.

(b) Core histone variants. Three core histone variants are present in the Neurospora genome (Table 4) (322). The H2Az variant is closely related to S. cerevisiae Htz1p and S. pombe Pht1, proteins involved in maintaining chromatin integrity, recruiting polymerase II to specific promoters, and protection from telomeric silencing (4, 130, 374, 521). The H3v variant (see "Centromere organization and kinetochore complexes" above) is a homolog of Cse4p from S. cerevisiae (143, 520, 759), CENP-A from humans (21, 827, 895), and cid from D. melanogaster (85, 331). There are no good matches to the H4 variant (H4v) in other fungi; this may be a pseudogene (322).

(c) Linker histones. Similar to other fungi (403), Neurospora has only one histone H1 gene (Table 4) (242). Studies with Neurospora, S. cerevisiae, Tetrahymena, Aspergillus nidulans, and Ascobolus immersus revealed that H1 is not essential in any of these organisms (44, 252, 640, 729, 817) but, rather, is implicated in the regulation of nitrogen and carbon metabolism (242, 329, 727).

(d) Histone fold motifs and HMG proteins. Short proteins with histone fold motifs are involved in transcriptional regulation in all eukaryotes (271, 639). The histone fold motif in CBFD/NF-YB/HMF is similar to domains found in archaebacteria (608). Three predicted *Neurospora* proteins containing histone folds are related to general transcription factors (Table 4): TATA binding factor (TBF) (NCU02017.1); CHRAC17 (NCU03073.1), a putative subunit of RNA polymerase II with homology to a subunit of the CHRAC chromatin remodeling factor from *Drosophila*; and HAP (NCU09248.1), a homologue of the CCAAT binding proteins Hap3p from *S. cerevisiae*, HAP-C from *A. nidulans*, Php3 from *S. pombe*, and NF-YB from humans.

Two sex-determining region, Y chromosome (SRY)-related high-mobility group (HMG) transcription factors, MATA-3 and MATa-1, have been characterized in *Neurospora* (233, 615). Three additional proteins have homology to mating peptides or carry a sterile alpha motif and an HMG-1-like box (Table 4). NCU03481.1 is related to a repressor of hypoxic genes (Rox1p) in *S. cerevisiae* and a virulence factor (Rfg1) in *Candida albicans*. NCU09387.1 and NCU02326.1 are related to *S. pombe* Ste11. Interestingly, the Ste11-related proteins from *Neurospora* are each other's closest homologues, which is unusual in *Neurospora*. HMG-like proteins are typically short, such as HMG1.2 (NCU09995.1; related to *S. cerevisiae* recombination proteins Nhp6A and Nhp6B) and HMG2A (NCU02819). The most unusual HMG protein predicted in *Neurospora* (NCU09120.1) has a HMG box at the C terminus and is most similar to human polyamine oxidases.

(ii) Histone modifications. Core and linker histones are extensively modified at the posttranscriptional level (826). Histone modifications have been studied both for their effects on the regulation of specific genes and for their importance to global regulatory phenomena. Recent work has uncovered an epigenetic "histone code" (384, 750, 761, 811) involved in transcriptional regulation (for reviews, see references 67 and 104) and other DNA transactions (for recent reviews, see references 245, 361, 691, and 810). Histone residues can be actively modified by acetylation and deacetylation of lysine; methylation of lysine and arginine; phosphorylation and dephosphorylation of serine, threonine, and histidine; ADP-ribosylation of glutamic acid, and ubiquitylation of the entire proteins (384, 826). Because the interrelationships between histone modifications are essential for the formation and maintenance of silent chromatin states (384, 445, 559), genes involved in Neurospora histone modification are described below (see "Genome defense, DNA repair, and recombination"). The combinatorial possibilities of the histone code are staggering, even without considering the facts that different modification states are possible on the same residue and that lysine and arginine can be mono-, di-, or trimethylated (see, e.g., reference 778).

(a) HATs. Histone acetyltransferases (HATs) transfer acetyl groups from acetyl coenzyme A acetyl-CoA to lysines, most often located within the amino-terminal tail of the core histones. Neurospora has representatives of many of the HATs involved in transcriptional activation and gene silencing (e.g., TAFII250, Gcn5p, Sas2p, Sas3p, Esa1p, and Elp3p [J. Dobosy and E. Selker, unpublished results]) but lacks homologues to S. cerevisiae Hpa1p and metazoan CBP and SRC (Table 5). The homologue of S. cerevisiae Gcn5p lacks a locus identifier because the predicted protein lies at the beginning of contig 3.38 (Table 5). Neurospora has three N-terminal acetyltransferases that correspond to the budding yeast proteins Nat1p, Nat3p, and Mak3p, all of which are involved in cell cycle control (Table 5) (415). Interestingly, three of the putative GNAT/RIM1-type acetyltransferases have bacterial proteins as their closest relatives. The range of substrates of the predicted Neurospora acetyltransferases remains unknown.

(b) HDACs. Histone deacetylases (HDACs) remove acetyl groups from lysine residues, e.g., histone tails. They are separated into three families (190). Neurospora has 11 predicted proteins related to known or putative HDACs in two of the families (Table 5). Four proteins (HDA-1 to HDA-4) have homologs in the Rpd3/Hos/Hda group (Dobosy and Selker, unpublished), compared to five such proteins in budding yeast and seven and nine in animals and plants, respectively. Seven predicted proteins (NST-1 to NST-7) are related to the NAD-dependent HDAC Sir2 and related proteins ("sirtuins" [G. Kothe, M. Freitag, and E. Selker, unpublished results]). Neurospora has homologues of all four S. cerevisiae sirtuins as well as three additional sirtuins most closely related to those in animals. Arabidopsis has just one sirtuin but contains a class of

unrelated HDACs, called HD2 type, which is not found in *Neurospora* or animals (593).

(c) HMTs. Histone methyltransferases (HMTs) add methyl groups supplied by SAM to lysines (Lys) or arginines (Arg) on core histones. Neurospora has a single member of all known HMT subfamilies (Table 5), whereas metazoans, plants, and in some cases S. cerevisiae and S. pombe have multiple proteins for each. Neurospora is predicted to have nine proteins (SET-1 to SET-8 and DIM-5; M. Freitag, K. Adhvaryu, and E. Selker, unpublished data) with SET domains, a motif first found in the Drosophila Su(var) 3-9, Enhancer of zeste, and Trithorax proteins. SET domains are characteristic of lysine protein methyltransferases (for a review, see reference 445), although not all SET domain proteins are HMTs. Neurospora lacks some SET proteins identified in S. cerevisiae (e.g., Set5p and Set6p) and in humans but has two proteins (SET-6 and SET-8) that are either novel or restricted to only a subset of fungi. Both proteins align well by their Zn finger and Jumanji (JmJi) domains, but the pairing of these domains with the SET motif is rare. Like its homologues in S. pombe, animals, and plants (641, 690), Neurospora DIM-5 is a histone H3 Lys9 HMT (776, 778). Budding yeast Set1p and Set2p methylate histone H3 on Lys4 (106, 553, 660) and Lys36 (762), respectively, and Neurospora has striking homologues (SET-1 and SET-2) to these two proteins. Neurospora SET-3 is related to Drosophila ASH1, which methylates histone H4 on Lys20 and histone H3 on Lys4 and Lys9 (59), and SET-7 is related to E(z), which methylates histone H3 Lys9 and Lys27 (546). Human G9A also methylates histone H3 on Lys9 and Lys27 (770, 771), and Neurospora SET-5 appears related to the G9A subgroup.

Neurospora has three putative arginine methyltransferases (PRMTs) (for reviews, see references 432 and 511) that are homologous to PRMT1, PRMT3 and PRMT5 of humans, respectively (Table 5). Interestingly, *Neurospora* does not have a recognizable homologue of either PRMT2 or PRMT4 (CARM1). CARM1 methylates histone H3 Arg2, Arg17, and Arg26 (51, 180). PRMT3 has not yet been shown to methylate histones (249, 781), but PRMT1 methylates histone H4 Arg3 in vivo (696, 843) and PRMT5 and its homologues can methylate both histone H2A and H4 in vitro (103, 227). PRMT5 is a homolog of *S. pombe* Skb1 and *S. cerevisiae* Hsl7p, which methylate the protein kinases Shk1 (43) and Swe1p (150), respectively. The *Neurospora* homolog is called PP-2 (P. Bobrowicz and D. Ebbole, unpublished data).

Neurospora has one homologue of *S. cerevisiae* Dot1p (M. Freitag, C. Matsen, J. Murphy, G. Kothe and E. Selker, unpublished data) (Table 5), an HMT which methylates Lys79 within the globular domain of histone H3 and which is important in telomeric silencing in *S. cerevisiae* (740) and humans (232, 568).

(d) Histone kinases. Like histone acetylation and methylation, histone phosphorylation has been intensively studied, and found to be important for chromosome codensation, the signaling of active versus silent chromatin states, transcription, regulated cell death, and DNA repair (145). All core and linker histone H1 can be phosphorylated in vitro (826), and all histone kinases that act on histones in vivo are involved in the control of cell cycle progression. Histone H3 Ser10 can be phosphorylated by at least two different kinases in *S. cerevisiae*, Snf1p (483) and Ip11p/Aurora B (172, 351). The *S. cerevisiae*

TABLE 5.	Neurospora	homologues of	proteins	involved	in	histone	modifications ⁶
----------	------------	---------------	----------	----------	----	---------	----------------------------

		Match found by BLAST					
Protein	Locus (gene)	Best match ^d	S. cerevisiae	S. pombe	Animal ^b	Plant ^c	
Histone acetyltransferases							
ScHat1	6472.1 (hat-1)	S. cerevisiae	4e-28	4e-14	4e-22	3e-15	
HsTAFII250	2556.1 (hat-2)	S. pombe	No match	1e-109	1e-49	3e-27	
ScElp3	1229.1 (elp-3)	S cerevisiae	0.0	0.0	0.0	0.0	
ScGen5	No ID (nof_{-1})	V linobica: 1e-138	1e-135	1e-116	7e-78	6e-79	
Section S	$5218 \pm (hg + 4)$	L. upolylicu, 10-156	10-100	10-110	0-06	2 00	
SCESAL	3218.1 (<i>nui-4</i>)	S. cerevisiae	16-122	10-122	96-90	26-90	
ScSas2	2249.1 (hat-5)	S. cerevisiae	2e-34	1e-25	3e-34	2e-34	
ScSas3	4782.1 (hat-6)	S. pombe	7e-78	1e-95	2e-84	1e-79	
Protein acetyltransferases							
SnArd1/ScNot3	1276.1	S nomba	10.26	20 11	69.40	10.32	
SpArd1/Scivals	2417.1	D more 2 75	1- 22	4 - 25	6-41	1- 22	
SCMak5	2417.1	P. anserina; se-75	16-33	46-35	66-41	16-35	
SpAts1	7589.1	P. aeruginosa; 8e-25	No match	7e-22	2e-14	4e-20	
GNAT-type	9914.1	B. iaponicum; 2e-5	No match	No match	No match	No match	
GNAT-type	4583.1	<i>B. cereus</i> ; 1e-13	No match	No match	No match	No match	
Deacetylases							
ScHda1/SpClr3	1525.1 (hda-1)	S. pombe	1e-137	1e-145	4e-80	4e-67	
ScHos2	$2795 1 (hda_2)$	A nidulans: 1e-143	1e-104	1e-106	1e-100	2e-95	
SeD nd2	08241 (hdg 2)	1. manuaris, 10 145	10 161	10 100	10 146	10 127	
SCKPUS	0.00000000000000000000000000000000000	A. manuns, 10-178	10-101	1C-144	10-140	16-12/	
SCH083	7018.1 (<i>naa-4</i>)	S. cerevisiae	96-85	NO IIIS	36-18	86-25	
ScHst1/HsSirtuin1	4737.1 (<i>nst-1</i>)	S. pombe	6e-80	5e-88	7e-62	No match	
ScHst2/HsSirtuin2	0523.1 (nst-2)	S. pombe	5e-56	7e-71	1e-65	No match	
ScHst4	3059.1 (nst-3)	S. pombe	5e-61	4e-73	5e-21	No match	
ScHst3/HsSirtuin3	4859.1 (nst-4)	S. cerevisiae	7e-28	3e-15	2e-6	0.001	
HsSirtuin4	0203.1 (nst-5)	A thaliana	No match	No match	5e-35	8e-42	
HeSirtuin5	5073.1 (nst 6)	M musculus	No match	No match	10.35	No match	
	5975.1 (<i>hst-0</i>)	M. musculus	NO match	NO match	10-33	1 40	
HsSirtuin/	/624.1 (<i>nst-</i> /)	A. thaliana	No match	No match	3e-43	1e-49	
Protein methyltransferases							
Lysine, SET domain							
SpClr4	4402.1 (dim-5)	S. pombe	No match	2e-43	7e-36	2e-26	
ScSet1	1206.1 (set-1)	S cerevisiae	3e-64	2e-53	3e-53	7e-45	
ScSet2	0260.1 (set 2)	S. pomba	10.104	10 116	30 50	30 13	
	0209.1 (sel-2)	S. pombe	1 20 ((2)	2 25	36-30	36-43	
DmAsn1	1932.1 (set-3)	R. norvegicus	1e-29 (set 2)	3e-25	20-37	2e-31	
ScSet3/Set4	4389.1 (set-4)	H. sapiens	9e-10	8e-8	9e-13	0.027	
HsG9a	6119.1 (set-5)	C. elegans	No match	No match	8e-12	No match	
NcSET-6	9495.1 (set-6)	S. pombe	2e-6	7e-36	9e-6	3e-6	
DmEn(z)/AtMedea	7496.1 (set-7)	M. musculus	No match	No match	4e-29	1e-25	
NcSET-8	1973.1 (set-8)	H. sapiens	5e-4	No match	4e-13	4e-6	
Lysine, non-SET							
ScDot1	6266.1	S. cerevisiae	3e-38	No match	5e-22	No match	
Arginine						• • • •	
HsPRM11/ScHmt1	7459.1 (prm-1)	S. pombe	1e-111	1e-116	1e-100	3e-98	
HsPRMT3	1669.1 (prm-3)	S. pombe	4e-62	6e-86	6e-67	2e-62	
HsPRMT5/ScHs17/SpSkb1	1613.1 (pp-2)	S. pombe	4e-69	1e-107	5e-86	3e-77	
Kinases							
ScSnf1	4566.1	F. oxysporum: 0	1e-134	1e-123	1e-103	1e-103	
ScSnf4	1471 1	K lactis: 2e-94	4e-92	4e-88	7e-48	8e-74	
SeCol93	3037 1	S correction	10 22	50 23	50 10	20 27	
SUJAIOS	3937.1	S. cerevisiue	10-27	30-23	JC-19	20-10	
Scipii/HsAurora B	0108.1	S. pombe	1e-64	1e-102	16-8/	4e-80	
<i>En</i> Nım-A	3187.1 (nim-1)	E. nidulans; 1e-172	6e-68 (Kin3)	3e-66 (Fin1)	8e-63 (Nek2)	1e-36	
Ubiquitylases	0731 1 (marco 9)	N haquatococce to 75	20.63	49.64	30.57	30 52	
SCRAUO	9751.1 (<i>mus-</i> ð)	w. naematococca; 4e-75	20-03	40-04	30-37	30-33	
ADP-Ribosylases	8852 1 (nam)	1 thaliana	No motob	No motoh	30.60	30 67	
LUCIU	0032.1 (parp)	A. Inaliana	ino maten	ino maten	36-00	36-07	

^{*a*} Histone phosphatases and histone kinases are listed in Tables 48 and 56, respectively. Note: Based on sequence comparisons with fungal Snf1p homologues, the Snf1 locus is misannotated at the WICGR but correct at MNCDB (CAD70761). ^{*b*} Caenorhabditis elegans, Drosophila melanogaster, Strongylocentrotus purpuratus, Xenopus laevis, Mus musculus, or Homo sapiens. ^{*c*} Arabidopsis thaliana, Vicia faba, or Oryza sativa. ^{*d*} Species listed as best matches are Podospora anserina, Aspergillus nidulans, Physarum polycephalum, Vicia faba, Sordaria macrospora, Fusarium oxysporum, Yarrowia lipolytica, Candida albicans, Nectria haematococca, and Hypocrea jecornia.

Ductoin	I ()		Match found by BLAST					
Floteni Locus (gene)		Best match ^c	S. cerevisiae	S. pombe	Animal ^a	Plant ^b		
CAF-1								
ScCac1 (Hsp150)	4198.1 (cac-1)	S. pombe	0.026	0.002	No match	No match		
ScCac2 (Hsp60)	8357.1 (cac-2)	S. pombe	1e-39	1e-52	7e-37	1e-31		
ScCac3 (Hsp48)	6679.1 (cac-3)	D. rerio; 1e-113	2e-45	1e-105	1e-113	1e-105		
RCAF								
ScAsf1	9436.1 (asf-1)	S. cerevisiae	2e-58	3e-52	8e-48	5e-43		
NAP								
NcNAF-1	1438.1 (naf-1)	S. pombe	6e-55	1e-67	5e-39	6e-34		
NcNAF-2	3769.1 (naf-2)	S. pombe	No match	5e-12	2e-8	3e-5		
Accessory factors								
<i>Hs</i> PCNA	9239.1 (pcn)	S. pombe	2e-62	7e-96	5e-76	3e-80		

TABLE 6. Neurospora homologues of proteins involved in nucleosome and chromatin assembly

^a Drosophila melanogaster, Anopheles gambiae, Caenorhabditis elegans, Strongylocentrotus purpuratus, Danio rerio, Mus musculus, or Homo sapiens.

^b Arabidopsis thaliana, Daucus carota, Vicia faba, or Oryza sativa.

^c Species listed as best match is Danio rerio.

Snf1p kinase is a heterotrimer, composed of the Snf1p α subunit, the Snf4p γ subunit and three different β subunits, Sip1p, Sip2p, or Gal83p (474). *Neurospora* has homologues to Snf1p and Snf4p but has only one β subunit, most closely related to Gal83p (Table 5). Phosphorylation of H3 Ser10p is a prerequisite of acetylation at histone H3 Lys14 and is usually required for gene activation (482, 572). Like H3, the centromeric H3 variant CENP-A can be phosphorylated, albeit at Ser7 (895). In addition to homologues of Snf1p and Ip1p/Aurora B, filamentous fungi have a cell cycle kinase, Nim-A (*Neurospora* NIM-1 [635]), that phosphorylates histone H3 Ser10 (192).

In animals, histone H3 Thr13 can also be phosphorylated by the Dlk/Zip kinase in vivo (629). This modification is found on both H3 and CENP-A at assembled centromeres, suggesting that Thr11 phosphorylation rather than Ser10 phosphorylation may be involved in the maintenance of silent chromatin and kinetochore attachment. Curiously, these threonine DAP (death-associated protein kinase)-like enzymes seem restricted to animals, since no homologue has been found in any fungus, including *Neurospora*.

Candidate histone phosphatases from *Neurospora* (PPP-1, PPH-1, and PZL-1) have been isolated by biochemical means, and their genes have been identified (see "Environmental sensing" below).

(e) Histone ubiquitylases. Our understanding of histone ubiquitylation is still fragmentary. Ubiquitylated histones H2A and H2B are the most abundant ubiquitylated proteins in eukaryotes (381). Ubiquitin is linked to histone H2A Lys119 (91) and H2B Lys120 or Lys123 in animals and *S. cerevisiae*, respectively (658, 789). All histone H2 variants known in animals can be ubiquitylated (381). *Neurospora* has several predicted subunits of ubiquitin-activating (E1) and ubiquitin-ligating (E2) proteins that are predicted to be involved in histone ubiquitylation. For example, the *Neurospora mus-8* gene encodes a Rad6p-like H2B ubiquitin-ligase (Table 5) (658, 749). This enzyme affects gene silencing via histone H3 methylation states in *S. cerevisiae*, an example of "trans-tail" regulation of histone modifications (765).

(f) Histone ADP-ribosylases. Of the histone modifications, ribosylation is currently the least well understood. One class of

enzymes thought to be involved in ribosylation and histone turnover are the poly(ADP-ribose) polymerases (PARPs) (16). In contrast to mammals, which have multiple PARPs, *Neurospora* has a single *parp* gene (G. Kothe and E. Selker, unpublished results; (Table 5). Whether any of the small GTP binding proteins with ADP ribosylation activity have effects on histones in vivo is unknown.

Chromatin assembly and remodeling. (i) CAFs. Eukaryotes use histone chaperones or chromatin assembly factors such as CAF and RCAF to guide histones prior to assembly into nucleosomes (for a review, see reference 517). As in other eukaryotes, Neurospora CAF-1 is predicted to be composed of three subunits (CAC-1 to CAC-3) (M. Freitag and E. Selker, unpublished data) (Table 6). Disruption mutants with mutations in the gene encoding the largest, least well conserved subunit, cac-1, are viable (Freitag and Selker, unpublished); this is similar to the situation in S. cerevisiae, where neither CAF nor the RCAF subunit of the antisilencing factor (Asf1p) are essential. These findings suggest the presence of additional chromatin assembly factors. CAC-2 and CAC-3 contain WD40 domains and are conserved among all eukaryotes studied. Neurospora CAC-3 is more closely related to retinoblastoma binding protein 48 (RBBP4; p48) from mammals than it is to S. cerevisiae Cac3p (Msi1p/Ira1p). In mammals, RBBP4/p48 also associates with HDAC complexes (653, 840). Neurospora contains one homologue of S. cerevisiae Asf1p (Table 6), a component of an alternative chromatin assembly factor first described in Drosophila as RCAF (813). asf1 mutants exhibit more drastic silencing effects in S. cerevisiae than do cac1 mutants, and the severity is further enhanced in double-knockout strains (740, 813). Drosophila ASF1 cooperates with the brahma chromatin-remodeling complex (see below), and mutation of ASF1 results in derepression of heterochromatic regions (541). These results suggest that both CAF and RCAF play a role in heterochromatin silencing in eukaryotes.

Neurospora has two predicted proteins with domains characteristic of nucleosome assembly proteins (NAPs) (Table 6). All other eukaryotes with sequenced genomes have at least two paralogues. NAF-1 (nucleosome assembly factor 1) is a canonical NAP, which probably functions as a chaperone for the

D ()		Match found by BLAST						
Protein	NCU no. (gene)	Best match ^c	S. cerevisiae	S. pombe	Animal ^a	Plant ^b		
INO80/DOMINO								
ScSwr1	9993.1 (crf1-1)	S. cerevisiae	0.0	0.0	1e-121	0.00		
ScIno80	8919.1 (crf2-1)	S. cerevisiae	0.0	1e-146	1e-117	1e-164		
Swi2/Snf2/BRM								
ScSwi2/Snf2	6488.1 (crf3-1)	S. pombe	0.0	0.0	0.0	00		
ISWI								
DmISWI	3875.1 (crf4-1)	S. cerevisiae	0.0	0.0	0.0	0.0		
ScIoc4	2684.1 (crf4-3)	S. pombe	0.035	6e-6	No match	No match		
ScItc1	164.1 (<i>crf4-2</i>)	S. cerevisiae	7e-21	No match	4e-6	5e-7		
Lsh/ScYFK8								
HsLSH	6306.1 (crf5-1)	A. fumigatus; 9e-95	1e-59	No match	2e-64	2e-56		
Mi-2								
ScChd1/HsCHD2	3060.1 (crf6-1)	S. pombe	0.0	0.0	1e-173	1e-160		
HsCHD3/Mi-2 AtGYMNOS	1406.1 (crf7-1)	S. pombe	1e-96 (Chd1)	1e-155	1e-111	1e-115		
TAF172/Mot1-like								
HsTAF172	7556.1 (crf8-1)	S. pombe	1e-69	0.0	0.0	0.0		
Ris1								
ScRis1	0631.1 (crf9-1)	S. pombe	6e-70	1e-118	2e-47	2e-59		
ScRis1-like/Dm LODESTAR-like	4786.1	S. pombe	5e-43 (Ris1)	4e-70	4e-50	3e-50		
Role in transcription?								
POLII term. factor	7358.1	S. pombe	No match	1e-13	5e-9	3e-12		
POLII term. factor	4445.1	H. sapiens	4e-21 (Ris1)	1e-39 (Rad8, Rhp16)	1e-39	9e-37 (Rad5)		
HsETL-like	7975.1	A. thaliana	3e-66 (Rad16)	1e-72 (Rad8)	2e-85	1e-110		
HsETL-like	5246.1	A. thaliana	No match	4e-57 (Rhp16)	9e-73	1e-84		
Viral activator	2913.1	C. fumiferana MNPV: 3e-42	No match	4e-31 (Rhp16)	9e-73	1e-33		
Activator protein	4424.1^{d}	A. thaliana	No match	No match	3e-24	6e-32		
Fun30/ETL1								
ScFun30/HsETL1	9106.1 (crf10-1)	S. pombe	1e-104	1e-122	1e-105	1e-74		

TABLE 7. Neurospora homologues of predicted chromatin-remodeling factors

^a Drosophila melanogaster, Anopheles gambiae, Mus musculus, or Homo sapiens.

^b Arabidopsis thaliana, Zea mays, or Oryza sativa.

^c Species listed as best match are Aspergillus fumigatus and Choristoneura fumiferana multicapsid nucleopolyhedrovirus (MNPV).

^d NCU04424.1 may be misannotated (the 5' region is too short), and a homologue of A. nidulans sonB, a nucleoporin (NCU04784.1) is annotated as containing a SNF2 domain.

histone H2A-H2B tetramer (139, 368). NAF-2 is related to phosphatase 2A inhibitor 2 (471), which is involved in chromatin decondensation in *S. pombe* and humans and may help balance competing kinase and phosphatase activities at histone H3 Ser10 (572).

Like *S. cerevisiae* and *S. pombe, Neurospora* has a single homologue of the proliferating-cell nuclear antigen protein (PCNA) (Table 6). The homotrimeric PCNA complex serves to mark newly replicated DNA at the replication foci to which the CAF complex localizes. PCNA is involved in many activities involving DNA, including replication, repair, and silencing (517).

(ii) CRFs. Chromatin-remodelling factors (CRFs) use the energy generated by their ATPase subunits to remove or position nucleosomes relative to the DNA substrate (for recent reviews, see references 58, 240, and 452). *Neurospora* has 24 predicted proteins related to the SWI/SNF ATPase/helicase domain (Table 7). In organisms in which they have been studied, these proteins have been implicated in chromatin remod-

eling, DNA repair, and activation of transcription and are typically found in large complexes. Ten of the predicted *Neurospora* proteins are homologues of previously identified CRFs. The remaining 14 may be involved in ATPase-dependent repair processes and transcription. One of these, MUS-25, is the previously identified *Neurospora* RAD54 homolog (312). The *Neurospora* complement of SWI/SNF ATPases is a subset of those found in *S. cerevisiae*, *S. pombe*, *Arabidopsis*, and animals.

The SWI/SNF domain is homologous to viral and prokaryotic helicases. Therefore, all proteins in this group were previously called helicases (217, 290), even though helicase activity had not been demonstrated. Several members of this family have now been shown to function in large chromatin complexes such as ATPases without apparent helicase activity (58). In combination, CRFs may be expected to be involved in regulating the activity of many, if not most, genes by either activation or repression. All known CRFs have at least two components in vivo, but some ATPase subunits can remodel chromatin in vitro alone, albeit with altered specificity (210).

CRFs were categorized by their putative conserved ATPase subunits alone. Usually, ATPase subunits of various CRFs from different organisms are more closely related to each other than to different CRF ATPases from the same organism. Compared to CRFs from the yeasts, *Drosophila*, and human, relatively few homologues to non-ATPase subunits can be identified in *Neurospora* (e.g., to the yeast RSC, SNF2, and ISW complexes, the *Drosophila* CHRAC and NURF complexes, and the human BRG and BRM complexes). While there are fewer CRF ATPases in the *Neurospora* genome than in the *S. cerevisiae* genome, it appears likely that one ATPase subunit may associate with different accessory proteins to form specific complexes, analogous to the situation in *Drosophila*, where ISWI is present in three complexes: ACF, NURF, and CHRAC (452).

Similar to *S. cerevisiae*, *Neurospora* has two ATPase homologues in the INO80 group of CRFs, CRF1-1 and CRF2-1 (Table 7). Ino80p is the only CRF that has helicase activity (728). CRF1-1 is a homologue of Swr1p, while CRF2-1 is a homolog of Ino80p. Both are related to DOMINO CRFs of *Drosophila* (667).

Arabidopsis, metazoans, S. pombe, and S. cerevisiae have several CRFs that are generally associated with global gene activation. In Drosophila, for example, the distribution of RNA polymerase II and the BRAHMA complex largely coincide (27). Like humans (BRG and BRM), S. cerevisiae has two such complexes (SNF2 and RSC) with the bromodomain ATPases Snf2p and Sth1p, respectively (392, 542). Strikingly, Neurospora has only one predicted bromodomain ATPase, CRF3-1 (Table 7). Similar to their homologues, Neurospora CRF3-1 and two polybromodomain proteins (PBD-1 and PBD-2 [Table 7]) predicted to be in RSC- or SNF2-like complexes contain bromodomains, which bind to acetylated lysine residues (581).

Most eukaryotes have more than one heterochromatinassociated complex of the Imitation Switch (ISWI) type. In S. cerevisiae, at least four ISWI complexes exist, two each with specific ATPase, Isw1p and Isw2p, whereas in Drosophila, the single ISWI protein is present in three separate complexes: ACF, NURF, and CHRAC (for a review, see reference 452). This may be similar to the situation in Neurospora, where only one ISWI homologue, CRF4-1, exists but where subunits related to a yeast ISW2 component and Drosophila CHRAC subunits (Table 4, "Histone-like proteins") can be identified (Table 7). CRF4-2, for example, is predicted to be a DNA binding protein with a DDT domain, similar to Drosophila ACF1, which is present in both ACF and CHRAC. As in S. cerevisiae, and in contrast to Drosophila, disruption of crf4-1 is not lethal in Neurospora (Freitag and Selker, unpublished). ISWI complexes are generally thought to serve as global repressors of gene expression because they are colocalized with silent heterochromatin in Drosophila polytene chromosomes and excluded from transcriptionally active regions (193).

Neurospora CRF5-1, a relatively short (882-amino-acid) CRF ATPase (Table 7), is related to similar-length proteins involved in DNA methylation: DDM1 from *Arabidopsis* (382) and LSH from humans (189). DDM1-like CRF subunits may represent a subgroup of the ISWI CRFs, since CRF5-1 is closely related to CRF4-1. *Arabidopsis* and animals have two chromodomain CRF ATPase groups with at least two representatives in each group, while *S. cerevisiae* apparently has a single Mi-2-like chromodomain protein, Chd1p (796). *Neurospora* has one representative for each of the two groups (Table 7). CRF6-1 is a homologue of *S. pombe* Hrp1 (888) and Hrp3 (377), *Drosophila* KISMET (179), and mammalian CHD2 (859), while the ATPase domain of CRF7-1 is more closely related to human Mi-2/CHD3 (794, 840) and *Arabidopsis* GYMNOS/PICKLE (223). In animals, Mi-2-containing NuRD complexes can bind to methylated DNA via proteins with methyl binding domains (560, 657, 839). While the ATPase domain of CRF7-1 most closely matches Mi-2, the chromodomain is a poor match to the chromodomain consensus sequence.

Three additional predicted ATPases may be subunits of CRFs involved in transcription in *Neurospora* (Table 7). CRF8-1, CRF9-1, and CRF10-1 are similar to Mot1p, Ris1p, and Fun30p from *S. cerevisiae*, respectively. Mot1p-like proteins repress genes by interaction with TATA binding factor but can also be involved in gene activation, presumably by chromatin remodeling during transcription preinitiation (178, 543). The *Neurospora* protein is more closely related to plant and animal homologues of Mot1, such as human TAF172. Ris1p has a "role in silencing" (286). Two RING finger domain-containing Ris1p homologs can also be found in *S. pombe* and *Arabidopsis*. Fun30p may be important for chromosome stability (580).

Seven poorly characterized yet closely related predicted *Neurospora* ATPases may be involved in transcription (Table 7). These proteins are similar to human RNA polymerase termination factor, plant transcription factors, and viral activator proteins. An additional seven putative helicases or SWI/SNF ATPases are known or predicted to be involved in DNA repair in *Neurospora* (Table 8). *Neurospora* has a single Rad5p homologue and two members each of the Rad16p, Rad26p and Rad54p groups (discussed in more detail in "Genome defense, DNA repair, and recombination below). Human Rad26 homologues have chromatin-remodeling activity (151). One of the RAD54 proteins has homology to ATRX, a protein involved in DNA methylation (281).

Transcription Factors

With the availability of genome sequences from many diverse organisms, it is now possible to conduct comparative genomics on proteins required for gene transcription and regulation. This analysis will contribute to our understanding of promoter evolution by providing information about the regulation of organismal complexity. A recent review addressing the gene number conundrum, which is exemplified by the fact that a simple nematode worm, C. elegans, has $\sim 20,000$ genes whereas a more complex organism, Drosophila, has $\sim 14,000$ protein-coding genes, suggests that it is not gene number but the multitude of regulatory combinations that determine complexity. This is accomplished by the increased elaboration of cis-regulatory elements controlling gene expression coupled with more complex transcription machinery. A simple eukaryotic promoter found in unicellular eukaryotes is contrasted with complex metazoan transcriptional control modules (470).

This section is a compilation of the sequence-specific DNA

D		BLAST match				
Protein	Locus (gene)	Best match ^c	S. cerevisiae	S. pombe	Animal ^a	Plant ^b
ScRad5/SpRad8 ScRad5	9516.1 (rad-5)	S. pombe	1e-124	1e-167	1e-75	1e-118
ScRad26/SpRhp26 HsCSA/ScRad26 Sp Rhp26/HsCSB	4229.1 (csa) 7837.1 (csb)	M. musculus S. pombe	6e-54 9e-64 (Sth1)	4e-58 (Rhp54, Rhp26) 0 (Rhp26)	1e-115 1e-153	1e-118 1e-157
ScRad16 ScRad16 ScRad16-like	3650.1 (<i>rad-16</i>) 3652.1	S. cerevisiae S. pombe	0.0 2e-74	0.0 1e-134	3e-61 2e-31	1e-114 4e-31
ScRad54 ScRad54 ScRad54-like	2349.1 (<i>mus-25</i>) 6190.1 (<i>atrx</i>)	M. grisea; 0.00 O. sativa	0.0 1e-48 (Rad54)	9e-91 7e-55 (Rhp54)	0.0 4e-71	1e-135 2e-72

TABLE 8. Neurospora homologues of predicted ATPases or helicases involved in DNA repair

^a Drosophila melanogaster, Anopheles gambiae, Mus musculus, or Homo sapiens.

^b Arabidopsis thaliana or Oryza sativa.

^c Species listed as best match is Magnaporthe grisea.

binding transcriptional regulatory proteins encoded by the *Neurospora* genome. This preliminary examination reveals that *Neurospora* gene regulation shares elements of both the simple and more complex metazoan models (Fig. 3).

 $Zn(II)_2Cys_6$ fungal binuclear cluster family. The largest class of transcription factors in the *Neurospora* genome belongs to the fungus-specific $Zn(II)_2Cys_6$ fungal binuclear cluster family. Gal4p, a transcriptional activator that regulates galactose utilization in *S. cerevisiae*, is the prototype for this transcription factor family. Structure studies revealed that Gal4p is a $Zn(II)_2Cys_6$ binuclear cluster protein (591). Three functional domains are characteristic of this transcription factor family: a C_6 zinc cluster, which is involved in DNA binding, a middle homology region, which is thought to be necessary for in vivo DNA binding specificity (685), and a third, less well understood activation domain.

The Neurospora genome contains 77 open reading frames (ORFs) with the $Zn(II)_2Cys_6$ motif. Twenty of these factors have no match to proteins in yeasts (Table 9). There are nine ORFs with homologues in other filamentous fungi. Interestingly, half of the putative $Zn(II)_2Cys_6$ motif proteins are most similar to other Neurospora $Zn(II)_2Cys_6$ fungal binuclear cluster proteins, a rare finding in the Neurospora genome. This

result suggests a regulatory mechanism involving sets of factors and several novel mechanisms for gene regulation. The factor set may recognize the same promoter elements, but with different binding affinities, depending on activation or repression requirements. Alternatively, the factors may interact with one another, thus preventing DNA binding.

Only two of the $Zn(II)_2Cys_6$ proteins, NIT-4 and QA1F, have been characterized at the protein level in *Neurospora* (see below). NIT-4, the most extensively studied of the two, is required for nitrate assimilation and interacts with NIT-2, a GATA factor (see below), to activate expression of nitrate and nitrite reductases (230, 257, 892). The QA1F factor, which regulates quinic acid utilization, is another example where protein function and DNA binding properties have been determined (52). The *fluffy* gene product (FL), required for macroconidiation, is also a member of the Zn(II)₂Cys₆ family (40) but has not been characterized at the protein level. Of interest, the closest homologue of FL is a hypothetical protein (NCU09205.1) from *Neurospora*.

C2H2 zinc fingers. There are 43 C2H2 zinc finger transcription factors, making this the second largest class encoded in the genome (Table 10). Unlike the $Zn(II)_2Cys_6$ factors, which are unique to fungi, the C2H2 factors are found in both prokary-



FIG. 3. Transcription factor distribution. (Left) proportion of analyzed *Neurospora* transcription factors in each of the indicated classes. (Right) Venn diagram showing the distribution of Zn_2Cys_6 , C2H2, BZIP, and bHLH transcription factors in bacteria, fungi, and animals.

	Characterized	Best BLAST match					
NCU no.	Neurospora						
	protein	Overall	Yeast"	Comment(s)			
08042.1		Arabidopsis thaliana; 3.00e-27	<i>S. pombe</i> NP_5953118.1; 1.00e-12				
06799.1		S. cerevisiae; 9.00e-11	Best match	A. thaliana; 5.00e-10			
0/669.1		A. nidulans positive regulator of	S. <i>cerevisiae</i> positive regulator of				
00054-1		purine utilization; 0.00	UKAI&3; 5.00e-21 Post match	A thaliana: 1,000,00			
01478 1		S. pombe NP 592804: 1 00e-22	Dest match	A mutunu, 1.000-09 Aspergillus niger FacB: 6 00e-19			
02934 1		N crassa NCU07669 1: 3e-21	S cerevisiae: 7 00e-11	Tisperginus niger Taeb, 0.00e-19			
08000.1		<i>N. hocca</i> cutinase T F 1: 0.00	5. 66.67.68.66, 7.6666 11	Aspergillus orvzae AmdR: 1.00e-56			
07007.1		S. pombe NP 588286; 1.00e-07	Best match				
08294.1	NIT-4	T. inflatum CAB71797 0.00	S. pombe NP 587679; 4.00e-30				
08652.1		Collectotrichum lindemuthianum; 1.00e-110	<i>S. pombe</i> NP_595318; 9.00e-07				
08651.1		N. crassa NCU07669.1; 2.00e-23	S. cerevisiae NP_013114; 1.00e-11				
06656.1	ACU-15	FacB homolog from <i>Aspergillus</i> <i>niger</i> ; 1.00e-153	S. cerevisiae CAT8; 2.00e-23				
02752.1		N. crassa NCU00217.1; 6.00e-64	S. pombe NP_595318; 4.00e-14	EAA27366			
07374.1		N. crassa NCU07705.1; 1.00e-11	S. cerevisiae CAT8; 5.00e-05				
06407.1		<i>S. pombe</i> CAA22853; 2.00e-51	Best match				
05383.1		S. cerevisiae NP_012329; 2.00e-13	Best match				
0021/.1		N. crassa NCU02/52.1; 3.00e-56	None $S_{\text{rescaled}} = C \Lambda \Lambda (2006, 7.00, 07)$				
04000.1		$N_{crassa} NCU02752.1; 5.00e-46$	S. Cereviside CAA05900, 7.00e-07				
05994 1		Aspergillus orvzae TamA: 1 00e-108	S cerevisiae Dal81n: 2 00e-80				
09033.1		<i>S. pombe</i> NP 587679: 7.00e-12	Best match				
05536.1		S. pombe NP 594098; 1.00e-05	Best match				
07705.1		N. crassa NCU00808.1; 4.00e-34	None				
04827.1		S. pombe NP_587726; 4.00e-22	Best match				
07788.1		Aspergillus oryzae; 3.00e-40	S. cerevisiae Malp; 3.00e-22				
06990.1	0.117	<i>S. pombe</i> NP_594000; 1.00e-23	Best match				
06028.1	QAIF	Podospora anserina; 1.00e-140	S. pombe BAA8/112; 4.00e-08	wD repeats; quinic acid utilization activator			
03120.1		A. nidulans; 9.00e-15	None				
01097.1		$N_{crassa} NCU06049.1; 5.00e-35$	None $S_{powbs} NP_{504160} = 3.00 \pm 0.5$	ΕΛΛ26713			
07392.1		Sordaria macrospora Pro1 0.00	<i>S. cerevisiae</i> Ume6p; 5.00e-06	<i>S. macrospora</i> gene required for fruiting-body formation			
08848.1		N. crassa NCU06407; 6.00e-08	None				
08726.1	Fluffy (FL)	N. crassa NCU09205.1; 2.00e-20	S. pombe NP_593001; 4.00e-06				
04359.1		A. nidulans; 2.00e-10	S. pombe NP_595318; 1.00e-05				
08049.1		N. crassa NCU01097.1 3.00e-55	None				
08899.1		N. crassa NCU02896.1; 6.00e-27	S. pombe; 2.00e-09	C time to be in the factor 1 hat			
03045.1		Nectria naemalococca; $1.00e-155$	S. pombe; 1.00e-14	Cutinase transcription factor 1 beta			
01560.1		Fusarium pseudograminearum:	S. pombe, 1.000-05 S. cerevisiae Ume6p/Car80n:	E 4 4 30506			
04390.1		2.00e-17 N crassa NCU02896 1: 9.00e-28	3.00e-06 S. pomba NP 593170: 2.00e-16	L74450500			
03931.1		<i>N. crassa</i> NCU05993.1: 6.00e-07	<i>S. cerevisiae</i> : 3,00e-06	Upc2p involved in sterol uptake			
2307.1		S. pombe NP 593160; 1.00e-10	Best match	N. crassa NCU04001.1; 9.00e-09			
07139.1		<i>S. pombe</i> NP_593605; 2.00e-09		Some homology to <i>A. nidulans</i> PrnA (3.00E-06)			
07945.1		S. pombe NP_592804; 9.00e-12	Best match	A. nidulans PrnA; 9.00e-11			
03320.1		Aspergillus parasiticus Apa-2; 2e-04	None				
08658.1		A. nidulans ArcA; 2.00e-32	None	Regulatory gene in the arginine catabolic pathway			
02094.1		N. crassa NCU00344.1; 5.00e-17	S. cerevisiae Leu3p; 5.00e-25, top hit	Leu3p regulates the levels of Leu1, Leu2, and Leu4			
00808.1		N. crassa NCU07705.1; 8.00e-27	<i>S. pombe</i> NP_596765; 5.00e-06				
00720.1		Aspergillus niger AmyR; 6.00e-24	S. cerevisiae NP_116603; 6.00e-06				
09/39.1		IV. CRASSA INCUUULI/.1; 3.00e-39	S. pombe INP_595100; 5e-12 S. pombe NP_505060; 1,002, 14				
00945 1		N crassa NCU03489 1. 2 00e-05	None				
09829.1		S. pombe NP 5960: 4e-04	Best match	N. crassa NCU01478.1: 5e-05			
06971.1		<i>Hypocrea jecorina</i> xylanase reg 1; 0.00	<i>S. pombe</i> NP_012136; 1.00e-06	Found in A. niger and A. oryzae			
02214.1 02896.1		N. crassa NCU00054.1; 7.00e-15 N. crassa NCU04390.1; 1.00e-46	<i>S. pombe</i> NP_588286.1; 2.00e-08 <i>S. pombe</i> NP_593170; 3.00e-14				

	TABLE 9.	Zn(II	$)_{2}Cys_{6}$	fungal	binuclear	cluster	family
--	----------	-------	----------------	--------	-----------	---------	--------

Continued on following page

NCU	Characterized	Best BLAST match				
no.	protein	Overall	Yeast ^a	Comment(s)		
02142.1		Candida albicans Fcr1p; 1.00e-08	<i>S. pombe</i> Ntf1; 3.00e-07			
02576.1		S. pombe NP 593467; 3.00e-04	Best match			
04001.1		<i>S. pombe</i> NP 593160; 3.00e-13	Best match	EAA32829		
05294.1		Bacillus subtilis acetyltransferase; 2.00e-28		Wbbj, acetyltransferase domain		
00017.1		N. crassa NCU09739.1; 2.00e-37	S. pombe; 5.00e-10	EAA27768		
03686.1		N. crassa NCU03931.1; 1.00e-12	S. cerevisiae Upc2p; 4.00e-08	CAD11364		
05993.1		S. pombe NP 59361; 4.00e-07	Best match	EAA29565		
09804.1		<i>S. pombe</i> NP 594497; 8.00e-18	Best match	EAA3482		
03417.1		N. crassa NCU09549.1; 4.00e-19	S. pombe NP_593170; 5.00e-08	EAA27628		
09549.1		N. crassa NCU04851.1; 2.00e-38	<i>S. pombe</i> NP_593170; 4.00e-10	EAA29532.1		
00289.1		Aspergillus fumigatus; 2.00e-54	None	EAA28534		
08901.1		N. crassa NCU09804.1; 8.00e-05	None	EAA29603.1		
04851.1		N. crassa NCU09549.1; 4.00e-34	S. cerevisiae Rdr1p; 5.00e-06	EAA28676		
02768.1		N. crassa NCU02896.1; 2e-32	S. pombe NP_593170; 3.00e-24	EAA36342		
08063.1		Nectria haematococca; 2.00e-06	None			
07535.1		<i>N. haematococca</i> flanks pea pathogenicity (PEP) cluster; 9.00e-12	S. pombe Ntf1; 1.00e-06	EAA29812.1		
03489.1			S. cerevisiae YKR064wp; 2.00e-05	EAA26640		
09205.1		A. nidulans NIRA; 2.00e-45	S. pombe NP 595069; 6.00e-12	EAA29913.1		
05051.1		N. crassa NCU03417.1; 5e-25	S. pombe NP 593170; 1e-16			
09529.1		N. crassa NCU07575.1; 8E-04	S. cerevisiae NP 013610; 0.003	EAA28818		
08289.1		None below 0.003	None below $0.0\overline{0}3$	EAA33266		

TABLE 9—Continued

^a S. cerevisiae or S. pombe.

otic and eukaryotic organisms (99). What is striking about this class of transcriptional regulators is that they can be easily separated into two distinct groups. About half are most homologous to *S. cerevisiae* and *S. pombe* protein sequences, while the remainder are most similar to proteins from filamentous fungi and animals. Several members of the latter group have homology to Krüpple-type ZNF transcription factors from mice, rats, and humans (Table 10). None of the C2H2 zinc finger transcription factor proteins have yet been characterized in *Neurospora*.

The Neurospora Ste12p-like transcription factor (NCU00340.1) has a C2H2 domain as well as a homeodomain, and it has five homologs in available databases: A. nidulans SteA (819), Magnaporthe grisea Mst12 (597), Gibberella zeae Fst12, Colletotrichum lagenarium Cst1 (805), and Penicillium marneffei St1A (96). Ste12p from S. cerevisiae has been extensively characterized and shown to play regulatory roles during mating for haploid cells (200, 222, 234) and filamentous growth of diploids (489). The S. cerevisiae protein lacks a C2H2 domain. The conservation of this protein in filamentous fungi is remarkable, suggesting that this regulatory element is critical and necessary for mating and development. Three of the genes encoding fungal homologues (SteA, CST1, and Mst12) have been deleted in their respective organisms. The steA deletion strain is sterile and exhibits defects in ascogenous tissue and fruitingbody development, but no effect on vegetative growth or morphology was detected (819). The cst1 deletion mutant is impaired in the production of infectious hyphae from appressoria, rendering it nonpathogenic (805). When the mst12 deletion mutants were tested on onion epidermal cells, appressoria appeared normal but could not penetrate and carry out infectious growth. These phenotypes reveal important roles for Ste12p-like proteins in developmental pathways and pathogenesis.

The dichotomous results found with C2H2 transcription factors, half with homology to proteins from unicelluar yeasts and the other half with sequences from filamentous fungi and animals, suggest that these factors represent a point of divergence in promoter evolution and gene regulation. A recent study of human and mouse C2H2 factors offers evidence supporting continuing evolution of C2H2 zinc finger proteins. There are several familial gene clusters encoding C2H2 zinc finger proteins in these vertebrates. A careful analysis of a syntenic region in humans and mice suggested that the analyzed clusters arose from tandem duplications with eventual divergence, resulting in a large assortment of this type of factor (722). Unlike other gene family clusters in mammals, specifically those involved in immunity and smell, these did not contain pseudogenes. This suggests that C2H2 zinc finger gene family is still evolving in mice and humans.

GATA factors. The founding member of the extensive family of GATA transcription factors, GATA-1, was cloned in 1989 and shown to regulate erythroid cell differentiation in vertebrates (226, 804). GATA transcription factors bind (A/T)GAT A(A/G) motifs and have one or two Cys-X₂-Cys-X₁₇-Cys-X₂-Cys zinc fingers. Unlike the C2H2 zinc finger proteins, GATA factors are found only in eukaryotic organisms. Although only six have been identified in the *N. crassa* genome, all but one has been cloned, making this class of transcription factors the most extensively characterized in this organism (Table 11). *Neurospora* GATA factors regulate critical processes, ranging from nitrogen utilization and iron uptake to light regulation and ascospore development.

TABLE 10. C2H2 zinc finger transcription facto	TABLE	inscription facto	C2H2 zinc finger	ctors
--	-------	-------------------	------------------	-------

NCU no	Best BLAST match					
NCU IIO.	Overall	Yeast ^a	Comment(s)			
03975.1	Mus musculus; 2.00e-33	None	Three domains			
10025.1	Mus musculus; 2.00e-33	None	Identical to above			
02699.1	S. cerevisiae; 6.00e-24	Best match	Zap1p DNA binding protein; zinc ion homeostasis			
02666.1	S. cerevisiae; 2.00e-15	Best match	Crz1p calcineurin responsive			
04179.1	S. cerevisiae NP 014756; 1.00e-49	Best match	Asparagine-rich zinc finger			
03421.1	Homo sapiens locus AAH07307; 4.00e-27	None	Two domains			
06907.1	S. pombe NP 594109; 4e-25	Best match	S. cerevisiae Ace2p; 2.00e-16			
00038.1	S. pombe NP 594670; 4.00e-36	Best match	TFIIIA from frog			
01629.1	Zebrafish NP 571798; 4.00e-12	None	Krupple-like factor			
07952.1	S. cerevisiae NP 014371; 2.00e-37	Best match	One domain; Crz1p calcineurin responsive			
02853.1	Colletotrichum lagenarium CMR1; 2.00e-12	S. cerevisiae NP 012661; 8.00e-08				
02671.1	Nectria haematococca MPV1; 3.00e-65	S. cerevisiae YER130cp; 6.00e-20	Cutinase G-box binding protein			
02173.1	Colletotrichum lagenarium CMR1; 1.00e-12	S. cerevisiae Adr1p; 3.00e-07	Two domains			
06503.1	Emericella nidulans AmdA: 3.00e-16	S. cerevisiae YM1081Wp; 2.00e-06	Two domains: positive-acting TF			
04561.1	Colletotrichum lagenarium CMR1: 6.00e-23	S. pombe NP 592812; 9.00e-08	, i i i i i i i i i i i i i i i i i i i			
08807.1	Trichoderma reesei Cre1; 4.00e-82	Debaryomyces occidentalis Mig1 protein: 5E-23	Two domains; CRE-1 carbon catabolite repressor			
05064.1	S. cerevisiae NP 015094; 7e-13	Best match	Up-regulated during starvation in S. cerevisiae			
10006.1	Homo saniens BAA91019: 4.00e-11	None	•F • • 8 · · · · · · · · · · · · · · · · ·			
00694.1	D, willistoni AAO01096: 5 00e-06	None	Two domains			
06487.1	Rattus norvegicus XP 234852: 1.00e-06	None	Two domains			
09576.1	Drosophila sp. Sp1/egr-like: 8.00e-08	S. cerevisiae MET32: 3.00e-06	Two domains: regulator of sulfur metabolism			
03043.1	Podospora anserina CAD12881; 1.00e-75	None	Two domains; <i>fle</i> gene coordinates male and female sexual differentiation			
02994.1	Ascobolus immersus CAA67549; 5.00e-45	None				
05285.1	Emericella nidulans AA024631; 1.00e-29	S. pombe 0.018				
03184.1	S. pombe CAB61785; 8.00e-10	Best match	Four domains			
01122.1	S. pombe NP_594996; 9.00e-28	Best match	Two domains			
03699.1	Ascobolus immersus CAA67549; 4.00e-62	None	Four domains			
05242.1	S. pombe NP_594996; 7.00e-47	Best match				
03206.1	S. pombe CAB59682; 4.00e-35	Best match	COG5048 SFP: putative transcriptional repressor regulating G ₂ /M transition			
00340.1	C. langenarium BAC11803.1; 0.00	S. cerevisiae NP_011952; 2.00e-54	STE12 has both a homeodomain and the C2H2 domain			
	M. grisea Mst12; 0.0 G. zeae AF509440; 0.0 E. nidulans SteA; 0.0 P. marafai AF284062: 0.0					
02713.1	S cerevisiae NP 009622: 2 00e-09	Best match	RG1 transcriptional repressor—invasive			
02713.1	B. COLUMN 111_000022, 2.000 00	N	growth			
05909.1	Homo saplens NP_114124.1; 4.00e-11	None	Two domains; BTE binding protein 4			
09355.1	hypocrea jecorina AAF 55280; 1.00e-180	None	promoter <i>chb1</i> in <i>Trichoderma reesei</i>			
09252.1	N. crassa NCU04628.1; 1.00e-07	None	I hree domains			
00090.1	Conetotrichum subineolum; Pac C 5.00e-67	S. cerevisiae KIN11; 5.00e-28	regulates melosis in yeast			
03552.1	Mus musculus LOC2446/4; 3.00e-06	None	i wo domains			
04019.1	Aspergulus fumigatus CAD2844/; 2.00e-31	None				
00385.1	N. crassa CAD3/059; 8e-14	INORE				
04(28.1	Asperguius niaulans AAF15889; 1e-33	5. pombe INP_594670; 4.00e-05	Four domains			
04028.1	N. crassa NCU09252.1; 2.00e-07	None	1 wo domains			
00285.1	N. crassa NCUU4619.1; 9.00e-14	None	Circular and in a			
00919.1	Kunus norvegicus INP_5/9846; /.00e-09	inone	Six domains			

^a S. cerevisiae, S. pombe, or Debaryomyces occidentalis.

NIT-2, a positive regulator of nitrate assimilation and nitrogen utilization, was shown to have one DNA binding motif (262). Careful analysis of NIT-2 binding to promoters of nitrogen-regulated genes revealed that two closely spaced GATA sites were required for high-affinity binding by this factor to the DNA (146, 231). Another GATA factor, SRE, was isolated using a probe generated from a PCR with degenerate primers to the conserved zinc finger region (899). This GATA factor has two zinc fingers and sequence similarity to several fungal proteins, SreP from *P. chrysogenum*, UrbS from *U. maydis*, and SreA from *A. nidulans*. All are negative regulators of siderophore synthesis (317, 318, 835). SRE negatively regulates siderophore production in *Neurospora* (898). Analysis of a *sre* null mutant (898) revealed the existence of other regulators involved in maintaining iron homeostasis. However, these components have not been identified. The cloning and sequencing of the genes that encode White Collar 1 (WC-1) and White Collar 2 (WC-2) revealed that they both contain a single GATA transcription factor motif (42, 478). These GATA factors mediate responses, including input for the circadian clock and light-induced expression of genes that regulate blue light processes, as well as functioning exclusively in

NCU no.	Characterized	Best BLAST match				
	protein	Overall	Yeast ^a	Comment(s)		
09068.1	NIT-2	Colletotrichum lindemuthiannum; 1.00e-139	<i>S. cerevisiae</i> YFL021w BAA09217; 8.00e-14	One GATA domain; Major nitrogen regulatory protein		
07039.1	ASD-4	Penicillium chrysogenum; 7.00e-41	S. cerevisiae Gzf3p/Dal80; 4.00e-19	One GATA domain		
07728.1	SRE	Botryotinia fuckeliana; 3.00e-44	S. cerevisiae UGA43; 9.00e-09	Two domains; siderophore regulation protein		
00902.1	WC-2	Nectria haematococca; 1.00e-153	None	One GATA domain; one PAS domain; palindrome-binding protein (PBP)		
02356.1	WC-1	Podospora anserina; 0.00	None	One GATA domain; three PAS domains; two PAC motifs		
01154.1		Penicillium chrysogenum; 2.00e-11	None	One GATA domain		

TABLE 11. GATA factors

^{*a*} S. cerevisiae or S. pombe.

the dark as the positive elements of the core feedback loop of the circadian clock (see "Photobiology and circadian rhythms" below). Each factor has PAS domains; WC-1 has three, and WC-2 has only one. One of the PAS domains in WC-1 is also a LOV (for "light, oxygen, or voltage") domain, a motif that was recently shown to be required for light sensing in *Neurospora* (144). The final characterized GATA factor is ASD-4, which is required for ascospore development (229). The exact role of ASD4 in this developmental pathway is not known.

There are four characterized GATA factors in *S. cerevisiae;* Gln3p, Nil1p/Gat1p, Dal80p, and Gfz3p/Nil2p (155, 175, 531, 753). All four participate in nitrogen regulation of gene expression, with the first two acting as positive regulators and the last two acting as negative regulators. It appears that the fungal GATA factors regulate processes that require exquisite balance in maintaining metabolic homeostasis. In nitrogen utilization, it is suboptimal for an organism to expend cellular energy taking up metabolites that it does not need. However, since nitrogen is essential, a critical level must be sustained. In addition to regulating nitrogen utilization in *Neurospora*, GATA factors regulate genes required for iron homeostasis and adaptation to blue light. Iron and light are necessary for normal growth and development in this organism; however, excesses are known to be deleterious.

bHLH transcription factors. The basic helix-loop-helix (bHLH) motif was first described in mammals (551) and is

unique to eukaryotes. This motif consists of conserved bipartite domains that dictate DNA binding and protein dimerization (551). A phylogenetic analysis based on 122 divergent bHLH sequences revealed that bHLH factors fall into four groups based on the DNA binding and protein interaction domains (32). Ten predicted bHLH or HLH proteins have been identified in the *Neurospora* genome sequence (Table 12); only one has been cloned and characterized, the *nuc-1* gene product, which was shown to regulate phosphorous utilization (399). All predicted *Neurospora* bHLH proteins are more similar to other fungal bHLH proteins than to animal bHLH proteins, and all are members of the B group (32). Interestingly, this family of transcription factors represents one of the largest found in the *A. thaliana* genome, with over 133 members (647).

B-ZIP transcription factors. The defining B-ZIP motif is a bipartite α -helix between 60 and 80 amino acids in length, with a DNA binding region of two basic clusters at the N terminus and a dimerization domain, formed by an amphipathic helix with a leucine every 7 amino acids, at the C terminus (833). The heptads, which comprise the leucine zipper, can be of different lengths and can participate in dimerization. These factors can homodimerize, heterodimerize, or homo- and heterodimerize, giving each transcription factor the capability of multiple functions.

NCU no.	Characterized		Best BLAST match	
	protein	Overall	Yeast ^a	Comment(s)
02957.1		Aspergillus nidulans; 2.00e-21	S. pombe NP 593230; 2.00e-08	bHLH sexual differentiation protein Esc1p
06744.1		No hits below 1.00e-05	None	BHLH; CAD70402
01871.1		N. crassa XP 328825; 9.00e-56	S. pombe NP 596545; 7.00e-44	HLH/MCM; Mcm7p
00749.1		Fusarium culmorum ANN73248; 8.00e-10	<i>S. pombe</i> NP_595229; 0.075	HLH I domain; USF protein
03077.1		N. crassa XP 331923; 3.00e-06	S. pombe NP 595229; 0.002	HLH; A thaliana H84860 1e-05
02724.1		A. nidulans AAG49357; 1.00e-24	<i>S. cerevisiae</i> NP_009447; 2.00e-06	bHLH; GLCD beta An; Rtg3pSc
05970.1		N. crassa XP 324088; 1.00e-13	S. pombe NP 595694; 1.00e-13	HLH
04731.1		N. crassa XP329061; 3.00e-23	<i>S. pombe</i> NP 595694; 5.00e-23	HLH
00144.1		Fusarium culmorum ANN73248; 1.00e-14	<i>S. cerevisiae</i> NP_014989; 0.42	HLH
09315.1	NUC-1	None	None	HLH; controls phosphorus aquisition

TABLE 12. bHLH transcription factors

^{*a*} S. cerevisiae or S. pombe.

NCU no.	Characterized	Best BLAST match				
	protein	Overall	Yeast ^a	Comment(s)		
04050.1	CPC-1	Cryphonectria parasitica; 5.00e-18	None	BZIP; general amino acid control		
03905.1		S. pombe NP 593662; 1.00e-05	Best match	BZIP; AP-1-like transcription factor		
08055.1		Cladosporium fulvum; 2.00e-09	None	BZIP		
01345.1		Claviceps purpurea; 7.00e-87	S. pombe Atf CREB family; 6.00e-10	BZIP; Cptf1 involved in oxidative stress		
01459.1		S. pombe Atf CREB family; 4.00e-11	4.00E-36	BZIP		
00499.1		None less than 1e-05	None	BZIP		
08744.1		None less than 1e-05	None	BZIP		
07900.1		None less than 1e-05	None	BZIP		
05637.1		None less than 1e-05	None	BZIP		
00329.1		None less than 1e-05	S. cerevisiae Adr1p; 3.00e-07	BZIP		
00233.1		Apergillus nidulans MeaB; 3e-41	None	BZIP		
08891.1		Apergillus oryzae; 2.00e-05	None	BZIP		
06399.1		None less than 1e-05	None	BZIP plus Rho guanine nucleotide exchange domain; EAA33001		
07379.1		None less than 1e-05	None	BRLZ; EAA29636		
04211.1		N. crassa NCU03905.1 5e-05	Candida albicans; 0.001	BZIP		
01994.1		None less than 1e-05	Candida albicans; 0.002	BZIP; EAA35623		

TABLE 13. B-ZIP transcription factors

^a S. cerevisiae, S. pombe, or C. albicans.

A total of 17 predicted B-ZIP proteins are present in the *Neurospora* genome sequence (Table 13). This number includes CYS-3, a well-characterized B-ZIP protein involved in regulating sulfur uptake and utilization (260, 396, 585), which escaped identification by the automated gene-calling program in the present WICGR assembly. Based on the reported number of unique B-ZIP proteins for other sequenced genomes, 53 for human (832), 27 for *Drosophila*, and 17 for *S. cerevisiae* (228), it appears that the majority of this class of transcription factors has been identified in *Neurospora*.

In addition to CYS-3, the *Neurospora* B-ZIP protein CPC-1, a Gcn4p ortholog, has been previously characterized. Gcn4p and CPC-1 were first identified as regulators of genes expressed during amino acid starvation or imbalance (46, 339, 589). The global nature of this regulation in *S. cerevisiae* was reported previously (28), and a recent review illustrates the central position of Gcn4p in regulating responses to environmental signals (340).

Half of the other identified *Neurospora* B-ZIP proteins have no significant homology (E values greater than 1e-05) to any protein sequences presently in GenBank. The remainder are most similar to proteins from other filamentous fungi and *S. pombe;* only NCU04050.1 (CPC-1) has highest homology to the *S. cerevisiae* Gcn4p protein (Table 13).

Miscellaneous factors. An additional 23 putative DNA binding proteins are listed in Table 14; they include 4 CBF CAAT binding factors, 3 forkhead domain proteins, 6 homeodomain, and 6 RING finger and WD repeat proteins. These factors play important roles in the biology of eukaryotic organisms. They link transcription to the cell cycle, RNA metabolism, meiosis, mitosis, cell death, DNA repair, chromatin remodeling, and nucleosome assembly. *Neurospora* shares four CBF CAAT binding factors with yeast. However, it has three winged-helix forkhead factors, one more than identified in *S. cerevisiae* (Table 14). Included in this list is RCO-1 (869), a protein similar to *S. cerevisiae* Tup1p, which plays a role in transcriptional repression. The identified homeodomain proteins have higher homology to filamentous fungi and metazoans than to unicellular yeasts. This is also true of proteins with the RING finger motif; of this group, only *Neurospora* UVS-2, similar to *S. cerevisiae* Rad18p, has been characterized (792).

Translation Factors

Many polypeptide components comprise the factors important for translation initiation, elongation, and termination (747). Analysis of the genome sequence shows that hypothetical *Neurospora* proteins homologous to cytoplasmic translation factors in other organisms (Table 15) are often incorrectly annotated (e.g., the predicted polypeptides contain N-terminal extensions, or the genes from which they are conceptually translated have incorrectly predicted intron structures). Nevertheless, several conclusions can be reached concerning the nature of eukaryotic initiation factors (eIFs), elongation factors (eEFs), and release (termination) factors (eRFs) in *Neurospora*.

The *Neurospora* genome contains essentially the same complement of translation factors as do the *S. cerevisiae* and *S. pombe* genomes. Each contains a gene for eEF3, while the animal and plant genomes lack a closely related polypeptide (20). Unlike animals and/or plants, these fungi lack eIF3j, eIF3k, and eEF1B β . One difference between *Neurospora* and the two yeasts is that the former has a protein similar to the eIF3l polypeptide that is known to be in the plant eIF3 complex, which is present in animals and plants but lacking in the yeasts (118). The function of this protein is unknown.

The predicted *Neurospora* translation factors more closely resemble those of *S. pombe* than those of *S. cerevisiae*. In the majority of cases, the *Neurospora* translation factor has a primary sequence more similar to the *S. pombe* factor. The *Neurospora* and *S. pombe* genomes, as well as animal and plant genomes, contain sequences for eIF3d-f and eIF3h, while *S. cerevisiae* lacks similar polypeptides. The *Neurospora* and *S. pombe* genomes both lack apparent homologues of eIF4B. This

NCU	Characterized	Best BLAST match					
no.	protein	Overall	Yeast ^a	Comment(s)			
03033.1 09248.1 03073.1		Hypocrea jecorina HapB; 9.00e-34 Hypocrea jecorina Hap3; 3.00e-49 Ratius norvegicus; 2.00e-06	<i>S. cerevisiae</i> Hap2p; 1.00e-17 <i>S. cerevisiae</i> Hap3p; 2.00e-26 <i>S. pombe</i> ; 1.00e-11	CBF CAAT binding factor CBF CAAT binding factor/HMF CBF CAAT binding factor/HMF; EAA34930			
02017.1		<i>R. norvegicus</i> TBP binding NC2; 1.00e-18	S. pombe NP_596283; 6.00e-29	CBF CAAT binding factor/HMF; EAA35646			
00019.1		S. cerevisiae Fkhp1: 5.00e-41	Best match	Two forkhead domains			
06173.1		S. pombe Hnf-3; 2.00e-28	Best match	One forkhead domain; regulation of septation			
08634.1		S. cerevisiae AAA99643; 1.00e-07	Best match	One forkhead domain			
03244.1		Branchiostoma floridae; 2.00e-72	S. pombe NP_595227; 8.00e-46	WD repeats; <i>D. melanogaster</i> (4.00e-71) will die slowly			
06205.1	RCO-1	Penicillium marneffei; 1.00e-167	S. pombe NP_592873; 1.00e-117	WD repeats; Transcriptional Repression			
06411.1		No hits below 1e-05	None	RING-type zinc finger			
03593.1		Podospora anserina CAC16792; 0.00	S. cerevisiae NP_010177/Pho2p; 9e-09	Homeobox			
03070.1		D. melanogaster AAD38649; 1.00e-07	S. cerevisiae CAA44264; 9.00e-04	Homeobox			
03266.1		M. musculus ACC53336; 7.00e-06	None	Homeobox			
05257.1		D. discoideum AA052126; 7.00e-10	None	Homeobox; Lycopersicon esculentum; 3e-08			
00097.1		H. sapiens AAC51243; 5.00e-13	S. cerevisiae NP_015148/Cup9p; 1.00e-08	Homeobox			
05250.1		Coprinus cinereus S71461; 2.00e-65	S. pombe NP_592917.1; 3.00e-66	Homeobox; sister chromatid cohesion molecule			
03962.1		H. sapiens BAB14697; 5.00e-06	S. pombe NP_594050; 1.00e-09	Human herpesvirus 6 immediate early protein			
05210.1	UVS2	A. nidulans CAA90033: 1.00e-63	S. pombe NP 595423: 1.00e-32	RING finger UVS2			
06213.1		<i>S. pombe</i> NP_593123; 6.00e-38	<i>S. cerevisiae</i> NP_010697; 1.00e-34	MIZ zinc finger			
07561.1		H sanians $BAA96066$; 200e-12	S nombe NP 588450: 2 00e-17	PHD-finger			
00631.1		<i>S. pombe</i> NP_596602; 1.00e-118	Best match	SNF2/RING/Helicases; A. thaliana			
				NP_564568; 1.00e-52			
01954.1		H. sapiens CAC42525; 1.00e-109	S. pombe NP_596357; 6.00e-79	RING finger; C3HC4			
03356.1		M. musculus P70191; 9.00e-05	None	Related to TRAF5; RING-type zinc finger			

FABLE
FABLE

^{*a*} S. cerevisiae or S. pombe.

polypeptide is poorly conserved between *S. cerevisiae* and higher eukaryotes and thus may have gone undetected because of low sequence similarity, since the closest *S. pombe* match to eIF4B, Sce3p, does not have an obvious homologue in *Neurospora*.

GENOME DEFENSE, DNA REPAIR, AND RECOMBINATION

Genome Defense Mechanisms

A basic tenet of classical genetics is the notion that the structure and behavior of an organism are determined by the structure and workings of its genome. In contrast, the idea that the structure and behavior of a genome may reflect, at least in part, the structure and behavior of the organism is relatively new. Some of the earliest and strongest evidence for active processes that can shape genomes has come from studies of *Neurospora*. This organism provided the first example of a eukaryotic genome defense system, RIP (Fig. 4) (709, 712). Like most filamentous fungi, plants, and many animals, but in contrast to *S. cerevisiae* and *S. pombe, Neurospora* can methylate its DNA (712). More recently, studies have shown that *Neurospora* employs at least two additional epigenetic systems

that appear well suited to help maintain its streamlined genome: quelling and meiotic silencing (Fig. 4) (24, 131, 732). Repeated sequences are detected and inactivated during haploid, vegetative growth by a vegetative RNA silencing mechanism called quelling (157, 158, 162). Quelling produces diffusible signals, small interfering RNAs, that interfere with the propagation of the repeated element within nuclei in the same cytoplasm (132). During meiosis, *Neurospora* cells use a process related to RNA interference to check for unpaired sequences; this process is called meiotic transvection (24, 25) and meiotic silencing (732). The many convenient features of *Neurospora*, including its streamlined genome, facile genetics, and well-developed molecular tools, have made this organism a preferred model for the study of genomic surveillance processes.

Heterochromatin silencing and DNA methylation. The terms "gene silencing," "transcriptional silencing," "silent chromatin," and "heterochromatic silencing" are often used interchangeably to describe processes that, in contrast to promoter- or gene-specific transcriptional regulation, act on larger chromosomal regions (for reviews, see references 34, 299, and 535). Stably inherited gene silencing is called "epigenetic" because phenotypes are maintained over many generations but

TABLE 15. Neurospora translation factors

Protein	NCU	CU BLAST matches				
class	no.	Closest ^c	S. cerevisiae	S. pombe	Animal ^a	Plant ^b
Initiation						
factors						
eIF1	01981.1	Sc	2e-31	7e-28	3e-25	4e-25
eIF1A	07437.1	Animal	2e-32	2e-37	2e-39	1e-29
eIF2α	08277.1	Sp	2e-88	3e-89	2e-62	4e-67
eIF2β	04640.1	Sc	4e-49	9e-49	2e-45	2e-43
eIF2γ	02810.1	Sc	0	0	0	0
eIF2Bα	04344.1	Sp	2e-30	6e-32	7e-20	6e-16
eIF2Bβ	04640.1	Sc	1e-49	2e-49	5e-46	4e-44
eIF2Bγ	03548.1	Sp	2e-18	2e-19	2e-16	
eIF2Bδ	01468.1	Sp	3e-31	4e-43	5e-31	1e-33
eIF2Be	02414.1	Sp	2e-80	4e-95	4e-59	6e-67
eIF3a	00040.1	Sp	4e-87	e-167	e-122	e-103
eIF3b	02208.1	Sp	e-114	e-167	e-120	e-114
eIF3c	07831.1	Sp	e-108	e-180	e-102	e-135
eIF3d	07380.1	Sp		e-136	e-87	e-90
eIF3e	05889.1	Animal		3e-85	3e-91	7e-85
eIF3f	01021.1	Sp		5e-62	7e-30	4e-26
eIF3g	08046.1	Animal	6e-19	2e-19	3e-27	8e-22
eIF3h	07929.1	Sp		3e-59	2e-41	9e-54
eIF3i	03876.1	Sp	1e-99	e-119	4e-93	8e-73
eIF3i		~r				
eIF3k						
eIF3l	06279.1	Animal			e-113	8e-73
eIF4A	07420.1	Sn	e-146	e-177	e-162	e-149
eIF4B	07.12011	SР	• 110	• 1//	0 102	• 1.0
eIF4F	02076 1	Sn	3e-37	1e-48	5e-27	4e-22
eIF4G	07868 1	Sp	3e-47	3e-82	4e-42	8e-26
eIF5	00366.1	Sp	4e-83	8e-88	7e-62	4e-34
eIF5A	$05274 1^d$	Sc	2e-51	2e-46	3e-48	$2e_{-40}$
eIE5B	05270.1	Sc	0	0	e-166	0
eIF6	09004 1	Sc	e-103	e-100	5e-07	7e-0/
PARP	04799 1	Sn	e-130	e-157	e-125	e-116
TADI	07/)).1	Sþ	C-150	0-157	0-125	C-110
Elongation						
eFF1A	02003 1	Sn	0.00	0.00	0.00	0.00
eEF1Re	06035.1	Sp Sp	20.00	5e 40	20.00	10.00
eFF1B0	00033.1	эр	20-20	50-47	20-44	10-50
eFF1Be	03826 1	Sc	$2e_{-81}$	2e-71	76-54	0e-13
oEF1D y	03820.1	Sc	0.00	20-71	0.00	0.00
eEF2	07022 1	Sc	0.00	0.00	$7_{e_{-}} 1 1$	1e-42
CEFJ	0/944.1	30	0.00	0.00	/0-44	10-42
Termination factors						
eRF1	00410.1	Animal	e-176	e-166	0.00	e-174
eRF3	04790.1	Sp	e-159	e-164	e-147	e-133

^a Anopheles gambiae, Drosophila melanogaster, Homo sapiens, Mus musculus, Oryctolagus cuniculus, Rattus norvegicus, or Xenopus laevis.

^b Arabidopsis thaliana, Oryza sativa, Triticum aestivum, or Zea mays.

^c Sc, S. cerevisiae; Sp, S. pombe.

^d BLAST results obtained using Protein Accession AAA61707.1.

are not determined by the DNA component alone. Two major, evolutionarily distinct mechanisms for chromatin-associated gene silencing that operate through posttranslational histone modifications have emerged, currently best characterized in the yeasts *S. cerevisiae* (Sir-mediated silencing) and *S. pombe* (HP1/Swi6-mediated silencing) (reviewed in references 34, 275, 299, 353, 384, 535, and 651). *Neurospora* contains homologues of most proteins previously implicated in gene silencing. Some proteins considered key in the silencing pathways of *S. cerevisiae*, *S. pombe*, plants, and *Drosophila* are absent from the *Neurospora* proteome, however. In contrast to *S. cerevisiae*, *S.* pombe, and Drosophila, Neurospora also uses DNA methylation to inactivate genes.

In *S. cerevisiae*, some chromosomal domains are transcriptionally silenced by the action of complexes that contain Sir proteins (275, 353). *Neurospora*, like *S. pombe*, plants, and metazoans, does not have homologues of *S. cerevisiae* Sir1p, Sir3p, Sir4p, and Net1p. Sir2p-like deacetylases (sirtuins) are the only members of Sir complexes conserved from bacteria to eukaryotes, and seven putative *Neurospora* sirtuins have been identified (see "Chromatin assembly and gene regulation" above).

Formation of heterochromatin in S. pombe, plants, and metazoans is achieved by HP1/Swi6-mediated silencing (reviewed in references 34, 299, 353, 535, and 651). As outlined in "Chromatin assembly and gene regulation" (above), Neurospora has components of this pathway, e.g., well-conserved HATs of the GNAT and MYST type, HDACs of the yeast Rpd3/Hos/ Hda group, bromodomain proteins which are involved in the recognition of acetylated lysines, at least 3 putative histone kinases and 3 phosphatases, and 12 known or putative HMTs. Most Su(var)3-9 lysine methyltransferases contain a chromodomain within their amino terminus, but Neurospora DIM-5 lacks this domain (776). Curiously, Neurospora has two short chromodomain proteins (CDP-1 and CDP-2 [Table 16]) that bear Su(var)3-9-like chromodomains. Neurospora also has a single HP1/Swi6 homologue, heterochromatin protein 1, (HP1), characterized by an N-terminal chromodomain and C-terminal chromoshadow domain. In fission yeast, the formation of heterochromatin at the silent mating-type locus and the centromeres is dependent on homologues of the RNA interference machinery (307, 837), a process that, at least in plants, may involve DNA methylation (277, 524).

Over the last 15 years, Neurospora has been developed as a model to investigate the control and function of DNA methylation in eukaryotes, an area which attracted increased attention with the realization that abnormal methylation is commonly associated with cancer and other diseases (54, 73, 387). Neurospora has proven to be particularly useful in efforts to elucidate the establishment of methylation. In Neurospora, about 1.5% of the cytosines (C's) are methylated (246, 669), but no methylation has been reported for any of the proteincoding genes that have been studied. Three naturally methylated regions have been characterized in great detail: the tandemly arranged rDNA (611), the 1.6 kb zeta-eta (ζ - η) region (526, 714-716), and the psi-63 (\u00c663) region (246, 500, 525, 527). The last two regions are relics of RIP. It is possible that almost all DNA methylation in the wild-type Neurospora genome, including the limited methylation of the rDNA, is a result of RIP. Indeed, a survey of methylated Neurospora sequences isolated by affinity chromatography, using the methyl binding domain of mammalian MeCP2, revealed clear evidence of RIP in nearly all sequenced fragments (717). It is clear, however, that sequences not exposed to RIP can be methylated in the genome. For example, some transforming sequences are subject to de novo methylation without going through the sexual cycle (114, 575, 594, 662, 706, 714, 715). Detailed analyses of short DNA segments constructed in vitro revealed that the methylation machinery preferentially recognizes AT-rich regions as short as 75 bp (526, 777). Methylation indirectly blocks transcription elongation in Neurospora (664)



FIG. 4. Neurospora silencing pathways. The stages of the life cycle of Neurospora are presented, indicating points where the various gene silencing pathways are active. Quelling is a post-transcriptional gene silencing (PTGS) pathway that is active in the vegetative phase of the life cycle, from germination of ascospores or conidia to formation of the mycelium and differentiation of conidiophores and conidia. RIP and meiotic silencing are silencing pathways that are specific to the sexual cycle but differ in their molecular mechanisms. RIP scans for the presence of duplicated copies of DNA fragments present in the genomes destined to participate in meiosis. Duplicated regions are inundated with a series of transition mutations, and most of the remaining nonmutated cytosine bases are methylated. This process occurs in the heterokaryotic ascogeneous tissue formed following fertilization but prior to karyogamy. Meiotic silencing, like quelling, is a PTGS-like mechanism that is activated when a discrete region of DNA fails to sense (i.e., trans-sense) an equivalent region in the opposite chromosome. This failure of trans-sensing in turn triggers the silencing of all genes contained in the loop of unpaired DNA.

and can silence transforming DNA (365). Hence, DNA methylation may serve as a genome defense system (707) distinct from quelling (131) and meiotic silencing of unpaired DNA (24, 732).

A connection between one of the steps in HP1/Swi6-mediated silencing, histone H3 methylation, and genome-wide DNA methylation was first shown in Neurospora (776). More recently, this connection has been established in Arabidopsis (375, 386, 496) and in animals (39, 264, 465). Neurospora has two cytosine DNA methyltransferase (DMT) homologues, DIM-2 and RID, but no homologues of plant chromomethylases, plant domain rearranged methylases (DRMs), or the S. pombe PMT1 and human DNMT2 candidates for DMT pseudogenes. DIM-2 is required for all known DNA methylation in Neurospora (247, 433, 712), whereas RID, a member of a family of putative DMTs known only in filamentous fungi, is required for RIP (253). Neurospora does not have proteins that bear canonical methyl-binding domains (MBDs), suggesting the existence of alternative proteins involved in the recognition and maintenance of methylated DNA, compared to the situation for Arabidopsis and mammals (712). As indicated above, SWI/ SNF-like putative CRFs from Arabidopsis (DDM1) and humans (LSH) are involved in DNA methylation (for reviews, see references 100 and 651). DDM1 is essential to silence transposons and transgenes via control of histone H3 lysine 9 methylation (277). While Neurospora has a putative homolog of LSH (CRF5-1), another SWI/SNF-like Arabidopsis protein involved in silencing, MOM (19), is apparently absent from the Neurospora proteome.

In conclusion, the picture emerging from a combination of experimental and bioinformatic analyses predicts that Neurospora relies on a variation of the HP1/Swi6 pathway to achieve a silenced chromatin state that is coupled directly or indirectly to de novo and/or maintenance DNA methylation (712). Whether DNA methylation is involved in all heterochromatin silencing remains to be determined.

	I	Match found by BLAST				
Protein	Locus (gene)	Best match	S. cerevisiae	S. pombe	Animal ^b	Plant ^c
Bromodomain proteins						
ScRsc1/Rsc2	2354.1 (pbd-1)	P. anserina; 0	6e-76	4e-68	2e-20	No match
SpBrd1	2078.1 (pbd-2)	M. musculus	No match	1e-6	1e-8	No match
<i>Sp</i> Bdf1-like	8809.1 (bdp-1)	D. rerio	No match	4e-12	8e-14	2e-12
<i>Sc</i> Spt7	2276.1 (bdp-2)	S. pombe	4e-38	1e-42	4e-14	No match
ScBdf1	8423.1 (bdp-3)	S. pombe	1e-18	3e-19	4e-18	2e-14
Chromodomain proteins						
DmHP1/SpSwi6	4017.1 (hpo)	S. pombe	No match	3e-8	0.005	No match
CDP-1	8362.1 (cdp-1)	H. sapiens	No match	0.001	5e-7	No match
CDP-2	738.1 (cdp-2)	H. sapiens	No match	No match	0.087	No match
CDP-3	1522.1 (<i>cdp-3</i>)	D. melanogaster	No match	No match	0.001	No match

TABLE 16. Neurospora bromodomain and chromodomain proteins^a

^a This table excludes CRF3-1 (bromodomain) and CRF6-1 and CRF7-1 (chromodomains). There are also several RIP-mutated relics of retrotransposons whose inactive POL genes match chromodomains in BLAST searches.

^b Drosophila melanogaster, Mus musculus, Homo sapiens, Danio rerio, or Caenorhabditis elegans.

^c Arabidopsis thaliana or Oryza sativa.

RIP. The prototypical genome defense system RIP was discovered in 1986 as a result of a detailed analysis of progeny from crosses of Neurospora transformants (Fig. 4) (298, 707, 708, 710, 711, 713). RIP detects duplicated sequences in the haploid genomes of special dikaryotic cells resulting from fertilization (711) and riddles both copies of the duplicated sequence with GC-to-AT mutations (123). In a single passage through the sexual cycle, up to $\sim 30\%$ of the GC pairs in duplicated sequences can be changed to AT pairs (124). Because RIP shows some sequence preference - the C-to-T mutations occur principally at CpA dinucleotides, thus generating an excess of TpA dinucleotides (708) - most genomic regions that have been subjected to RIP are readily recognizable (500). Frequently, but not invariably, cytosines in sequences altered by RIP become methylated de novo. It is possible that the mutations induced by RIP occur by enzymatic deamination of 5-methylcytosines, but they could also occur in other ways, such as by deamination of cytosines followed by DNA replication (306, 509, 708, 845). As a result of the Neurospora genome sequencing projects, a putative DNA methyltransferase (RID) has been recently identified and demonstrated to be essential for RIP (253). RIP appears tailor-made to counter "selfish" DNA such as transposons. Hence, it is not surprising that the great majority of Neurospora strains appear devoid of active transposable elements (417, 419) and that Neurospora contains numerous examples of nonfunctional transposable elements bearing hallmarks of RIP (418, 500, 717).

RNA-dependent silencing. In *Neurospora*, several genes involved in RNA silencing have been identified through a combination of mutant screens and candidate gene approaches. Three genes, *qde-1*, *qde-2*, and *qde-3*, coding respectively for an RNA-dependent RNA polymerase (RdRP), an Argonaute-like protein, and a RecQ-like helicase (131, 132, 159–161, 163), provided the backbone of the vegetative silencing pathway called quelling (Fig. 4). Analysis of these genes was fundamental to our understanding of the RNA silencing machinery in all organisms. A second RdRP, *Suppressor of ascus dominance 1* (*Sad-1*), was identified (732) as a suppressor of a meiotic phenomenon called meiotic silencing (24, 459). Recently, Argonaute-like (related to translation initiation factors) and Dicer-

like (double stranded-specific endonuclease related to the SFII helicase-RNase III fusion protein of the carpel factory) genes, *Suppressor of meiotic silencing 2* (Sms-2) (458) and *Suppressor of meiotic silencing 3* (Sms-3) (M. McLaughlin and R Aramayo, unpublished data), respectively, have been implicated in meiotic silencing. These observations suggested the existence of more than one RNA silencing pathway in *Neurospora*.

Quelling belongs to the broad category of posttranscriptional gene silencing (PTGS) mechanisms (132, 295), and all existing experimental evidence suggests a similar molecular mechanism for meiotic silencing (Fig. 4) (458, 459, 732). These two pathways are thought to be derived from ancestral natural defense systems directed against invading nucleic acids (858). Surprisingly, mutations in all known components of the meiotic silencing pathway (i.e., *sad-1, sms-2*, and *sms-3*) and in some genes of the quelling pathway (e.g., *qde-1*) affect sexual development and/or meiosis, suggesting that in addition to being a defense system, RNA silencing plays a critical role in controlling development.

Pathways similar to quelling and meiotic silencing are also found in plants (PTGS) and animals (RNA interference or RNAi [309]), where they participate in controlling transposon mobilization and development (302, 376, 769). Cosuppression, RNAi, quelling, and meiotic silencing are all related mechanisms that share similar molecular components. These include RdRPs and Argonaute-like and Dicer-like proteins, in addition to many other ATP-dependent DNA and ATP-dependent RNA helicases and double-stranded RNA binding proteins, whose specific function in the silencing pathways remain to be elucidated (359).

This two-pathway hypothesis predicts (i) that there are paralogues in the *Neurospora* genome for genes involved in RNA silencing and (ii) that these paralogues, if functioning in different pathways, should have diverged from a presumed ancestral pathway into phylogenetically distinguishable clades. This hypothesis was tested by searching the *Neurospora* genome for homologues of known components of the RNA silencing machinery. Three RdRPs, two Argonaute-like translation initiation factors, two Dicer-like RNases, and two RecQlike helicases were identified (Table 17). The presence of these

	TABLE	17.	RNA-si	lencing	pathways
--	-------	-----	--------	---------	----------

		01 5		
Predicted protein ^a	N. crassa ^b	A. fumigatus ^c	S. $pombe^d$	Pathway ^e
RNA-directed RNA polymerase	<i>qde-1</i> (NCU07534.1) <i>Sad-1</i> (NCU02178.1) <i>rrp-3</i> (NCU08435.1)	<i>rrpA</i> (contig158) <i>rrpB</i> (contig472)	<i>rdp1</i> ⁺ (SPAC6F12.09)	Quelling Meiotic silencing Unknown
Argonaute-like, related to translation initiation factors	<i>qde-2</i> (NCU04730.1) <i>Sms-2</i> (NCU09434.1)	<i>ppdA</i> (contig720) <i>ppdB</i> (contig196)	ago1+ (SPCC736.11)	Quelling Meiotic silencing
Dicer-like, related to SFII-RN aseIII RNase of the carpel factory	dcl-2 (NCU06766.1) Sms-3 (NCU08270.1)	<i>dclB</i> (contig618) <i>dclA</i> (contig310)	<i>dcr1</i> ⁺ (SPCC584.10C)	Quelling Meiotic silencing
RecQ helicase-like, related to Bloom's and Werner syndrome helicases	<i>qde-3</i> (NCU08598.1) <i>rqh-2</i> (NCU03337.1) ^f	<i>rqhA</i> (contig443) <i>rqhB</i> (contig58) ^f	hus2+ (SPAC2G11.12)	Quelling Unknown

^{*a*} Function predicted based on protein homology.

^b Neurospora crassa Genome Project at http://www-genome.wi.mit.edu.

^c Unfinished Aspergillus fumigatus Genome Project at http://www.tigr.org.

^d Schizosaccharomyces pombe Genome Project at http://www.genedb.org.

^e Pathway assigned based on either known experimental data for *qde-1*, *qde-2*, and *qde-3* (quelling pathway); *Sad-1*, *Sms-2*, and *Sms-3* (meiotic silencing pathway); or predicted based on phylogenetic analysis.

^{*f*} *Rec*Q helicase-like (rqh).

paralogues fulfills the first prediction of the two-pathway hypothesis.

Alignment of the *Neurospora* RdRP, Argonaute-like, and Dicer-like paralogues to their homologues in plants, animals, and other fungi (using neighbor joining [672]) and the Bayesian or most posterior probability methods (354, 355) resulted in trees in which all of the *Neurospora* paralogues clustered with different fungal homologs into two separate clades. This result is consistent with a clade representing independent quelling and meiotic silencing pathways and was used to predict a role for the other fungal homologues (Table 17).

Surprisingly, the predicted *A. fumigatus* homologues also separate into the different clades, suggesting that two mechanisms similar to quelling and meiotic silencing also exist in this organism. This observation is striking for an organism that is not known to have a sexual phase during its life cycle and suggests that perhaps a sexual stage does exist but has missed detection or that such ability was recently lost. *S. pombe* appears to have only one pathway, which clustered consistently with the known *Neurospora* meiotic silencing genes, *Sms-2* and *Sad-1*. Extrapolating this observation onto the phylogenetic results for the Dicer homologs, the genes *Sms-3* and *dcl-2* were predicted to be part of the meiotic silencing and quelling pathways, respectively (Table 17).

One of the *Neurospora* RdRPs (RRP-3) could not be clearly placed in either the quelling or the meiotic silencing pathways. Its presence suggests either that this protein might function in a yet to be discovered silencing pathway or that it forms the remains of an ancestral silencing pathway that may be present in other fungi but that has been lost in *Neurospora* (Table 17).

Thus, *Neurospora* contains paralogues for the fundamental RNA silencing machinery present in plants and animals and these genes have evolved into at least two separate pathways: quelling (vegetative) and meiotic silencing (developmental).

DNA Repair

Repair of DNA damage is essential for the maintenance of genome integrity. DNA damage results from exposure to UV, ionizing radiation, and various chemical mutagens and from errors in DNA replication. Isolation and characterization of strains sensitive to mutagens has led to an understanding of the range of DNA repair mechanisms. These include excision repair, recombination repair, photoreactivation repair, postreplication repair, and mismatch repair, all of which exist in *Neurospora* (287, 364, 692). Another mechanism, known as checkpoint control, affects the efficiency of DNA repair. Checkpoint control mutants are also sensitive to DNA damage.

This section covers all of the above repair systems (recombination repair is described in more detail in the section on meiotic recombination, below). Some of the genes involved in DNA repair were cloned by isolating DNA fragments that complement the sensitivity of damage-sensitive mutants. Other genes were identified by searches of the genome database by using candidate genes from other organisms. In some cases, roles in repair have been confirmed by reverse genetics, while some genes remain as tentative orthologues.

Photoreactivation. Photoreactivation repair of UV damage to DNA is carried out by DNA photolyase (674). The *Neurospora phr* gene encodes a photolyase specific for the cyclobutane pyrimidine dimer (CPD 867). Unlike *Drosophila, Xenopus*, and zebrafish, *Neurospora* does not have a photolyase that repairs a TC(6-4) photoproduct (730).

Excision. The eukaryotic excision repair system has been characterized primarily in S. cerevisiae and in humans and is one of the most fundamental systems that utilizes doublestranded DNA in the repair process (254). Two different excision repair systems have been reported: nucleotide excision repair (NER) and base excision repair (BER). NER, which deals with UV damage and the bulky DNA lesions produced by many chemical mutagens, is processed by a sequentially assembled protein complex. This system is highly conserved across kingdoms from bacteria to humans. Neurospora has two NER systems (319); one is the system conserved in all organisms, and the other is found only in Neurospora and the fission yeast S. pombe. BER, in which a DNA glycosylase specific for a particular lesion produces an apurinic or an apyrimidic site (AP-site) as a first repair step, has been characterized in many organisms.

(i) NER. In S. cerevisiae and human cells, mutants defective in NER genes show extremely high sensitivity to UV and to some chemicals. *Neurospora* NER mutants have been difficult to characterize because *Neurospora* has an additional NER system that is represented by the *mus-18* gene. *Neurospora*

LABLE	18	Nucleotide	eversion	ropair	aonos
IABLE	10.	Nucleotide	excision	repair	genes

-

S. cerevisiae gene	Function	Neurospora NCU no. (gene)	Contig	E value	Best match to <i>Neurospora</i> protein in SP + TrEMBL	E value (BLASTP)
RAD1	5' endonuclease	07440.1 (<i>mus-38</i>)	3.434	e-140	S. pombe Rad16 M. musculus XPF Human XPF	0.0 e-141 e-140
RAD2	3' endonuclease	07498.1 (<i>mus-40</i>)	3.439	1e-72	S cerevisiae Rad1p S. pombe Rad13 S. cerevisiae Rad2p	e-134 1e-87 9e-68
RAD3	5'-3' helicase	01625.1	3.67	0.0	Human XPG S. pombe Rad15 S. cerevisiae Rad3p Human XPD	5e-55 0.0 0.0
RAD4	GGR-specific damage recognition factor	06585.1	3.381	2e-28	S. pombe Rad4 homologue Drosophila MUS210 Human XPC	2e-62 5e-26 8e-22
RAD7	DNA damage recognition	03649.1	3.199	4e-11	S. cerevisiae Rad4p S. pombe Rad7 homologue S. cerevisiae Rad7p Human FBL6	3e-20 2e-41 4e-14 3e-06
RAD10	5' endonuclease	07066.1 (<i>mus-44</i>)	3.413	8e-16	<i>S. pombe</i> Swi10 Rice Rad10 homologue Human ERCC1	3e-53 2e-43 5e-42
RAD14	Damaged DNA binding activity	08742.1 (<i>mus-43</i>)	3.541	3e-34	S. cerevisiae Rad10p S. pombe Rad14 homologue S. cerevisiae Rad14p Humon XPA	2e-13 1e-42 4e-31 3e 15
RAD16	DNA damage recognition, DNA-dependent ATPase activity	03650.1	3.199	0.0	S. cerevisiae Rad16p S. pombe Rhp16 Human Rad16 homologue	0.0 0.0 2e-50
RAD23	DNA damage recognition	07542.1	3.442	8e-31	S. pombe Rad23 homologue A. thaliana Rad23 homologue Human HHR23B Human HHR23A S. cerevisiae Rad23n	9e-58 1e-39 8e-32 3e-21 2e-29
RAD25/SSL2	DNA helicase activity, TFIIH complex	06438.1	3.371	0.0	S. pombe Rad25 homologue S. cerevisie Rad25p Human XPB	0.0
RAD26	DNA-dependent ATPase activity	07837.1	3.471	0.0	S. pombe Rhp26 S. cerevisie Rad26p Human FRCC6/CSB	0.0 0.0 e-154
RAD28 S. pombe uve1 ⁺	Unknown UV damage-dependent endonuclease	None 08850.1 (<i>mus-18</i>)	3.550	8e-99	S. pombe Uve1 Bacillus UVSE	0.0 1e-30
Human DDB1	UV-damaged DNA binding protein	06605.1	3.382	1e-92	<i>O. sativa</i> OSUVDDB <i>A. thaliana</i> AT4G05420 Green monkey DDB1 Human DDB1	e-122 e-108 7e-87 2e-86

MUS18 and *S. pombe* UVDE are UV damage-specific endonucleases able to nick DNA to the 5' side of UV damage to initiate excision repair (868, 887). After nicking of singlestranded DNA, a polynucleotide, including the damaged base, is removed by a flap endonuclease, Fen1, in *S. pombe* (889). In *S. pombe*, UVDE is involved in repair of DNA damage in mitochondria, as well as in the nucleus (880). However, the roles of Fen1 in excision repair and of MUS18 in repair of mitochondrial DNA damage remain to be established in *Neurospora*.

Genes in the primary NER pathway have been characterized in *Neurospora* only in the last 10 years, after the discovery of the second, *mus-18* NER pathway. *mus-38* and *mus-40* respectively encode homologues of the *S. cerevisiae* genes *RAD1* and *RAD2*. A mutation in either of these genes results in mild sensitivity to UV radiation in a normal genetic background but a synergistic sensitivity to UV light in *mus-18* strains. All other NER genes that comprise the conserved pathway are present in *Neurospora*, and their amino acid sequences are similar to those of the two sequenced yeasts, particularly *S. pombe* (Table 18).

(ii) **BER.** BER has not been studied in *Neurospora*, since no suitable mutants have been isolated. *Neurospora* orthologues of BER genes have been identified on the basis of sequence homology, although homologies are usually not high (Table 19). Indeed, sequences of these genes appear highly divergent from organism to organism. Confirmation of a role in BER for the *Neurospora* orthologues awaits the analysis of disruption mutants.

Recombination repair. Recombination repair genes are separated into two groups: homologous recombination (HR) and nonhomologous/end joining (NHEJ). Homologous recombination is covered in the Meiotic Recombination section below.

S. cerevisiae gene	Function	<i>Neurospora</i> NCU no.	Contig	E value	Best match to <i>Neurospora</i> protein in SP + TrEMBL	E value (BLASTP)
APN1	AP endonuclease	10044.1	3.748	2e-78	S. cerevisiae Apn1p	9e-81
					Plasmodium Apn1	5e-69
MAG1	3-Methyladenine DNA	08938.1	3.559	6e-21	O. sativa Mag1 homologue	1e-18
	glycosylase				Halobacterium Mag1 homologue	7e-18
					S. cerevisiae Mag1p	6e-12
NTG1	DNA N-glycosylase	06654.1	3.385	2e-44	S. pombe Nth1	5e-62
					Drosophila Nth1 homologue	6e-57
					Human NTH1	2e-50
OGG1	8-Oxo-guanine DNA glycosylase	03040.1	3.154	3e-33	Rat OGG1	3e-39
					M. musculus OGG1	2e-38
					Human OGG1	6e-38
					S. cerevisiae Ogg1p	3e-31
UNG1	Uracil N-glycosylase	07482.1	3.439	4e-65	S. pombe Ung1	1e-86
					S. cerevisiae Ung1p	5e-68
					Human UNG	1e-52
Human PARP-1	Poly(ADP-ribose) polymerase	08852.1	3.550	2e-63	A. thaliana app	8e-66
	activity, protein modification				Dictyostelium ADPRT2	2e-62
	-				Human ADPRT	4e-60

TABLE 19. Base excision repair genes

The human Nijmegen breakage syndrome (NBS1) protein has weak homology to the XRS2 gene product of S. cerevisiae (508). While a possible Neurospora orthologue of human NBS1 was identified, the similarity is very poor (see Table 24), consistent with the observation that Xrs2 orthologues are highly divergent between species. mus-45 mutants have a similar phenotype to that of uvs-6 (yeast rad50 homologue) and mus-23 (S. cerevisiae mre11 homologue) mutants, and mus-45 is likely to be a true orthologue of NBS1 (H. Inoue, unpublished data). Orthologues of the KU70 and KU80 proteins, which bind to DNA double-strand breaks (DSBs), are also found in *Neurospora*. However, the DNA protein kinase (DNA-PKcs) gene involved in NHEJ in humans is not present in *Neurospora* or yeasts.

Postreplication repair. The postreplication repair pathway includes two ubiquitin-conjugating enzymes and two ubiquitin ligases which have ring finger motifs, suggesting that the main function of this repair system may be related to a ubiquitin-control system. *Neurospora* has orthologues of the budding

S. cerevisiae	Function	<i>Neurospora</i> homologue	Contig	E value	Best match to <i>Neurospora</i>	E value (BLASTP)
		nomologue				(BE#1011)
RAD5	RING finger protein, DNA	NCU09516.1 <i>mus-41</i>	3.611	e-120	S. pombe Rad8	0.0
	helicase				S. cerevisiae Rado	e-143
DID	тп	NCI 100721 1 0	2 (2)		Human HIP116	9e-76
<i>KAD</i> 6	ubiquitin-conjugating enzyme, required for histone ubiquitination, protein monoubiquitination, and ubiquitin-dependent protein catabolism	NCU09/31.1 mus-8	3.634	/e-6/	N. haematococca NhKADb	66-83
					E. nidulans UVSJ	5e-80
					S. pombe Rhp6	2e-70
					S. cerevisiae Rad6	5e-69
					Human HHR6B	4e-61
					Human HHR6A	2e-60
RAD18	RING finger protein, ATPase	NCU05210.1 uvs-2	3.293	7e-31	E. nidulans NUVA	1e-74
	activity, single-stranded DNA				S. pombe Rhp18	3e-40
	binding activity				S. cerevisiae Rad18	4e-29
					Human hRAD18	8e-27
UBC13	Ubiquitin-conjugating enzyme,	NCU02113.1	3.95	9e-63	Catharanthus CrUBIE2	2e-62
	heterodimer with Mms2				A. thaliana AY049261	3e-62
					S. cerevisiae Ubc13	3e-61
					S. pombe Spu13	3e-60
					Human UBE2N	4e-60
MMS2	Ubiquitin-conjugating enzyme,	No number	Excluded contig		K. delphensis MMS2	2e-39
	heterodimer with Ubc13		3.72		S. pombe Spm2	1e-37
					S. cerevisiae Mms2	2e-35
					Human hMMS2	4e-34
					Human CROC-1B	4e-34

TABLE 20. RAD6 DNA repair genes

S. cerevisiae gene	Function	Neurospora homologue	Contig	E value	Best match to <i>Neurospora</i> protein in SP + TrEMBL	E value (BLASTP)
SRS2	DNA helicase; recombination	NCU04733.1	3.260	3e-76	S. pombe Srs2	e-117
	suppression, DSBR via NHEJ				L. citreum PcrA	7e-93
					E. coli UVRD	1e-78
					S. cerevisiae Srs2	4e-70
MGS1	DNA-dependent AAA(+)	NCU08706.1	3.541	5e-95	S. pombe putative protein	e-107
	ATPase; helicase activity				A. thaliana AC002396	3e-94
					Human WHIP	9e-92
					S. cerevisiae Mgs1	1e-82
POL30/PCNA	DNA polymerase processivity	NCU09239.1	3.579	1e-67	S. pombe PCNA	e-100
	factor				D. carota PCNA	9e-83
					Human PCNA	1e-78
					S. cerevisiae Pol30	4e-60
Human p66	DNA polymerase δ subunit	NCU07998.1	3.481	1e-07	Human p66	7e-06
					M. musculus p66	2e-05
					S. pombe Cdc27	0.28
CTF4	Polα accessory subunit, DNA	NCU08484.1 mus-27	3.513	1e-50	E. nidulans SepB	0.0
	binding activity				S. pombe Ctf4 homolog	e-154
					Human AND1	1e-40
					S. cerevisiae Ctf4	2e-36
RAD27	Flap endonuclease activity,	NCU02288.1	3.108	e-113	S. pombe Rad2	e-126
	Okazaki fragment processing				Xenopus Rad27	e-113
					S. cerevisiae Rad27	e-113
					Human FEN1	e-108

TABLE 21. DNA replication-related repair genes

yeast *RAD6* and *MMS2/UBC13* genes, which encode ubiquitinconjugating enzymes, and of *RAD18* and *RAD5*, which are ubiquitin ligase genes (Table 20). An orthologue of budding yeast Srs2p, a helicase that functions in postreplication repair, is present in *S. pombe* and *Neurospora* but not in plants and animals (Table 21).

The postreplication repair group also includes specific DNA polymerases that function in translesion DNA synthesis. Among these are human polymerase κ (kappa), polymerase ζ (zeta), polymerase ε (epsilon), and polymerase ι (iota) (628). Although an orthologue of DNA polymerase κ is absent from *S. cerevisiae* (but present in *S. pombe*) and both yeast species lack polymerase ι , orthologues of these polymerases exist in *Neurospora*. Some *Neurospora* polymerase orthologues are more similar to those of human than those of *S. cerevisiae* (Table 22). Except for the *pol* ζ homologue *upr-1* (673), the functions of *pol* gene products are not well understood in *Neurospora*.

Checkpoint control. Neurospora has orthologues of almost all yeast DNA damage checkpoint genes, although some of them do not show a high degree of sequence similarity (Table 23). Some of the mutagen-sensitive Neurospora mutants that have not yet been characterized at the sequence level may be included in this group. The multinucleate nature of Neurospora cells makes it difficult to detect damage checkpoint defects, since nuclear divisions are not synchronized, even in the same cellular compartment. However, the similarity between deduced amino acid sequences and observed mutant phenotypes made it possible to designate Neurospora mus-9, uvs-3, and mus-21 as orthologues of MEC1, DDC2, and TEL1, respectively, in S. cerevisiae (C. Ishii, unpublished data).

In conclusion, many *Neurospora* repair proteins are more similar to those of *S. pombe* than to *S. cerevisiae*, while others are more closely related to proteins from higher organisms such as humans. In some cases, the phenotype of a *Neurospora* mutant deficient in repair differs from that of a yeast strain carrying a mutation in the orthologous gene (for example, *srs-2*, *fen1*, and *mus-21* [H. Inoue, unpublished]). Some members of repair gene families that are found in other organisms are not found in *Neurospora* (for example, Rad59p of *S. cerevisiae*), likely due to constraints imposed upon gene duplication by RIP.

Meiotic Recombination

S. cerevisiae exhibits very high rates of recombination and has a highly tractable genome; much of what is known about both the mechanism and enzymology of meiotic recombination derives from studies of this species. Indeed, organisms other than *S. cerevisiae* appear to harbor very few meiotic recombination genes that it does not also possess (Table 24). The reverse is not true, however, since *S. cerevisiae* appears to contain quite a few genes not found in other organisms. Moreover, it is likely that numerous recombination genes remain to be identified in nonyeast organisms because, in relative terms, less effort has gone into their cloning.

Meiotic recombination is thought to begin with a DSB in the DNA duplex. This is followed by 5'-to-3' degradation either side of the DSB, yielding single-stranded tails. These singlestranded tails then invade the intact homologue, and the formation of a recombination intermediate, which holds homologues together, is thought to follow. The formation of a proteinaceous structure, the synaptonemal complex, between homologues signals synapsis. Sometime prior to the separation of homologues at anaphase I, the recombination intermediate is resolved. Resolution may yield homologues with or without a crossover. While all of this is going on, mismatched bases are detected and usually corrected. Although this account of meiotic recombination is not necessarily chronologically accurate, it does provide a framework around which a discussion of the numerous genes known to be involved in meiotic recombina-

S. cerevisiae gene	Function	Neurospora NCU no.	Contig	E value	Best match to <i>Neurospora</i> protein in SP + TrEMBL	E value (BLASTP)
POL1	Polα catalytic subunit, priming activity in DNA replication	07870.1	3.473	0.0	S. pombe Poll C. cinereus Poll	$\begin{array}{c} 0.0\\ 0.0\end{array}$
POL4	Polβ catalytic subunit, BER in nuclear DNA	07461.1	3.453	3e-11	<i>S. cereviciae</i> Pollp Human POLL <i>M. musculus</i> POLL	0.0 5e-52 1e-49
	Pola meiosis-associated DNA repair	01321.1	3.51	9e-07	<i>S. cerevisiae</i> Pol4p <i>S. pombe</i> Polβ-like Human POLM	2e-09 1e-39 5e-36
MIP1	Poly, BER in mitochondrial DNA	00279.1	3.12	0.0	S. cerevisiae Pol4p S. pombe Mip1 P. pastoris Mip1 S. cerevisiae Mip1p	0.002 0.0 0.0 0.0
POL3	Polò catalytic subunit, NER and MMR	01192.1	3.45	0.0	Human POLG S. pombe Pol3 C. albicans Pol3	e-118 0.0 0.0
POL2	Pole catalytic subunit, NER and MMR	04548.1	3.233	0.0	S. cerevisiae Pol3p Human POL3 E. nidulans NimP S. pombe Cdc20	$0.0 \\ 0.0 \\ 0.0 \\ 0.0 \\ 0.0$
REV1	dCTP transferase, TLS	02053.1 mus-42	3.90	3e-63	<i>S. cerevisiae</i> Pol2p Human POLE1 Human REV1 <i>M. musculus</i> REV1	0.0 0.0 e-100 4e-95
	Роlк, TLS	02457.1	3.131	1e-09	<i>S. cerevisiae</i> Rev1p <i>P. anserina</i> CAD60612 <i>S. pombe</i> O74944	6e-57 0.0 2e-92
REV3	Polζ catalytic subunit, TLS	01951.1 upr-1	3.86	1e-62	Human POLK E. nidulans UvsI S. cerevisiae Rev3p	1e-43 0.0 0.0
REV7	Polζ accessory subunit	06577.1 mus-26	3.381	0.017	S. pombe Mad2-like M. musculus Mad2L	0.0 7e-13 3e-08 2e.07
RAD30	Poly, TLS	01936.1 polh	3.84	2e-54	S. pombe Eso1 S. cerevisiae Rad30p Human XPV	2e-07 2e-95 9e-53 2e-40
	Polt, TLS	06757.1 <i>poli</i>	3.390	5e-07	Human POLI M. musculus POLI S. pombe Q74944	4e-32 4e-31 2e-15
Human POLQ	Polθ, DNA cross-link repair	07411.1	3.432	5e-85	Drosophila Mus308 Human POLQ M. musculus POLO	e-112 e-106 e-104
TRF4	Polo, nucleotidyltransferase, sister chromotid cohesion	05588.1	3.312	6e-38	S. cerevisiae Trf4 S. cerevisiae Trf5p Human TRF4-1	3e-37 2e-33 2e-26
	Polo, TRF5?	00538.1	3.20	3e-09	S. pombe Cid13 S. pombe AL031154 Human KIAA0191 S. cerevisiae Trf5p	1e-77 5e-60 1e-27 4e-8

TABLE 22. Repair-related DNA polymerases

tion can be organized. However, it should be borne in mind that the influence of some genes is not restricted to a given step.

Before the DSB. While the generation of DSBs is thought to be the initiating event of meiotic recombination, it appears that in *S. cerevisiae*, premeiotic DNA replication is a necessary precondition for DSB genesis. Smith et al. (743) showed that mutation of the B-type cyclins *CLB5* and *CLB6* prevented premeiotic DNA replication, which in turn blocked the formation of DSBs. The situation seems to be similar in *Arabidopsis*. Mutation of a meiosis-specific cyclin Solo dancers (*sds*) in *Arabidopsis* results in defects in homologue pairing and in recombination (36). While both of the *S. cerevisiae* cyclins (Clb5/Clb6) and the sds protein have limited homology to

several hypothetical *Neurospora* proteins (Table 24), homology is largely restricted to mitosis-specific cyclins from other organisms. A fully functional Spo22 appears to be required for premeiotic DNA synthesis in *Coprinus* (523). Thus, it remains to be established as to whether one or more of the hypothetical *Neurospora* cyclins has any effect on meiotic recombination.

DSB generation. Spo11p is a novel type II topoisomerase (68) that is thought to cleave DNA, generating the DSBs considered to be the universal initiating event of meiotic recombination. *SPO11* was first identified in *S. cerevisiae* (408) and has orthologues in fission yeast (*rec-12*), *Coprinus, Neurospora, Sordaria*, nematodes, *Drosophila* (*mei-W68*), *Arabidopsis* (*Spo11-1*), mice, and humans. In fact, it seems that wherever *SPO11* has been sought in eukaryotes, it has been found. It is

S. cerevisiae gene	Function	<i>Neurospora</i> homologue	Contig	E value	Best match to <i>Neurospora</i> protein in SP + TrEMBL	E value (BLASTP)
RAD17	DNA damage checkpoint, control protein, 3'-5' exonuclease (putative)	NCU00942.1	3.37	9e-05	Xenopus RAD1 M. musculus REC1 S. pombe Rad1	9e-17 4e-16 5e-15
MEC1	DNA damage checkpoint, inositol/ phosphatidylinositol kinase activity	NCU00625.1 mus-9	3.23	e-135	S. cerevisiae Rad1/ E. nidulans UVSB S. pombe Rad3 Human ATR	0.009 0.0 0.0 e-146
DPB11	S-phase checkpoint, epsilon DNA polymerase activity	NCU09503.1	3.610	8e-04	S. cerevisiae Mec1 S pombe Rad4 Human KIAA0259 S. cerevisiae Dpb11	e-130 2e-55 6e-15 1e-08
S. pombe rad9 ⁺	Cell cycle checkpoint	NCU00470.1	3.15	2e-18	S. octosporus homolog Human RAD9 homolog	5e-16 7e-16 5e-09
RAD24	DNA damage checkpoint, DNA clamp loader activity	NCU00517.1	3.18	2e-22	M. musculus RAD17 Green monkey RAD17 Human RAD17 S pombe Rad17 S cerevisiae Rad24	7e-2 2e-22 3e-22 2e-14
E. nidulans uvsD (DDC2/PIE1/ LCD1)	DNA damage checkpoint, damaged DNA binding activity, protein binding activity	NCU09644.1 uvs-3	3.627	8e-30	E. nidulans UVSD	4e-22
CHK1	DNA damage checkpoint, protein kinase activity	NCU08346.1 un-1	3.501	2e-68	S. pombe Chk1 S. cerevisiae Chk1 Human CHK1	1e-82 5e-63 3e-43
RAD9	DNA damage checkpoint, protein binding activity	NCU08879.1	3.553	1e-06	S. pombe RHP9 Leishmania PPG3 S. cerevisiae Rad9	1e-12 2e-07 6e-05
RAD53/MEC2/ SPK1/LSD1	DNA repair, protein threonine/ tyrosine kinase	NCU02751.1	3.140	6e-50	S. cerevisiae Rad53 S. pombe Cds1 Human CAMK1	1e-46 4e-36 2e-32
DUN1	DNA damage checkpoint, protein kinase activity	NCU02814.1	3.146	9e-72	S. cerevisiae Dun1 Human CHK2	7e-72 2e-71
S. pombe hus1 ⁺	DNA damage checkpoint, DNA binding activity	NCU03820.1	3.203	8e-30	A. nidulans NUV101 S. pombe Hus1 M. musculus HUS1	2e-56 9e-44 1e-10
TEL1	Response to DNA damage, inositol/ phosphatidylinositol kinase activity	NCU00274.1 mus-21	3.12	e-112	S. pombe Tell Xenopus ATM Human ATM S. cerevisiae Tell	e-158 e-116 e-104 5e-97
PTC2	DNA damage checkpoint, protein phosphatase type 2C activity	NCU04600.1	3.239	2e-79	S. corevisiae Ptc3 S. pombe Ptc3 S. cerevisiae Ptc3 Human PPM1A	e-121 e-117 4e-79 4e-51
RFX1/CRT1	DNA damage checkpoint, specific transcriptional repressor activity	NCU06701.1	3.387	6e-29	Cephalosporium CPCR1 Penicillium RFX1 S. cerevisiae Rfx1 Human RFX1	e-169 e-102 2e-36 2e-30
ASF1	Histone binding, DNA damage response, signal transduction	NCU09436.1	3.602	8e-65	S. cerevisiae Asf1 S. pombe Cia1 Human HSPC146	1e-61 2e-54 2e-45

TABLE 23. DNA damage checkpoint genes

perhaps a little surprising that with such a widely conserved and important function the amino acid sequence is not highly conserved. *Neurospora spo-11* is quite similar to its *Coprinus*, *S. pombe*, and human orthologues but has limited homology to budding yeast Spo11p (Table 24). However, all Spo11 orthologues possess five conserved motifs containing a number of invariant amino acids. While the predicted amino acid sequence for the *Neurospora* orthologue appears to lack important residues in motif 3, this is probably an artifact of the gene prediction algorithm. Sequencing of part of the *Neurospora* cDNA indicates that these "missing" residues are in fact present (F. J. Bowring, P. J. Yeadon, R. G. Stainer, and D. E. A. Catcheside, unpublished data). *S. cerevisiae* Spo11p requires a number of accessory proteins such as Mei4p, Mer2p, Rec102p, Rec104p, and Rec114p (reviewed in reference 407) for the formation of DSBs. Apart from modest similarity between part of the Rec114p protein sequence and *S. pombe* Rec7, these accessory proteins do not appear to have counterparts in any of the other Spo11p-containing organisms. Possibly some of the *S. cerevisiae* Spo11p function is carried out by one or more of these accessory proteins, or perhaps one or more of these proteins is responsible for the rather high recombination frequencies typical of this yeast. Rec103p, probably also required for DSB generation (273), does have orthologues in *Neurospora* and other organisms (Table 24).

S. cerevisiae gene ^b	Neurospora homologue	E value	Best match to protein in SP + TrEMBL	E value (BLASTP)
Before the DSB CLB5 CLB6 sds (Arabidopsis)	NCU02758.1 hyp protein	3e-51 5e-59 2e-11	<i>A. nidulans</i> G ₂ /mitosis-specific cyclin B <i>S. pombe</i> G ₂ /mitosis-specific cyclin <i>cdc13</i> <i>S. cerevisiae</i> G ₂ /mitosis-specific cyclin 2 Human G ₂ /mitosis-specific cyclin B2	2.1e-161 3.6e-129 1.0e-106 7.9e-75
DSB generation				0.00
HOP2	None		S. pombe MEI13 P. falciparum hyp. protein Human BRCA1-associated protein	8e-08 8e-04 9e-04
MEI4	None	0.04	None	1 50
MEK1/MRE4	NCU02814.1	2e-36	S. cerevisiae DUNI Human CHK2	1e-72 4e-72
			S. cerevisiae MEK1	1.5e-27
MER1	None		None Similarity of the second se	
MER2/REC10/ MER3	None NCU09793.1 hvp. protein	e-148	Similarity only to myosin neavy-chain genes S. cerevisiae HFM1/MER3	7.4e-170
101110	recover hyp. protein	0 110	A. thaliana genomic DNA	7.1e-93
			D. melanogaster GH18520 full-length cDNA	1.3e-65
MRF2/NAM8	NCU00768 1	8e-55	Human mRNA for putative RNA helicase, 3' end A thaliana at5x54900	1.0e-52 3e-67
	110000700.1	00 33	Nicotiana tabacum putative RNA binding protein	4e-65
			S. cerevisiae NAM8	2e-58
REC102	None		None	5e-37
REC103/SK18	NCU03517.1 hyp. protein	2.00e-17	S. pombe rec14	1.0e-71
DEC104	NT		Human <i>REC14</i>	1.2e-25
REC104 REC114	None		Some homology to <i>S</i> pombe Rec7	
RED1	None		Kluyveromyces lactis red1	2e-70
(0.0.11		0 10	Human KIAA0874 protein	4e-06
SPOII	NCU01120.1 S. pombe	2e-10	Coprinus cinereus spoll S. pombe recl?	3e-29 2e-28
	reerz related protein		Human SPO11	3e-25
			S. cerevisiae SPO11	1e-08
Removal of Spo11 protein				
from DNA				
MRE11	NCU08730.1 mus-23	e-123	Magnaporthe grisea (NK73)	e-178
	(AD002350)		S. cerevisiae MRE11	e-118
			Human MRE11	e-103
RAD50	NCU00901.1 uvs-6 (AB055069)	e-175	S. cerevisiae RAD50 Mouse R4D50	e-175 e-159
	(11005000))		Human RAD50	e-159
SAE2/COM1	None		None	
Resection of ends				
XRS2	None		P. falciparum e1-e2 putative ATPase/hydrolase	8e-06
Human NDS1	NCU04220 1	20.05	Human neurofilament, heavy polypeptide	4e-05
	NCU04329.1	36-03	S. cerevisiae LP19	3e-110
Strand invasion	None		Widely conserved including mammals	
RAD51	NCU02741.1 mei-3	e-123	A. nidulans uvsC	e-178
			Penicillium paxilli pp rad51	e-174
			Human RAD51 S. cerevisiae R4D51	e-135 e-131
RAD52	NCU04275.1 mus-11	5e-46	M. grisea NK72	4e-95
			A. nidulans radC	2e-67
			S. cerevisiae RAD52 Human RAD52	6e-28
RAD54	NCU02348.1 (AB032901)	0.0	M. grisea NK74	0.0
	mus-25		S. pombe RAD54	0.0
			S. cerevisiae RAD54	0.0
			11ulliali IVAD34	0.0

TABLE 24. Genes involved in DSB initiation and processing^a

Continued on following page

S. cerevisiae gene ^{b}	Neurospora homologue	E value	Best match to protein in SP + TrEMBL	E value (BLASTP)
RAD55	NCU08806.1	7e-05	Methanopyrus kandleri rada S. pombe rhp55	7e-04 0.001
RAD57	NCU01771.1 <i>RAD57</i> homologue	2e-23	S. cerevisiae RADSS S. pombe RHP57 S. cerevisiae RAD57	0.003 2e-32 1e-16
RDH54/TID1	NCU02348.1 <i>RAD54</i> homologue	e-107	Human XRCC3	6e-12
	NCU07837.1	1e-57	S. pombe RHP26 S. cerevisiae RAD26 Human ercc-6	0.0 0.0 e-156
RFA1	NCU03606.1	e-115	S. cerevisiae RDH54 S. pombe RFA1 Human RFA1	6.8e-51 e-170 e-119
RFA2	NCU07717.1	2e-23	S. cereviside RFA1 A. fumigatus possible Rfa S. pombe RFA2 Human RFA2	e-114 2e-34 1e-28 2e-16
RFA3 SAE3	None None		S. cerevisiae RFA2 None None	7e-13
Synapsis and synaptonemal complex formation				
HOP1	None		<i>K. lactis HOP1</i> Poorer matches (6e-14 to e-07) to proteins involved in synapsis in plants and animals	1e-93
MND1			Human GAJ. Also mouse, A. thaliana, and S. pombe	1e-17 to 1e-08
ZIP1	NCU00658.1	8e-24	Rat Golgi complex-associated protein Human centromeric protein S. cerevisiae ZIP1	2e-75 3e-69 2e-21
ZIP2	None		None	
Regulation of crossover frequency				
ME15 MLH1	None NCU08309.1	e-159	Human ninein (<i>GSK3B</i> -interacting protein) <i>S. cerevisiae MLH1</i> Human <i>MLH1</i>	2e-04 e-168 e-167
MLH3	NCU08309.1 <i>mlh-1</i> NCU08020.1 <i>pms-1</i> NCU09272.1 <i>mlh</i> -2	5e-23 9e-18		
	NCU05385.1	2e-09	S. cerevisiae MLH3 A. thaliana PMS1 Mouse PMS2 Human PMS2	3.4e-13 2e-09 3e-08 2e-07
MSH4	NCU02230.1 msh-2 NCU08115.1 msh-3 NCU08135.1 msh-6 NCU09384.1 msh-5 NCU07407.1 msh-1	1e-40 8e-32 8e-27 3e-22 6e-18		
	Contig 3.27 (scaffold 2)	2e-36 (TBLASTN)	Mouse MSH4 Thermotoga maritima MutS S. cerevisiae MSH4	4e-39 4e-39 2e-35
MSH5	NCU09384.1	1e-58	Human <i>MSH4</i> S. cerevisiae MSH5 Human <i>MSH5</i>	3e-34 1e-55 4e-42
TAM1/NDJ1 Drosophila mei-9	None NCU07440.1 <i>mus-38</i>	e-119	None S. pombe chromosome III cosmid Mouse DNA repair endonuclease XPF Human excision repair S. cerevisiae RAD1	2e-270 5.3e-199 2.2e-197 5.9e-170
Drosophila mei-218	None		A. thaliana 5' repair endonuclease None	1.5e-139
Mismatch repair <i>MLH2</i>	NCU08020.1 pms-1	6e-19		

TABLE 24—Continued

Continued on following page

S. cerevisiae gene ^b	Neurospora homologue	E value	Best match to protein in SP + TrEMBL	E value (BLASTP)
		6 16		
	NCU08309.1 <i>mlh-1</i>	6e-16	Human BMS1	40.24
	NC009373.1 min-2	46-15	Phaeosphaeria nodorum hyp. protein	40-24 5e-24
			S cerevisiae MLH2	5 0e-15
MSH2	NCU02230.1 msh-2	0.0	S. cerevisiae MSH2	0.0
1110112	110002230.1 man 2	0.0	Human <i>MSH2</i>	0.0
MSH3	NCU08115.1 msh-3	e-115	Human MSH3	e-168
			Mouse MSH3	e-164
			S. cerevisiae MSH3	e-105
MSH6	NCU08135.1 msh-6	0.0	S. pombe MSH6	0.0
			S. cerevisiae MSH6	0.0
			Human MSH6	e-157
PMS1	NCU08020.1 pms-1	8e-82	S. pombe PMS1	5e-83
			S. cerevisiae PMS1	8e-81
			Human PMS2	4e-80
Resolution of recombina-				
tion intermediates				
MMS4/SLX2	NCU04047.1	2e-04	Xenopus laevis nucleolar phosphoprotein	5e-04
			Human nucleolar phosphoprotein	0.002
SLXI	NCU01236.1	2e-26	S. cerevisiae SLX1	2e-23
			S. pombe hyp. protein	5e-17
CI 322/10/1001		0 50	Human hyp. protein MGC5178	7e-12
SLX3/MUS81	NCU0/45/.1	2e-52	S. cerevisiae MUS81	8.2e-50
			S. pombe hyp. protein	Se-76
SI VA	None		Human MUS81	2e-36
SLX4 SLX9	None NCLI02272 1	2. 07	Human SMC4	2e-05
SLAO	NC003872.1	2e-07	S. pombe zinc inger protein	7- 20
			S. cerevisiae Chr All sequence	/e-39
			$\begin{array}{c} Fullman \ K N F 10 \\ S \ commission \ S I \ V S \end{array}$	40.05
HEV2/SI V5	None		S. CELEVISIUE SLAO	46-03
TOP1/MAK1/MAK17	NCU00118 1	0.0	A nidulans TOP1	0.0
1011/maki/maki/	NC009118.1	0.0	A. manunis 1011 S. pombe TOP1	0.0
			S. cerevisiae TOP1	0.0
			Human TOP1	e-114
TOP2/TOR3/TRF3	NCU06338 1	0.0	A niger TOP?	0.0
1012/1010/1103	100000001	0.0	Penicillium chrysogenum TOP2	0.0
			S cerevisiae TOP2	0.0
			Human TOP2	0.0
TOP3/EDR1	NCU00081.1	e-114	S. pombe topoisomerase 3	e-115
,			C. elegans topoisomerase I	e-105
			Human topoisomerase III	e-104
			S. cerevisiae TOP3	e-101
Nonhomologous end joining				
LIF1	None		Human hyaluronan-mediated motility receptor	4e-05
			(RHAMM) isoform A	
LIG4	NCU06264.1	7e-97	S. pombe DNA ligase 4	e-143
			C. albicans LIG4	1e-94
			Human ligase IV	3e-89
			S. cerevisiae LIG4	1e-88
YKU70/HDF1/NES24	NCU08290.1	1e-27	S. pombe putative DNA helicase	3e-91
			Gallus gallus ku70	1e-58
			Human <i>ku70</i>	6e-51
			S. cerevisiae YKU70	4e-25
YKU80/HDF2	NCU00077.1	2e-14	S. pombe putative DNA helicase	3e-73
			Mouse Ku80	2e-26
			Human DNA helicase	9e-26
			S. cerevisiae HDF2	8e-12

TABLE 24—Continued

^{*a*} Searches of the literature and of the Saccharomyces database (http://genome-www4.stanford.edu/cgi-bin/SGD/SAGE/querySAGE) were used to identify candidate genes. The resulting amino acid sequences were used to find orthologues in the *Neurospora* predicted protein database. Where this approach yielded nothing, a TBLASTN search of the whole *Neurospora* genome sequence was performed, thus allowing us to locate *msh-4*. The amino acid sequence of the best match for each gene was used to search the Swiss Prot + TrEMBL nonredundant protein databases. For *msh-4*, the *Neurospora* nucleotide sequence was used in a BLASTS search against Swiss Prot + TrEMBL. *S. cerevisiae* genes with no *Neurospora* orthologue were used to search Swiss Prot + TrEMBL or the human genome protein database (http://www.ncbi.nih.gov/genome/seq/HsBlast.html).
Red1p, a component of the synaptonemal complex, and Hop2p, a protein that discourages recombination between nonhomologous chromosomes, are both required for normal levels of DSB formation in *S. cerevisiae*. Neither has an orthologue in *Neurospora* or higher eukaryotes. Mek1p is a kinase that phosphorylates Red1p and is also required for full levels of DSBs in *S. cerevisiae*. Curiously, while both *Neurospora* and humans have a Mek1p equivalent, both apparently lack an orthologue of its substrate, Red1p.

As indicated above, Mer2p appears to be unique to *S. cerevisiae*. Splicing of *MER2* and *MER3* RNA is regulated by two genes, *MER1* and *MRE2* (534, 555). *MER3* codes for a putative helicase thought to be involved in the processing of DSBs and in interference (555) and has orthologues in *Neurospora* and other organisms (Table 24). Although *MER1* appears to be unique to budding yeast, there are *MRE2* orthologues in *Neurospora*, *Arabidopsis*, and humans (Table 24). It is possible that Mre2p plays a role in the splicing of transcripts from their *MER3* equivalents in *Neurospora* and other organisms lacking *MER2*.

Removal of Spo11 protein from DNA. The *S. cerevisiae* Rad50p, Mre11p, and Sae2p/Com1p proteins are implicated in the removal of Spo11p from DNA following DSB genesis. While Rad50p and Mre11p are widely conserved (Table 24), Sae2p/Com1p appears to be unique to *S. cerevisiae*.

Resection of ends. Rad50p and Mre11p are also required for post-scission 5'-to-3' degradation, yielding 3' single-stranded tails on either side of the DSB. While it is not known whether Xrs2p participates in resection, Rad50p, Mre11p, and Xrs2p are known to form a complex. Apart from weak homology to a putative protein from *Plasmodium*, Xrs2p does not appear to have potential orthologues in other organisms. Human NBS1 and budding yeast Xrs2p have little sequence homology, but NBS1 is similar in size to Xrs2p and also forms a complex with the human Rad50 and Mre11 proteins. Partly because of this, the possibility that NBS1 is functionally equivalent to Xrs2p has been considered. NBS1 shows weak homology to a putative *Neurospora* protein (Table 24).

Strand invasion. During meiosis, the E. coli RecA orthologues Dmc1p and Rad51p colocalize on chromosomes at foci that are the probable sites of recombination in S. cerevisiae. Both are thought to play a role in strand invasion, possibly via different pathways (596). While many organisms possess both Dmc1p and Rad51p orthologues, it appears that only Rad51p is represented in the Neurospora genome. Although budding yeast Dmc1p has homology to the hypothetical Neurospora protein NCU02741.1 (2.00E-83), NCU02741.1 is a closer match to S. cerevisiae Rad51p (Table 24). dmc1 and sae3 mutants have a similar phenotype and are thought to act in the same pathway (596). SAE3 appears to be unique to budding yeast. Because Tid1p/Rdh54p is generally needed for S. cerevisiae sporulation, affects spore viability, and interacts with Dmc1p in two-hybrid experiments, it is thought that it may be part of the Dmc1p pathway. It is possible that Neurospora also lacks an orthologue of this protein, since the two putative Neurospora proteins with Tid1p/Rdh54p homology are more similar to S. cerevisiae Rad54p and Rad26p, respectively (Table 24). Given the constraints imposed by RIP on the evolution and maintenance of gene families, it would not be surprising if Neurospora lacked this putative Dmc1p/Sae3p/Rdh54p pathway.

Localization of Dmc1p and Rad51p to chromosomal foci during meiosis appears to require Rad52p, Rad55p, and Rad57p, and *Neurospora* has an orthologue of each (Table 24). Rfalp forms a trimeric single-stranded DNA binding protein with Rfa2p/Rfa3p and is known to colocalize with Rad52p at chromosomal foci. While other organisms, including *Neurospora*, carry both *RFA1* and *RFA2* (Table 24), *RFA3* appears to be unique to *S. cerevisiae*.

Synapsis and SC formation. Synaptonemal complex (SC) formation in *S. cerevisiae* depends on recombination. It is thought that this is also the case for mammals but not for the worm or the fly. Zip1p and Zip2p are required for synaptonemal complex formation (148) but also modulate crossover frequency in *S. cerevisiae* (806).

Of the *S. cerevisiae* genes required for synapsis and SC formation, *Neurospora* has an orthologue only of *ZIP1* (Table 24). Although *S. cerevisiae* Zip1p is not highly similar to *Neurospora* ZIP-1, each matches human Zip1 to the same extent (Table 24), suggesting that this protein diverges between species. In support of this, human and mouse Zip proteins are only 65% identical. Of the other proteins required for synapsis in *S. cerevisiae* (Hop1p, Hop2p, Red1p, Mnd1p, and Mer1p) (186, 346, 469, 742, 855), all but Red1p and Hop1p have no close orthologues in any sequenced species. Red1p and Hop1p have close orthologues only in other yeasts, suggesting that these proteins may be species or lifestyle specific, with functions performed by different proteins in different groups of organisms.

Regulation of crossover frequency. *S. cerevisiae MSH4* and *MSH5* are *E. coli mutS* homologues with no role in mismatch repair (MMR). *S. cerevisiae MLH1* is an *E. coli mutL* homologue with MMR function. Mutation in any of these three genes decreases crossover frequency in *S. cerevisiae* (26, 570). The *S. cerevisiae MER3* gene, whose transcript is spliced by Mre2p/Mer1p, encodes a novel helicase required for transition of DSBs to intermediates (554). *mer3* mutants have fewer crossovers, and those that remain are randomly spaced along chromosomes (555), indicating a requirement for Mer3p in interference. A mutation in *TAM1/NDJ1* also abolishes interference in *S. cerevisiae* (167). Zip1p and Zip2p, as described above, are components of the SC, itself required for crossover regulation, and *S. cerevisiae mei5* mutants have normal levels of gene conversion but no crossovers (280).

It has been suggested (97) that a heterodimer of Msh4 and Msh5 proteins binds to Holliday junctions and that a heterodimer of Mlh1p and another MutL protein, possibly Mlh3p, subsequently binds to the Mshp-DNA complex. The complex of Mlh and Msh proteins influences resolution of the conversion intermediate and is required for crossover formation.

Neurospora has close orthologues of the *S. cerevisiae MSH4*, *MSH5*, *MLH1*, and *MLH3* genes. Of these, Mlh1p is highly conserved in all sequenced eukaryotes (Table 24). Msh4 and Msh5 are moderately conserved, with the *Neurospora* MSH-4 protein somewhat more closely related to the mouse than to the *S. cerevisiae* Msh4p, and Mlh3p much less so (Table 24). Although the *Neurospora* orthologues of *S. cerevisiae PMS1* and *MSH5* are unambiguously identifiable, it is difficult to determine which coding sequences are orthologous to *MSH4* or *MLH3* (Table 24). Therefore, relating *S. cerevisiae* phenotypes to the specific roles of these genes in crossing over in *Neurospora* is not likely to be useful. *Neurospora* has a close Mer3p orthologue (Table 24). Orthologous DNA helicases exist in other eukaryotes including humans, although both *Neurospora* and *S. cerevisiae* Mer3 proteins are closer matches to other eukaryotic RNA helicases than to DNA helicases (Table 24).

All other *S. cerevisiae* proteins required for crossing over and interference (Tam1p/Ndj1p, Rec102p, Rec104p, Rec107p/ Mer2p, and Mei5p/Lph6p) (77, 167, 270, 659) have no orthologues in any sequenced organism and thus may be species specific.

In addition, there are two genes in Drosophila, mei-9, encoding an orthologue of S. cerevisiae RAD1 (703), and mei-218 (515), each of which reduces crossing over by 95% but has no effect on gene conversion. Hence, these proteins may also play a role in resolution of conversion intermediates. Mei-218p appears to be Drosophila specific, since it has no orthologue in any sequenced species, but S. cerevisiae rad1 mutants have only 1-h delay in crossing over (166), clearly a different phenotype from mei-9. MUS-38 is the Neurospora Rad1p/Mei-9 orthologue, superficially a better match to Rad1p (Table 24) than to Mei-9 (Table 24). However, MUS-38 and Mei-9 proteins are more similar in length, each lacking both the amino and carboxy termini of Rad1p, and share more similarities in the carboxy termini of the proteins. It is possible, therefore, that MUS-38 has a function closer to that of Drosophila Mei-9 than of S. cerevisiae Rad1p.

Mismatch repair. Msh and Mlh proteins function in a highly conserved pathway for removing mismatches in DNA duplexes. Msh2p, as a heterodimer with Msh3p or Msh6p, binds to the mismatch. The protein-DNA complex attracts a heterodimer of Mlh proteins, Pms1p-Mlh1p, Pms1p-Mlh2p, or Mlh1p-Mlh2p. The identity of the Msh and Mlh proteins in the complex determines the type of mismatch recognized (97) and possibly influences the direction of correction (596). Mlh1 may be the only MMR protein responsible for restoration, if it ever occurs.

Mlh1p, Msh2p, Msh3p, and Msh6p are very highly conserved in eukaryotes, including *Neurospora*, while Pms1p is slightly less so (Table 24). In contrast, Mlh2p is much less highly conserved and cannot be unambiguously identified in *Neurospora* (Table 24). This suggests that although the pathway for correction of single-base mismatches, involving Msh2p-Msh6p and Mlh1p-Pms1p (97), is probably invariant across higher organisms, the way in which other types of mismatches are corrected may vary between species and must be determined by analysis of null mutations in each organism.

Interestingly, *Neurospora* MSH-3 is much more similar to the human than to the budding yeast orthologue (Table 24). If the Msh3p heterodimers interact with those including the less highly conserved Mlh2p, as suggested (97), MMR in *Neurospora* may have more similarities to MMR in mammals than in *S. cerevisiae*.

Resolution of recombination intermediates. Of the *S. cerevisiae* proteins thought to be involved in resolution of recombination intermediates (Slx1p, Mms4p/Slx2p, Slx3p/Mus81p, Slx4p, Hex3p/Slx5p, and Slx8p), *Neurospora* has orthologues of Slx1p and Slx3p/Mus81p (Table 24). In *S. cerevisiae*, Mus81p forms a complex with Mms4p to yield an endonuclease with roles in both meiotic recombination and in resolution of stalled

replication forks (395). Slx1p forms a heterodimer with Slx4p, while Slx5p forms a heterodimer with Slx8p (544).

Mus81p shows a moderate level of conservation in eukaryotes, but its partner in the complex, Mms4p, has no close orthologue either in Neurospora (Table 24) or in other organisms. Slx1p may be somewhat conserved, although only in fungi, but its partner, Slx4p, has no orthologue in sequenced species (Table 24). The closest match to Slx8p in Neurospora may be another DNA binding protein with a different function, since it is very similar to a S. pombe zinc finger protein and a closer match to a different yeast chromosomal region than that of the SLX8 sequence (Table 24). Once again, the partner, Slx5p, has no identifiable orthologue. In conclusion, because in each case only one member of each S. cerevisiae heterodimeric protein has an orthologue in Neurospora, the processes involved in this stage of recombination in other species cannot be predicted using knowledge gained from experiments with S. cerevisiae.

If recombination proceeds by a synthesis-dependent strandannealing mechanism (596), recombination intermediates may be resolved by a topoisomerase (282). All three *S. cerevisiae* topoisomerases (Top1p/Mak1p/Mak17p, Top2p/Tor3p/Trf3p, and Top3p/Edr1p) belong to a highly conserved protein family, and *Neurospora* is no exception (Table 24).

Nonhomologous end joining. NHEJ is a mechanism for the repair of DSBs without homologous recombination. Of the *S. cerevisiae* proteins required for this process (Lig4p, Yku80p/Hdf2p, Lif1p, and Yku70p/Hdf1p/Nes24p [326, 334, 855]), only Lif1p is not conserved in eukaryotes, and it is also the only protein lacking a *Neurospora* orthologue.

METABOLIC PROCESSES AND TRANSPORT

Extracellular Digestion

Neurospora is a saprophyte, obtaining nutrition from a substrate that is, for the most part, polymeric and insoluble. To assimilate this resource, polymers must first be broken into small, soluble units: monomers or small oligomers. Carbon nutrition is derived from polysaccharides, proteins, lipids, and nucleic acids; proteins and nucleic acids are nitrogen sources, sulfur is obtained from proteins, and phosphorus sources are predominantly nucleic acids. To exploit these resources, *Neurospora* possesses a number of genes for glycosyl hydrolases, proteases, lipases, nucleases, and phosphatases. In spite of the existence of RIP, the glycosyl hydrolases, proteases, and lipases are present in families; however, as with the sugar transporter family (269) (see "Transporters" below), there is no evidence of recent duplication. Therefore, it is probable that these families arose by gene duplication prior to the evolution of RIP.

Secreted cellulases can be detected by clearing of a cellulose halo and staining with Congo red (786), a starch halo is seen after iodine staining (760), proteases are detected by clearing of an opaque casein halo (10), and a nucleic acid halo is assayed by staining with α -naphthyl phosphate and Diazo blue B (393). Using such methods following mutagenesis, strains with altered halo sizes have been obtained, but many of these are regulatory mutants, affecting whole classes of secreted enzymes. For example, general carbon catabolite repression affects glycosyl hydrolases, lipases, proteases, and nucleases, the

Enzyme	EC no.	NCU no.	Cloned gene	Enzyme	EC no.	NCU no.	Cloned gene
Family 3 Xylan 1, 4-β-xylosidase	3.2.1.37	00709.1		Family 25 Lysozyme	3.2.1.17	00701.1	
Family 5 Endoglucanase	3.2.1.2	00762.1		Family 28 Endo-polygalacturonase Exo-polygalacturonase	3.2.1.15 3.2.1.67	02369.1 06961.1	
Family 6 Cellobiohydrolase	3.2.1.91	03996.1 07190.1 09680.1		Family 31 α-Glucosidase	3.2.1.20	02583.1 04674.1 09281.1	
Family 7 Cellobiohydrolase	3.2.1.91	04854.1 05104.1 05955.1		Family 32 Invertase Family 35	3.2.1.26	04265.1	inv
Endoglucanase	3.2.1.4	07340.1 05057.1 08227.1	cbh-1	β-Galactosidase	3.2.1.23	00642.1 04623.1	
Family 10 Endo-1,4-β-xylanase	3.2.1.8	04997.1 05924.1 07130.1 08189.1		Family 45 Endoglucanase Family 55 β-1,3-Exoglucanase	3.2.1.4 3.2.1.58	05121.1 04850.1 04947.1	
Family 11 Endo-1,4-β-xylanase	3.2.1.8	02855.1 07225.1				07523.1 08097.1 09791.1	
Family 13 α-Amylase	3.2.1.1	08131.1 09805.1		Family 61 Cellobiohydrolase Cellobiohydrolase Endoglucanase	3.2.1.91 3.2.1.91 3.2.1.4	00836.1 02240.1 02916 1	
Family 15 Glucoamylase	3.2.1.3	01517.1	gla-1	Cellobiohydrolase Endoglucanase Cellobiohydrolase	3.2.1.91 3.2.1.4 3.2.1.91	07760.1 08760.1 09764.1	
Family 16 Mixed linked glucanase 1,3-,1,4-β-glucanase	3.2.1 3.2.1.8/73	01353.1 08746.1		Family 81 Glucan 1,3-β-glucosidase	3.2.1.58	07076.1	
Family 18 Endochitinase	3.2.1.14	02814.1 04500.1					

TABLE 25. Glycosyl hydrolases

nitrogen metabolite repression genes *nit-2* and *nmr* regulate proteases and nucleases, and the transcriptional activator *nuc-1* regulates nucleases and phosphatases.

Glycosyl hydrolases. Halo methods have demonstrated the presence of extracellular cellulase and amylase activities in *Neurospora*. Four endoglucanases, three cellobiohydrolases, and one β -glucosidase were identified in spent medium (886); only one extracellular cellobiohydrolase had been cloned and sequenced prior to the completion of the genome sequence (775). An extracellular glucoamylase has also been cloned and sequenced (760), and an extracellular invertase has been identified (680). RIP inactivation of both the cellobiohydrolase and the glucoamylase left significant residual activity, suggesting the existence of isozymes of both. Mutants with quantitatively altered halos have demonstrated regulatory genes, e.g., *exo-1* (296) and *sor-4* (548).

Analysis of the genome predicted genes for approximately 100 glycosyl hydrolases. Those with the required function, the

necessary secretion signal sequence, and no other predicted destination have resulted in the list of candidate extracellular glycosyl hydrolases shown in Table 25. This group includes the previously sequenced *gla-1* glucoamylase, *cbh-1* cellobiohydrolase, and *inv* invertase. The enzymes are categorized according to the glycosyl hydrolase classification of Henrissat and Bairoch (332).

Proteases. Some mutational work has been done on extracellular proteases; the PTS-1 extracellular alkaline protease was identified based on electrophoretic variants in wild isolates (3, 313). Three inducible extracellular acid proteases have also been described (475). The genome sequence predicts a total of approximately 55 proteases. Analysis of these with regard to the required function and possession of the necessary signal sequence to permit secretion yields a total of 1 putative extracellular serine protease, 10 aspartyl proteases, and 1 metalloprotease (Table 26).

Nucleases and phosphatases. Several extracellular phosphatases have been characterized, including the PHO-3 repressible

TABLE 26. Proteases

Enzyme	NCU no.	EC no.
Serine protease		
Subtilase (lactocepin)	00263.1	3.4.21.96
Aspartyl protease		
Endothiapepsin	00994.1	3.4.23.22
1 1	02059.1	3.4.23.22
Aspergillopepsin	00338.1	3.4.23
Podosporapepsin	09484.1	3.4.23
1 1 1	07533.1	3.4.23
Candidapepsin	03168.1	3.4.23.24
1 1	09155.1	3.4.23.24
	07063.1	3.4.23.24
Pepsin?	02956.1	3.4.23
1	00249.1	3.4.23
Metalloprotease		
Thermolysin	05756.1	3.4.24

acid phosphatase and the PHO-2 repressible alkaline phosphatase. Two secreted phosphate-repressible alkaline DNases have been described (394), as well as a DNase A and a singlestrand-specific endonuclease (250). There are more than 30 nucleases and phosphatases in the *Neurospora* genome. These were further analyzed to identify those with the necessary function and signal sequence (Table 27). The group includes two non-specific S1 RNases, a DNase, a guanyl-specific N1 RNase, the PHO-2 alkaline phosphatase, and the PHO-3 acid phosphatase.

Lipases. A single extracellular triacylglycerol lipase associated with conidia has been described to date (441). Examples of extracellular lipases have been characterized in a number of other fungal species. Analysis of the genome and predicted genes for lipases reveals a total of 19, of which 7 triacylglycerol lipases have a secretion signal sequence, but 4 of these are probably membrane proteins (see also "Lipids" below). The

TABLE 27. Nucleases and phosphatases

Enzyme	NCU no.	EC no.	Cloned gene
Nonspecific endo-RNase S1	09194.1	3.1.30.1	
Nonspecific endo-RNase S1	08648.1	3.1.30.1	
DNase 1	01173.1	3.1	
Guanyl-specific RNase N1	01045.1	3.1.27.3	
Acid phosphatase	08643.1	3.1.3.3	pho-3
Alkaline phosphatase	01376.1	3.1.3.1	pho-2

remaining three candidates for extracellular lipases are listed in Table 28.

Transporters

Based on genome analysis, Neurospora encodes approximately 25% more transporter systems than does S. cerevisiae (605), primarily due to the presence of an increased number of major facilitator superfamily (MFS) and ATP binding cassette (ABC) family transporters (Fig. 5). A complete listing of predicted Neurospora membrane transporters categorized by gene family and function is available at http://www .membranetransport.org/. Neurospora appears to possess an enhanced capacity in terms of predicted drug efflux genes. It encodes more than twice as many ABC and MFS drug efflux systems as S. cerevisiae or S. pombe and also encodes two RND family efflux proteins, a type of efflux pump not seen in either S. cerevisiae or S. pombe. This expanded inventory of efflux systems relative to other fungi could conceivably play roles in (i) secretion of secondary metabolites, (ii) mediation of resistance to plant-produced secondary metabolites or other toxic compounds, or (iii) secretion of signaling molecules perhaps related to hypha morphogenesis. The first two possibilies correlate with the described repertoire of Neurospora secondary metabolite biosynthesis genes and cytochrome P450 detoxification genes and with the known abilities of plants and soil organisms to secrete a broad range of toxic compounds.



FIG. 5. Relative numbers of *Neurospora*, *S. cerevisiae*, and *S. pombe* transporters in various families. *Neurospora* was compared to *S. cerevisiae* (605) and *S. pombe* (http://www.membranetransport.org) with respect to transporters in the major facilitator superfamily (MFS), ATP binding cassette (ABC) superfamily, amino acid/polyamine/choline (APC) superfamily, mitochondrial carrier (MC) family, P-type ATPase family, and other families.

In terms of metabolite uptake, the capabilities of Neurospora appear to resemble those of S. cerevisiae, although Neurospora does encode an increased number of MFS uptake transporters of unknown specificity. Speculatively, these may be involved in the usage of plant-derived aliphatic and aromatic compounds. Neurospora has fewer predicted amino acid transporters than does S. pombe or S. cerevisiae. Neurospora has a similar number of predicted sugar MFS transporters as S. cerevisiae. However, phylogenetic analysis of fungal sugar transporters (269) indicates that the Neurospora sugar transporters are more divergent than those of S. pombe or S. cerevisiae. Consistent with the effects of RIP, there are no close paralogues of Neurospora sugar transporters, whereas the majority of the S. cerevisiae HXT hexose transporters and the S. pombe GHT transporters represent two relatively recent and independent expansions and include very recent duplicated genes such as HXT15 and HXT16. Hence, the Neurospora sugar transporters are much more divergent, and few, if any, are clear paralogues of characterized S. cerevisiae sugar transporters. This may indicate that the Neurospora predicted sugar transporters are more functionally diverse and able to utilize a much broader range of sugars, which would correlate with the expanded number of sugar polysaccharide hydrolases encoded in the Neurospora genome (605).

Glycolysis, Fermentation, and Gluconeogenesis

Glycolysis and the pentose phosphate cycle. Glycolysis is the process whereby sugars are metabolized to acetyl-CoA or pyruvate before oxidation during the citric acid cycle or fermentation to ethanol or lactate (reviewed in references 184, 241, and 300). Three glycolytic pathways have been described: the Embden-Meyerhof-Parnas (EM), hexose monophosphate (HM), and Entner-Doudoroff (ED) pathways (reviewed in reference 300). The EM and HM pathways are prevalent in fungi, while evidence for the ED pathway has been demonstrated for only a few species (Fig. 6) (300). The EM and HM pathways share glucose-6-phosphate, fructose-6-phosphate, and glyceraldehyde-3-phosphate. However, these two pathways play very different roles during metabolism (184, 241, 300). The EM pathway utilizes NAD as an electron acceptor to produce NADH, which can then be reoxidized by fermentation or oxidative phosphorylation. In contrast, HM glycolysis reduces NADP to provide NADPH for a host of biosynthetic reactions. EM glycolysis produces ATP and pyruvate that can be converted to acetyl-CoA for entry into several key pathways, including the citric acid cycle and fatty acid biosynthesis. The latter portion of the HM pathway, termed the pentose phosphate cycle, consists of a series of freely reversible sugar-phosphate interconversions that provide glyceraldehyde-3-phosphate and fructose-6-phosphate to enter the EM pathway, as well as precursors of sugar alcohols, nucleic acids, and aromatic amino acids (184, 241, 300).

Consistent with the action of RIP, *Neurospora* usually possesses one copy of each gene in a given pathway, in contrast to the two (or more) often found in the *S. cerevisiae* genome (Table 29; Fig. 6). In addition, with the exception of a single transaldolase (see below), the predicted *Neurospora* proteins had other fungal enzymes as their best match. Of these, the highest similarity was to corresponding proteins from other

TABLE 28. Lipases

Enzyme	NCU no.	EC no.
Triacylglycerol lipase	03639.1	3.1.1.3
Triacylglycerol lipase	03301.1	3.1.1.3
Triacylglycerol lipase	08752.1	3.1.1.3

filamentous fungi; in only one case (an alcohol dehydrogenase) was the closest homologue another *Neurospora* protein. Finally, only two enzymes in the pathways analyzed, phosphoglucomutase and glucose-6-phosphate isomerase, appear to be represented by known mutations in *Neurospora* (see below).

(i) Hexose phosphorylation. The glycolytic process begins with the phosphorylation of hexoses. Glycogen phosphorylation and cleavage to form glucose-1-phosphate is accomplished by glycogen (starch) phosphorylase. Similar to yeasts (241), Neurospora contains one form of this enzyme (Table 29). Glucose and fructose monomers can be phosphorylated by hexokinase at the C-6 position, while glucokinase phosphorylates glucose but not fructose. S. cerevisiae contains two hexokinases and one glucokinase (241). Evidence had previously been presented for four hexokinase activities in Neurospora (reviewed in reference 184). Analysis of the genome sequence is superficially consistent with these observations, since Neurospora has one good match each to glucokinase and hexokinase and two other genes with similarity to hexokinases. However, the protein corresponding to one of these other genes (NCU06996.1) does not appear to function as a hexokinase in Aspergillus nidulans but, instead, somehow regulates extracellular proteases during carbon starvation (404).

(ii) EM glycolysis. Previous work had suggested that the predominant route for glycolysis in *Neurospora* is entry of glucose-6-phosphate into the EM pathway (80 to 90%), with the remainder diverted to the HM pathway (reviewed in reference 184). The analysis of the genome sequence is consistent with intact EM and HM glycolytic pathways in *Neurospora* (Table 29).

To begin the EM pathway, the glucose-1-phosphate produced by the action of glycogen phosphorylase on endogenous glycogen is converted to glucose-6-phosphate by phosphoglucomutase; mutation of the *Neurospora rg* gene (533) causes deficiencies in this enzyme. Similar to *S. cerevisiae*, *Neurospora* contains one phosphoglucomutase gene. The presumed allelic *gpi-1* and *gpi-2* mutants (547) are lacking in glucose-6-phosphate isomerase, the enzyme that converts glucose-6-phosphate into fructose-6-phosphate. Analysis of the genome sequence showed that *Neurospora*, like yeasts (241), contains one glucose-6-phosphate isomerase gene.

Phosphofructokinase is a key regulatory enzyme in glycolysis, catalyzing the nonreversible phosphorylation of fructose-6-phosphate to produce fructose-1,6-bisphosphate, consuming a molecule of ATP in the process. *Neurospora* possesses one phosphofructokinase gene, in contrast to the two found in *S. cerevisiae* (241). *Neurospora* and yeasts both contain one fructose-bisphosphate aldolase gene (241), encoding the next enzyme in the pathway, which cleaves fructose-1,6-bisphosphate to form one molecule each of glyceraldehyde-3-phosphate and dihydroxyacetone phosphate.

Triose-phosphate isomerase interconverts glyceraldehyde-3phosphate and dihydroxyacetone phosphate. Analysis of the



FIG. 6. Glycolysis and alcoholic fermentation. *Neurospora* contains genes encoding enzymes of the EM (blue arrows) and HM (red arrows) pathways of glycolysis. Enzymes required for hexose phosphorylation are indicated above the black arrows, while those involved in the fermentation pathway are indicated by green arrows. Abbreviations for enzyme names are presented in Table 29. NCU numbers for predicted *Neurospora* proteins corresponding to each enzyme are indicated alongside each arrow.

genome sequence shows that *Neurospora* possesses one good match to this gene. The protein encoded by another gene (NCU04399.1) shows some similarity to triose-phosphate isomerases in the carboxy terminus but is most similar to a putative ribose 5-phosphate isomerase from *Streptomyces coelicolor*. The weaker similarity, coupled with a large intron in the amino terminal region, lessens the likelihood that this gene is a true triose-phosphate isomerase. In addition, *S. cerevisiae* possesses only one triose-phosphate isomerase gene (241).

Glyceraldehyde-3-phosphate dehydrogenase converts glyceraldehyde-3-phosphate into 1,3-diphosphoglycerate, with accompanying reduction of NAD to NADH and H⁺. For glycolysis to continue, the reduced NADH must be reoxidized (241). *Neurospora* contains one glyceraldehyde-3-phosphate dehydrogenase gene, in contrast to the three found in *S. cerevisiae* (241). It has been shown that glyceraldehyde-3-phosphate dehydrogenase is allelic to *ccg*-7, a clock-controlled gene in *Neurospora* (63, 731).

			•			•	
	EC	NCU			BLAST match		
	no.	no.	Best overall	S. cerevisiae	S. pombe	Animal ^a	Plant^{b}
Hexose phosphorylation Glycogen phosphorylase (GP)	2.4.1.1	07027.1	A. fumigatus GP; 0.00	Gph1p; 0.00	None	Liver GP; 0.00	Os α -1,4-glucan phosphor-
Hexokinase (HK) Hexokinase-like Hexokinase-like	2.7.1.1	02542.1 06996.1 04728.1	 A. oyzae HxkA; 1.00e-167 A. nidulans XprF; 1.00e-100 K. lactis RAG5 hexokinase; 	Hxk2p; 2.00e-117 Glk1p; 1.00e-17 Hxk2p; 2.00e-31	HK 1; 3.00e-112 HK 1; 6.00e-20 HK 1; 7.00e-32	Dm HK-t2; 1.00e-63 Hs MGC:29452; 5.00e-20 Dm HK-t2; 5.00e-26	ytase; 0.00 Os HK II; 8.00e-54 Os putative HK 1; 4.00e-22 At HK; 2.00e-25
Glucokinase (GK)	2.7.1.2	00575.1	4.00e-34 A. niger GK; 1.00e-155	Glk1p; 3.00e-77	HK 2; 1.00e-76	Dm HK-t2; 1.00e-51	Os putative HK 1; 3.00e-60
EM glycolysis Phosphoglucomutase (PGM) Glucose-6-phosphate isomerase (GPI) ⁶ 6-Phosphofructokinase (PFK) Fructose-bisphosphate aldolase (FBA) ⁶	5.4.2.2 5.3.1.9 2.7.1.11 4.1.2.13	10058.1 07281.1 00629.1 07807.1	 A. oyzae PgmA; 0.00 A. oyzae PgiA; 0.00 A. niger PFK; 0.00 P. brasiliensis AAL34519.2; 	Pgm2p/Gal5 p; 0.00 Pgi1p; 0.00 Pfk2p; 0.00 Fbalp; 1.00e-128	NP_59615 3.1; 0.00 NP_59663 5.1; 0.00 NP_59594 6.1; 0.00 FBA_NP_59569 2.1;	Hs PGM1; 1.00e-158 Dm AAA6367 2.1; 1.00e-168 Hs 6-PFK; 0.00 Ag EAA0269 2.1; 5.00E-88	At PGM; 1.006-155 At BAB1763 7.1; 1.006-111 At BAB5549 9.1; 1.006-10 AT T10022.2 4; 2.006-08
Triose-phosphate isomerase (TPI) ^c Glyceraldehyde-3-phosphate dehydrogenase	5.3.1.1 1.2.1.12	$07550.1 \\ 01528.1$	P. brasiliensis TPI; 2.00e-83 S. macrospora Gpd; 1.00e-178	Tpi1p; 5.00e-79 Tdh1p; 1.00e-119	1.00e-118 TPI; 7.00e-72 GAPDH; 1.00e-127	(not in mammais) Hs TPI 1; 2.00e-77 Mm NP_03211 0.1; 2.00e-135	Os TPI; 3.00e-76 At NP_17807 1.1; 1.00e-136
OATDIT) Phosphogycerte kinase (PGK) ^c Enolase (ENO) ^f Enolase ^c Pyruvate kinase (PYK)	2.7.2.3 4.2.1.11 4.2.1.11 2.7.1.40	07914.1 10042.1 01870.1 06075.1	T. viride PGK; 0.00 A. oyzae ENO; 0.00 S. japonicum ENO; 8.00e-129 T. reesei PYK; 0.00	Pgklp; 1.00e-154 Enolp; 1.00e-173 Enolp; 1.00e-120 Pyklp; 4.00e-173	NP_59673 0.1; 8.00e-163 NP_59590 3.1; 1.00e-155 NP_59590 3.1; 1.00e-120 PYK NP_59434 6.1; 9.00e-179	Hs PGK1; 1,00e-160 Dm NP 72272 1.1; 2.00e-149 Mm ENO 1; 1.00e-123 Ag EAA1055 5.2; 2.00e-140	At NP_17601 5.1; 1.00e-105 At ENO F2P9.10; 1.00e-140 At ENO F2P9.10; 1.00e-122 At PYK NP_19436 9.1; 4.00e-107
HM glycolysis/pentose phosphate pathway Glucose-6-phosphate 1-dehydrogenase //DDD41	1.1.1.49	09111.1	A. niger GPDH; 0.00	Zwf1p; 1.00E-160	Zwf1; 1.00E-147	GPDH X-linked; 1.00e-138	At GPDH; 1.00e-132
(JCDD) 6-Phosphogluconol aconase (PGL) 6-Phosphogluconic dehydrogenase (PGD) 6-Phosphogluconic dehydrogenase (PGD) Ribulose-Phosphate 3-epimerase (RPE) Transketolase (TK) Transaldolase (TA) Transaldolase?	3.1.1.31 1.1.1.44 1.1.1.44 5.1.3.1 2.2.1.1 2.2.1.2 2.2.1.2	00087.1 03100.1 00837.1 00519.1 01328.1 02136.1 06142.1	 S. pombe Sol1; 8.00e-65 A. niger PGD; 0.00 A. niger PGD; 1.00e-120 S. cerevisiae Repelp; 8.00e-51 A. niger TK; 0.00 K. thyrevi Tallp; 1.00e-106 H. sapiens transaldolase; 1.00e-15 	Sollp; 4.00e-57 Gndlp; 0.00 Gnd2p; 2.00e-66 Top hit Tk11p; 0.00 Tallp; 5.00e-10 Tallp; 5.00e-10	Top hit PGD; 0.00 SPBC660 16; 5.00e-67 SPAC31G 5.05c; 5.00e-48 SPBC2G5.05; 0.00 Tal1; 1.00e-106 Tal1; 2.00e-09	Mm PGL; 4.00e-42 Hs PGD protein; 0.00 Mm PGD; 1.00e-62 Hs MGC:226 37; 9.00e-44 Hs TK; 2.00e-42 Ag XP 32071 5.1; 2.00e-88 Top hit	At PGL-like; 4,00e-39 At PGD; 1,00e-115 At PGD-related; 9,00e-72 RPE; 9,00e-48 At TK precursor related; 0,00 At TOTAL2; 7,00e-32 None
Fermentation Pyruvate decarboxylase (PDC) Pyruvate decarboxylase (PDC)	4.1.1.1 4.1.1.1	02193.1 02397.1	S. pombe SPAC186.09; 0.00 P. anserina CAD60727.1; 0.00	Pdc6p; 6.00e-57 Pdc1p; 1.00e-141	Top hit C13A11.06; 1.00e-163	Dm CG11208-PA; 2.00E-11 Ce 2-hydroxyphytanoyl-CoA	At Pdc1; 1.00e-119 At PDC-related protein;
Alcohol dehydrogenase (ADH) Alcohol dehydrogenase (ADH)	1.1.1.1 1.1.1.1	01754.1 02476.1	A. flavus Adhl; 1.00e-133 N. crassa NCU01754.1; 1.00e-126	Adh3p; 1.00e-105 Adh3p; 6.00e-97	Adh1; 9.00e-97 Adh1; 1.00e-87	iyaes, 1.000-11 Ce probable ADH; 1.006-73 Ce ADH; 2.006-70	2.002-03 Os CAD3990 7.1; 3.00e-26 At cinnamyl-ADH; 8.00e-30
Gluconeogenesis Fructose-bisphosphatase (FBP) Phosphoenolpyruvate carboxykinase (PEPCK)	3.1.3.11 4.1.1.49	04797.1 09873.1	 A. nidulans FBP; 5,00e-163 A. nidulans PEPCK; 0.00 	Fbp1p; 1.00e-112 Pck1p; 0.00	P09202; 1.00e-102 None	Dm CG31692-PA; 1.00e-83 None	At 1g43670; 6.00e-93 At 4g37870; 0.00

TABLE 29. Glycolysis, the pentose phosphate cycle, alcoholic fermentation, and gluconeogenesis

BORKOVICH ET AL.

42

^a Caenorhabditis elegans (Ce), Drosophila melanogaster (Dm), Anopheles gambiae (Ag), Mus musculus (Mm), or Homo sapiens (Hs). ^b Arabidopsis thaliana (At) or Oryza sativa (Os). ^c Also a component of gluconeogenic pathway.

A high-energy phosphate from 1,3-diphosphoglycerate is transferred to ADP to produce ATP and 3-phosphoglycerate by the enzyme phosphoglycerate kinase. This is the first of two ATP-generating steps in glycolysis (241). Like *S. cerevisiae*, the *Neurospora* genome sequence contains one phosphoglycerate kinase gene. *S. cerevisiae* PGK1 is abundantly expressed; there are also numerous ESTs for the corresponding *Neurospora* gene.

There were no good matches in the genome database to phosphoglycerate mutase, the enzyme that allows the interconversion between 3-phosphoglycerate and 2-phosphoglycerate. The best match to both the rat and fungal enzymes, NCU01921.1, is interrupted by several introns and is most similar to an uncharacterized ORF from *S. cerevisiae*. Because phosphoglycerate mutase is a highly conserved enzyme, it is likely that the true phosphoglycerate mutase gene is absent from the current sequence assembly. *S. cerevisiae* contains one phosphoglycerate mutase gene (241).

Enolase catalyzes the production of phosphoenolpyruvate from 2-phosphoglycerate. *Neurospora* and *S. cerevisiae* (241) each contain two enolase genes. The two *S. cerevisiae* genes are differentially regulated (241); in this regard, it may be significant that NCU10042.1 is encoded by several ESTs while NCU01870.1 is represented by only one EST in available *Neurospora* databases.

The final step of EM glycolysis is the conversion of phosphoenolpyruvate and ADP to pyruvate and ATP by pyruvate kinase. This is the both the second ATP-generating and irreversible step of the EM pathway. *Neurospora* possesses one pyruvate kinase gene, in contrast to the two found in *S. cerevisiae* (241).

(iii) HM and ED glycolysis and the pentose phosphate cycle. The HM and ED pathways begin with glucose-6-phosphate being converted to glucono-1,5-lactone-6-phosphate by glucose-6-phosphate dehydrogenase. Similar to *S. cerevisiae* (241), *Neurospora* contains a single version of this enzyme (Table 29). The second step of the HM and ED pathways, production of 6-phosphogluconate from glucono-1,5-lactone-6-phosphate, is catalyzed by 6-phosphogluconolaconase. *S. cerevisiae* possesses genes encoding four such proteins, while *Neurospora* contains only one.

At this point, the HM and ED pathways diverge. The next reaction in the ED pathway is the conversion of 6-phosphogluconate to 2-dehydro-3-deoxygluconate-6-phosphate by phosphogluconate dehydratase. This enzyme is not found in S. cerevisiae (http://www.yeastgenome.org/). The best Neurospora match to the E. coli phosphogluconate dehydratase is an enzyme of amino acid biosynthesis, a dihydroxy acid dehydratase (Ilv3p; http://www.yeastgenome.org/) found in both S. cerevisiae and S. pombe. Similarly, the final enzyme in ED glycolysis, 2-dehydro-3-deoxyphosphogluconate aldolase, is not present in the *Neurospora*, *S. cerevisiae*, or *S. pombe* genome sequences (http: //www.yeastgenome.org/; http://www.sanger.ac.uk/Projects /S pombe/). This enzyme catalyzes the production of glyceraldehyde-3-phosphate and pyruvate from 2-dehydro-3-deoxygluconate-6-phosphate. Thus, Neurospora, like the two sequenced yeasts, lacks the final two (and the only pathway-specific) enzymes of ED glycolysis.

Continuing along the HM glycolytic route is the enzyme 6-phosphogluconate dehydrogenase, which catalyzes the production of ribulose-5-phosphate and carbon dioxide from 6-phosphogluconate. Like *S. cerevisiae* and *S. pombe, Neurospora* possesses two genes encoding this enzyme. Similarly, *S. cerevisiae* (241) and *Neurospora* each contain one gene encoding ribulose-phosphate 3-epimerase, the enzyme that converts ribulose-5-phosphate to xylulose-5-phosphate.

The next two enzymes in the HM pathway, transketolase and transaldolase, comprise the pentose phosphate cycle. Transketolase interconverts ribose-5-phosphate and xylulose-5-phosphate to produce sedoheptulose-7-phosphate and glyceralde-hyde-3-phosphate. In some species, this enzyme can utilize erythrose-4-phosphate and xylulose-5-phosphate to produce fructose-6-phosphate and glyceraldehyde-3-phosphate. *Neurospora* has one transketolase gene, in contrast to the two found in *S. cerevisiae* (241).

Transaldolase reacts sedoheptulose-7-phosphate and glyceraldehyde-3-phosphate to produce erythrose-4-phosphate and fructose-6-phosphate. *S. cerevisiae* and *S. pombe* contain one transaldolase gene (241). *Neurospora* has two matches to transaldolases; one is most similar to fungal isozymes (NCU02136.1), while the other (NCU06142.1) shows highest identity to transaldolase from humans but still exhibits significant similarity to transaldolases from fungi. Further analysis is needed to determine whether this second protein possesses transaldolase activity and contributes to pentose phosphate metabolism in *Neurospora*.

Alcoholic fermentation. Pyruvate produced from the glycolytic pathway can be oxidized to CO_2 through the action of the citric acid cycle, or it can be converted to ethanol by the alcoholic fermentation pathway (184, 241). Fermentation in fungi begins with the decarboxylation of pyruvate to yield acetaldehyde and CO_2 by the enzyme pyruvate decarboxylase (184). The *Neurospora* genome contains one pyruvate decarboxylase gene, while *S. cerevisiae* possesses at least three such genes (241).

To complete fermentation, acetaldehyde is reduced to ethanol using NADH + H⁺ by alcohol dehydrogenase (184). *Neurospora* possesses two proteins with high similarity to *S. cerevisiae* alcohol dehydrogenases (yeast has a total of four [241]), as well as several other predicted proteins with lower BLASTp scores (Table 29; Fig. 6). One of the two *Neurospora* alcohol dehydrogenases has the other *Neurospora* protein as its closest homologue.

Gluconeogenesis. Gluconeogenesis allows organisms to utilize noncarbohydrates as energy sources (Fig. 6). The gluconeogenic pathway is essentially the reversal of EM glycolysis and shares several enzymes that catalyze freely reversible reactions, including enolase, phosphoglycerate kinase, glyceraldehyde-3-phosphate dehydrogenase, triose-phosphate isomerase, fructose-bisphosphate aldolase, and glucose-6-phosphate isomerase. However, two of the reactions catalyzed by EM glycolytic enzymes, pyruvate kinase and 6-phosphofructokinase, are not reversible due to unfavorable levels of reactants in the cytoplasm (241). Thus, organisms must utilize phosphoenolpyruvate carboxykinase which catalyzes the conversion of oxaloacetate to phosphoenolpyruvate with net consumption of ATP) and fructose bisphosphatase (which hydrolyzes fructose-1,6-diphosphate to form fructose-6-phosphate) to circumvent the two nonreversible steps. Neurospora and S. cerevisiae

(241) each contain one copy of the phosphoenolpyruvate carboxykinase and fructose bisphosphatase genes (Table 29).

Mitochondrion and Energy Metabolism

The mitochondrial genome of Neurospora is a circle of 64,840 bp. It contains two rRNAs, at least 27 tRNAs, and 26 ORFs. Three tRNAs (Met-1, Met-2, and Cys) have been duplicated. Thus, including tRNA Met-3, a total of five methionine tRNAs are present. The duplicated tRNAs Met-2 and Cys are part of a 3-kb duplication and are located upstream and downstream, respectively, of two duplicated concatenated fragments of the nd2 gene. The entire duplicated region gives rise to one polycistronic mRNA (9). As in most eukaryotes, 14 subunits of complexes involved in oxidative phosphorylation are encoded in the mitochondrial DNA. A gene for ATPase subunit 9 is present in the nucleus as well as in the mitochondrial genome (823). The nuclear gene accounts for the bulk of the subunit in vegetative cells (373, 699, 814), but the mitochondrial version seems to be expressed in germinating spores (83). As mentioned above, part of the nd2 gene is duplicated, resulting in an additional ORF of unknown significance.

A total of 10 intronic ORFs are present in the single introns of the atp6, nd1, nd3, nd4, and nd4L genes, the two introns in each of nd5 and cob, and the intron in the rnl gene that encodes the large subunit rRNA. The latter intronic ORF codes for ribosomal protein S5, a homologue of the S. cerevisiae Var1p protein (115). All introns in the protein-coding genes are group I introns that encode putative maturases/endonucleases (136). The ORFs in the introns of the atp6 and nd1 genes are free-standing, while the other intron ORFs are continuous, in frame with the preceding exon. The ORF in intron 1 of the cob gene shows an exci-endo-N domain encoding a putative GIY-YIG motif endonuclease. All other introns encode putative endonucleases of the LAGLIDADG (dodecapeptide) type (448). However, the ORF in the nd3 gene is truncated by a stop codon interrupting what may have been the coding sequence for a maturase/endonuclease.

The only ORF in the *Neurospora* mitochondrial genome lacking a homologue in other organisms encodes a hypothetical 70-kDa protein. The ORF is located upstream of the *cox1* gene and is cotranscribed with the latter (117). No experimental data are available regarding its expression. Reviews of studies of the content and expression of the *Neurospora* mtDNA have been published elsewhere (301, 410).

Mitochondrial proteins encoded in the nucleus have to be imported into mitochondria. The majority of mitochondrial proteins contain N-terminal presequences that serve as targeting signals that are cleaved after import. Others lack a cleavable N-terminal import signal, and their targeting information is part of the mature protein sequence (567, 646, 838). Sequence features conserved in N-terminal targeting sequences have been used to predict the mitochondrial localization of proteins deduced from DNA sequence. However, the predictions are hampered by poor conservation of these targeting signals. Target P, a widely used software tool for the sequence based identification of N-terminal presequences (220), predicts that 18%, or 1,962, of the 10,900 deduced *Neurospora* ORFs would be mitochondrial. To judge the usefulness of this prediction for compiling a set of mitochondrial ORFs in *Neuro*-

TABLE 30. *Neurospora* ORFs assigned to mitochondria by different means

ORF	Total no.	No. (%) assigned to mitochondria by Target P
Manually annotated	8,350	1,515 (18)
Manually annotated assigned to any cell compartment	1,096	269 (25)
Manually annotated assigned to mitochondria	301	174 (58)

spora, the performance of Target P was tested using 1,096 ORFs assigned to various cellular compartments based on their close relationship to known proteins from other organisms (Table 30) (498). Of 269 ORFs predicted by Target P to be mitochondrial, 95 (36%) had been assigned to a nonmitochondrial compartment. Only 58% of the 301 ORFs assigned to mitochondria were identified by Target P (Table 30). ORFs not identified by Target P to be mitochondrial include those lacking an N-terminal targeting signal, those containing an N-terminal targeting signal not recognized by Target P, and those incorrectly suggested to contain an ER targeting signal by the program. When the prediction of Target P is adjusted by considering these two counteracting sources of error, the number of different mitochondrial proteins predicted for Neurospora reaches about 2,200. We are not aware of any bias in the set of 1,096 ORFs used for the comparison that would result in a significant error when extrapolated to the entire gene set. Estimates for the number of mitochondrial proteins in other organisms cover a wide range. For example, for S. cerevisiae, an estimate based largely on the results of a high-throughput immunolocalization screen suggested that 13%, or 793, of the 6,100 predicted yeast proteins would be mitochondrial (439). Based on protein phylogenetic profiles, Marcotte et al. (499) calculated about 630 mitochondrial proteins for yeast and 660 for C. elegans. Larger numbers were calculated for A. thaliana (2,000) (437) and humans (1,500) (784), based on data obtained from two-dimensional (2D) gel electrophoresis.

An alternative approach to the identification of mitochondrial proteins deduced from the genomic sequence of Neurospora relies on sequence comparison. This works quite well for generally known mitochondrial proteins, since the classification can be drawn from the comparison to homologs from various organisms. Thus, by manual annotation of 8,350 of the 10,900 Neurospora ORFs predicted in the genome sequence, 301 Neurospora ORFs were assigned to the mitochondrion due to their similarity to known mitochondrial proteins (Table 30). Significantly less reliable are conclusions drawn from comparisons to mitochondrial proteins from single, distantly related organisms. This is illustrated by a comparison to a large set of mitochondrial proteins compiled from data obtained by 2D gel electrophoresis of human mitochondria (785). Of 615 proteins in this set, 178 proteins did not match ($E > 10^{-5}$) a *Neurospora* ORF. Among these were ribosomal proteins and subunits of complexes involved in protein import and oxidative phosphorylation which probably did not yield a Neurospora homologue due to weak sequence conservation. Furthermore, detection of high similarity to human proteins identified by the 2D gel approach can also be misleading. Among 71 known Neurospora

proteins that are closely related to human proteins in the data set, 22 have been shown experimentally to exist outside mitochondria. This could be due to contamination of the human mitochondria that were analyzed or it could be because some proteins that show a relationship to mitochondrial proteins are actually localized to a different cellular compartment.

More reliable conclusions can be drawn from comparisons to closely related organisms. In the S. cerevisiae database, a set composed of 412 protein sequences is annotated as mitochondrial (MIPS database: http://mips.gsf.de/proj/yeast /CYGD/db; SGD database: http://genome-www.stanford.edu /Saccharomyces). By searching the entire Neurospora gene set deduced from the genomic sequence for close homologues, we identified 350 Neurospora ORFs. When these were combined with the 301 ORFs attributed to mitochondria by manual annotation, a set of 446 Neurospora ORFs resulted. For 321 of the predicted Neurospora mitochondrial protein ORFs, a 1:1 match to an ORF of S. cerevisiae was found. A total of 36 Neurospora ORFs were found in a 2:1 ratio relative to S. cerevisiae. That is, 18 pairs of Neurospora ORFs were found where both members of the pair were most closely related to a single S. cerevisiae ORF. In one case, a set of three Neurospora ORFs was most closely related to a single S. cerevisiae ORF (3:1 ratio). Conversely, 17 Neurospora ORFs were found in a 1:2 ratio relative to yeast ORFs. That is, 17 pairs of S. cerevisiae ORFs were found where both members of the pair most closely matched one Neurospora ORF. Four Neurospora ORFs each matched three different ORFs of S. cerevisiae (1:3 ratio). There were 65 Neurospora mitochondrion-assigned ORFs and 46 S. cerevisiae mitochondrion-assigned ORFs that were not detected in the deduced sequences of the other organism.

ORFs identified as mitochondrial in S. cerevisiae but lacking a significant Neurospora match are involved mainly in the expression of mitochondrial genes and the processing of gene products. A considerable number are likely to have a Neurospora homologue that was not detected in this analysis due to insufficient sequence conservation. For example, there are nine subunits of S. cerevisiae mitochondrial ribosomes that lack a matching ORF in the Neurospora genome. Similarly, 22 proteins involved in the processing and translation of mitochondrial mRNAs and the assembly of respiratory complexes, four subunits of respiratory complexes, and two subunits of the translocation channel of the outer membrane are known in yeast, but a Neurospora homologue has not yet been identified. Three S. cerevisiae proteins, Imp1p, Imp2p, and Som1p, are involved in the processing of proteins in the inner membrane. The other nonmatched S. cerevisiae proteins include a protoporphyrinogen oxidase (Hem14p), a protein required for autophagy (Apg14p), RNase P (Rpm2p), a 3'-5' exonuclease (Rex2p), and Lag2p, which is involved in determining longevity.

Most of the *Neurospora* mitochondrial proteins not found in *S. cerevisiae* are involved in respiration. *S. cerevisiae* and other fermentative yeasts lack respiratory complex I, and so the 31 subunits of complex I (24 nuclear and 7 mitochondrially encoded) are not present. In addition, five intron-located ORFs in *nd* genes of the mitochondrial genome and two assembly factors for complex I identified in *Neurospora* (438) are absent from yeast. Also missing in *S. cerevisiae* is the alternative oxidase that catalyzes the oxidation of ubiquinol by oxygen, thus

bypassing respiratory complexes III and IV. Alternative oxidase is present in the mitochondria of all higher plants, many fungi, many eukaryotic algae, and some protists (828). In Neurospora, the enzyme is present only under conditions that compromise normal mitochondrial function (450, 451). The aod-1 gene (4nc285_080 [Table 31]) codes for the Neurospora alternative oxidase (472). Mutant screens also identified the aod-2 gene, which is thought to encode an as-yet-unidentified regulatory factor required for *aod-1* expression (72, 449). A search of the Neurospora genome revealed a close relative of aod-1. This gene, termed aod-3 (xnc010 210 [Table 31]) specifies a second alternative oxidase, but the conditions required for its expression are unknown (782). Finally, yeast mitochondria lack a transhydrogenase for coupling the redox state of the internal and external NAD⁺-NADH pool, which is found in Neurospora and many other organisms (13e11_040 [Table 31] (446).

A number of ORFs assigned to mitochondria in Neurospora but not found in S. cerevisiae are involved in the degradation of amino acids. A probable isovaleryl-CoA dehydrogenase and a probable β -subunit of methylcrotonyl-CoA carboxylase are found adjacent to each other on linkage group I (1nc356 060 and 1nc356 070) in a head-to-head orientation, while the α subunit of methylcrotonyl-CoA carboxylase (b22i21 180) is located on linkage group II. All three participate in the degradation of leucine. In addition, ORF 29e8 120 is related to 3-hydoxybutyrate dehydrogenase, involved in the degradation of valine. The amino acids valine, leucine, and isoleucine share related degradation pathways. Enzymes involved in the degradation are found in mitochondria as well as the cytosol. Neurospora uses the same enzymes as plants and animals for this purpose, while S. cerevisiae and S. pombe lack these enzymes.

Other ORFs found in *Neurospora* but not in *S. cerevisiae* are rather distantly related to known proteins, and their function is therefore still unresolved (Table 31). These include another two acyl-CoA dehydrogenases. Also listed are seven mitochondrial carrier proteins of unknown specificity for which a close yeast homologue has not been identified. This is in contrast to the majority of mitochondrial carrier proteins, for which a 1:1 ratio to a yeast homologue is apparent. The closest *S. cerevisiae* homologues listed in Table 31 are significantly more closely related to a different *Neurospora* protein.

A peculiarity of filamentous fungi appears to be the presence of a specific succinyl-CoA synthetase (SCS). The known SCS is a heterodimer of two subunits (105). The genome of *Neurospora* reveals two ORFs for these subunits (bj10_140 and 8d4_130 [Table 31]), as well as a third ORF (1nc250_090 [Table 31]) in which the N-terminal half is related to the beta subunit, while the C-terminus is homologous to the alpha subunit. This additional SCS is also found in *A. nidulans* and *M. grisea* but has not yet been found in yeast, plants, and animals.

As yet, there is no comprehensive set of mitochondrial proteins compiled for *Neurospora* or any other organism. Since fewer than 500 known or deduced proteins have been assigned to mitochondria, this suggests that only one-quarter of the expected 2,000 different mitochondrial proteins can be extracted using the genomic sequence information alone. Although the proteins required for well-characterized mitochon-

	NCU	MIPS			BLAST match		
Gene name	no.	code	Best overall	S. cerevisiae	S. pombe	Animal	Plant
<i>aod-3</i> Alternative oxidase (<i>aod-1</i>) Probable isovaleryl-CoA dehydrog-	04874.1 07953.1 02126.1	xnc010_210 4nc285_080 1nc356_060	2.0e-140, T:AF321004_1 1.6e-251, TN:AY140655_1 1.0e-145, PIR:D95929	None None None	None None None	None None 5.7e-130, PIR:A37033	1e-53, PIR:T07947 1e-50, PIR:T07947 9.0e-143, SP:IVD2_SOLTU
enase Probable methylcrotonyl-CoA	02127.1	1nc356_070	3.2e-234, TN:AE003790_19	None	None	2.0e-221, SN:MCCB_HUMAN	1e-159, T:AF386926_1
carboxylase beta chain Probable methylcrotonyl-CoA	00591.1	b22i21_180	2.5e-165, TN:AE003779_43	None	None	1.3e-122, SN:MCCA_HUMAN	7.4e-161, SN:MCCA_ARAT H
carboxylase alpha chain Probable dienoyl-CoA isomerase	06647.1	$18a7_070$	4e-62, PIR:T16494	None	None	4e-62, PIR:T16494	2e-49, T:AB017070_10
Related to 3-hydroxyisobutyrate	ND^b	29e8_120	6.6e-52, PIR:G96013	None	None	7.4e-22, T:AC007130_1	8.1e-52, PIR:D86317
Related to long-chain-specific acyl-	08924.1	20h10_010	1.1e-109, T:AY033936_1	None	None	6.0e-26, TN:AB083302_1	None
CoA denydrogenase Related to short-chain 3-hydroxy-acyl-	08058.1	1nc100_110	1e-22, T:AP000996_69	None	None	5e-19, SP:HCDH_RAT	4e-4, PIR:T08956
Cox translation protein CYA5 CoxI translation protein CYA5 Related to nicotinamide nucleotide	$08692.1 \\ 01140.1$	$4nc677_080$ 13e11_040	1.0e-10, T:AB008268_15 0.0, PIR:T15521	None None	None None	None 0.0, T:BTNAD_1	1.0e-10, T:AB008268_15 None
transhydrogenase Related to light-induced alcohol	01107.1	b1308_020	5.2e-29, TN:CEE04F6_10	None	3.3e-23, PIR:T41570	None	9.9e-27, T:AC020666_24
denydrogenase bir-4 Related to kinetoplast-associated	04650.1	5f3_190	3.6e-30, PIR:A44937	None	None	None	None
Protein NAT Related to lysophosphatidic acid	06460.1	3nc220_480	1.6e-55, PIR:T40420	None	1.6e-55, PIR:T40420	1.5e-08, PIR:JH0610	None
puospiiatase Related to succinate-CoA ligase	09810.1	1 nc250_090	1.7e-78, PIR:S61696	1.7e-78, PIR:S61696	None	1.8e-68, SP:SUCA_PIG	5.0e-70, PIR:T51816
Mitochondrial carrier protein	06662.1	100h1_060	3.4e-34, PIR:T38879	None	3.4e-34, PIR:T38879	1.7e-33, SN:MCAT_MOUSE	3.7e-33, T:AC060755_11
Mitochondrial carrier protein Mitochondrial carrier protein	01564.1	b/n14_120 b2108_150	0.0e-07, FIK:S50285 3.9e-136, T:AF419344 1	None 3.9e-136, T:AF419344 1	None 2.4e-97, T:SPBC12D12 6	1./e-51, SF:FMI34_HUMAIN 4.3e-62, PIR:T50686	0.1e-51, FIK:F84825 6.9e-81, PIR:T49871
Mitochondrial carrier protein	07578.1		2.5e-21, PIR:F84823	1.3e-16, PIR:S69019 -	None	7.7e-13, T:XLA289240_1 6.6a 114_TN:DC024042_1	2.5e-21, PIR:F84823
Mitochondrial carrier protein	01810.1	b8b8_140	7.0e-44, PIR:T40082	8.8e-18, PIR:S60997	7.0e-44, PIR:T40082	1.1e-22, T:BC037680_1	1.4e-24, T:AC060755_11
Mitochondrial carrier protein	02352.1	7nc520_030	5.0e-27, PIR:B96830	1.1e-19, PIR:S61660	7.4e-21, PIR:138879	1.4e-26, TN:BC037680_1	5.0e-27, PIR:B96830
^a Not included are 40 proteins related	to respirate	ory complex I (subunits, assembly proteins and	l intron coded proteins). E-	values and codes of BLAST hi	its are given (T, TREMBL; TN, TF	REMBLNEW; SP, SWISSPROT;

TABLE 31. Known and deduced Neurospora mitochondrial proteins lacking a close homologue in S. cerevisiae^a

46 BORKOVICH ET AL.

SN, SWISSNEW). ^b ND, not determined.



FIG. 7. Pathway of sulfur acquisition leading to sulfur assimilation and cysteine biosynthesis in *Neurospora*. Potential sulfur sources from the environment or internal stores are indicated. The alkylsulfonate and cysteic acid conversions are predicted from the genome analysis. EC designations are shown; in some cases, putative multiple forms are indicated from the analysis (see Tables 32 and 35). Corresponding NCU numbers are as follows: EC 3.1.6.1 (arylsulfatase, NCU06041.1), EC 3.1.6.7 (choline sulfatase, NCU08364.1), EC 2.7.7.4 (ATP sulfurylase, NCU01985.1), EC 2.7.1.25 (adenylyl sulfate kinase, NCU0896.1), EC 1.8.99.4 (PAPS reductase, NCU02005.1), EC 4.1.1.15 (cysteic acid decarboxylase, NCU06112.1), EC 1.14.11.17 (taurine dioxygenase, NCU07610.1, NCU07819.1, NCU0978.1, NCU09800.1), EC 1.14.14.5 (alkanesulfonate monooxygenase, NCU05340.1, NCU10015.1), EC 1.8.1.2 (sulfite reductase, NCU04077.1, NCU05238.1), and EC 2.5.1.47 (cysteine synthase, NCU02564.1, NCU03788.1, NCU06452.1). Abbreviations: APS, adenosine-5'-phosphosulfate; PAPS, 3'-phosphoadenosine-5'-phosphosulfate.

drial functions have been assigned, these represent only a subset of mitochondrial metabolism.

Sulfur Metabolism

Sulfur acquisition and processing. When a nutrient such as sulfur is limiting, fungal cells will exhibit responses targeted to specifically alleviate the nutrient deficiency. In Neurospora, the responses include induction of transport systems (permeases) and enzymes that can release sulfur from internal or external sources (506, 584). These responses allow Neurospora to scavenge sulfur from a variety of possible sources. Previously unknown capabilities related to Neurospora sulfur acquisition are revealed from the genomic data. Of particular interest are two putative alkanesulfonate monooxygenases that can use aliphatic sulfonates (e.g., ethanesulfonic acid and isethionic acid but not compounds such as taurine) to generate sulfite, which can be used in the synthesis of cysteine (Fig. 7). The two putative alkanesulfonate monooxygenases show similarities only to bacterial alkanesulfonate monooxygenases (Table 32) (214) and have no significant matches to known eukaryotic proteins, they may represent a case of horizontal gene transfer.

The presence of these putative alkanesulfonate monooxygenases would give *Neurospora* the versatility to use an important class of sulfur compounds typically found in the environment. Sulfonates and sulfate esters represent the bulk of the sulfur content of aerobic soils (411).

Both cysteic acid and taurine are also potential sulfur sources for Neurospora. Analysis of the Neurospora genome suggests that the metabolism of cysteic acid is likely to occur by conversion into taurine. Extensive iterative database searching did not reveal other metabolic routes for either cysteic acid and taurine besides those shown in Fig. 7 (e.g., no homologues of sulfinoalanine decarboxylase, cysteine lyase, hypotaurine dehydrogenase, or other related enzymes involved in taurine and hypotaurine metabolism in a variety of other organisms were identifiable). Cysteic acid decarboxylase (also known as glutamate decarboxylase), which can convert cysteic acid to taurine, is present. Surprisingly, the putative Neurospora cysteic acid/ glutamate decarboxylase is most homologous to human GDC isoform 67 and has no identifiable homologs in yeast or plants (Table 32). This area of metabolism is relatively unexplored in Neurospora. In mammalian systems, taurine is an abundant

TABLE 32. Sulfur acquisition and processing

	FO	NOL		BLAST match			
Enzyme	EC no.	NCU no.	Best overall	S. cerevisiae	S. pombe	Animal ^a	Plant ^b
Arylsulfatase (ARS-1)	3.1.6.1	06041.1	P. anserina Arylsulf./0.0	c	_	2e-49	
Choline sulfatase	3.1.6.7	08364.1	P. aeruginosa Chol. Sulf./e-162	_	2e-15	1e-29	_
Alkanesulfonate monooxygenase	1.14.14.5	10015.1	P. putida MsuD/7e-126	_			_
Alkanesulfonate monooxygenase	1.14.14.5	05340.1	R. solanacearum SsuD/3e-36	_			_
Cysteic acid decarboxylase	4.1.1.15	06112.1	H. sapiens Gdc67/3e-74	_		3e-74	3e-17
Taurine dioxygenase	1.14.11.17	09738.1	S. pombe SCPB1C11.04C/6e-73	_	6e-73		_
Taurine dioxygenase	1.14.11.17	09800.1	Y. pestis Tau. Diox./3e-28	5e-27			_
Taurine dioxygenase	1.14.11.17	07610.1	E. coli TauD/3e-27	4e-26			_
Taurine dioxygenase	1.14.11.17	07819.1	S. cerevisiae Y11057cp/4e-36	4e-36			_
Cysteine dioxygenase	1.13.11.20	06625.1	H. sapiens cys. diox./3e-37	_		3e-37	_
Sulfite oxidase	1.8.3.1	04474.1	H. sapiens SUOX/1e-48	_		1e-48	1e-47
Sulfite oxidase	1.8.3.1	06931.1	C. elegans XQ117/9e-88	—	—	9e-88	3e-46

^a Caenorhabditis elegans, Drosophila melanogaster, Mus musculus, or Homo sapiens.

^b Arabidopsis thaliana or Oryza sativa.

^c Values below e-10 are represented as —

T	TO	NCU		BLAST match			
Transporter	IC no.	NCU no.	Best overall	S. cerevisiae	S. pombe	Animal ^a	Plant ^b
Sulfate permease I (CYS-13)	2A.53.1.2	03235.1	P. chrysogenum SutB/0.0	e-137	e-158	2e-56	8e-52
Sulfate permease II (CYS-14)	2.A.53.1.2	04433.1	P. chrysogenum SutB/e-179	e-112	e-134	2e-30	1e-27
Sulfate permease	2.A.53.1.2	09642.1	S. cerevisiase Ypr003cp/e-112	e-112	3e-80	4e-24	3e-41
Sulfate permease	2.A.53.1.2	02632.1	S. pombe SPAC24H6.11c/0.0	e-146	0.0	c	_
Methionine permease	2.A.3.8.4	02195.1	S. cerevisiae Mup1p/e-75	1e-75	_	4e-17	_
Methionine permease	2.A.3.8.4	04942.1	S. cerevisiae Mup1p/4e-94	4e-94	_	3e-25	_
Methionine Permease	2.A.3.8.4	07754.1	S. cerevisiae Mup1p/6e-87	6e-87	_	4e-32	_
Chromate resistance efflux	2.A.51.1.3	01055.1	A. aceti ChrA/7e-22	_			_

TABLE 33. Sulfur transporters

^b Arabidopsis thaliana or Oryza sativa.

^c Values below e-10 are represented as ---.

intracellular free amino acid with incompletely defined and disparate roles in brain development, osmolarity, and bile function (244). Taurine metabolism in Neurospora appears to be routed solely by conversion into sulfite by taurine dioxygenase. The sulfite can be subsequently converted to sulfide by sulfite reductase, and the sulfide can be used by cysteine synthase to generate cysteine (Fig. 7). An unusual finding is that four putative Neurospora taurine dioxygenases appear to be present (Table 32), two with best matches to bacterial enzymes and two with best matches to fungal enzymes (213, 815, 824). No matches for the putative taurine dioxygenases were found in animals or higher plants (Table 32).

Interestingly, the Neurospora genome encodes only two sulfatases. Arylsulfatase (ARS-1; NCU06041.1) (584) releases sulfate from aromatic sulfate compounds, while a choline sulfatase (NCU08364.1) releases sulfate from choline-O-sulfate (which also may serve as a sulfur storage compound). Homologues to the two Neurospora sulfatases were not observed in S. cerevisiae, S. pombe, or higher plants (Table 32). The Neurospora arylsulfatase could serve as a unique and useful model for the homologous multiple and specialized sulfatases in mammalian systems.

Sulfate can be transported into the cell by the sulfate permeases encoded by cys-13 (primarily conidial expression,) and cys-14 (primarily mycelial expression) (412). Since cys-13 cys-14 double mutants cannot use sulfate for growth (504), it was unexpected that the genomic data reveals the presence of two additional sulfate permeases that are homologous to those in other fungal and yeast species (Table 33). Presumably, the two additional putative sulfate permeases have specialized functional roles during other phases of the Neurospora life cycle that have not yet been studied with regard to sulfate transport. By comparison, only two genes encoding sulfate permeases in S. cerevisiae are known (SUL1 and SUL2) and have been found to encode high-affinity sulfate transporters (788). In contrast, A. thaliana has 14 isoforms in the sulfate transporter family, probably reflecting a variety of specific roles in plants (320).

The identification of a Neurospora gene showing homology to the ChrA chromate resistance gene of Pseudomonas aeruginosa (and Acetobacter aceti), which functions by chromate efflux (616), is intriguing. No significant homology to any eukaryotic genes is observed for this putative member of the chromate ion transporter (TC 2.A.51.1.3) family (Table 33), and this may represent another potential case of horizontal gene transfer. The capacity for chromate efflux in eukaryotes is currently unknown. The connection to sulfur metabolism is that both sulfate and chromate (which is toxic) are transported into the cell by sulfate permeases (504). On a related note, the bacterial ChrA family homologues are sulfur regulated (569).

Finally, analysis of the Neurospora genome reveals three methionine permeases (Table 33). The three putative methionine permeases show significant homology to MUP1 of S. cerevisiae, which encodes a high-affinity methionine permease (366). Available data from studies of S. cerevisiae suggest that MUP1 is also involved in cysteine uptake (429). MUP2 and MUP3 in yeast encode low-affinity permeases (788). Thus, Neu-

F	NOU		BLAST mat	ch		
Enzyme	NCU no.	Best overall	S. cerevisiae	S. pombe	Animal ^a	Plant ^b
bZIP activator (CYS-3)	None	A. nidulans MetR/3e-19	c	1e-19		_
F-box/WD-40 (SCON-2)	None	A. nidulans SconB/0.0	3e-114	3e-86	5e-37	3e-21
Skp1 homologue (SCON-3)	08991.1	M. canis SconC/5e-73	4e-35	3e-45	2e-36	5e-34
Rbx1 homologue	06224.1	S. salar Shop21/2e-43	4e-28	7e-38	2e-43	2e-40
Cullin ^d	05204.1	A. nidulans Cu1A/0.0	9e-107	2e-170	0.0	2e-91
Cullin ^d	00272.1	D. melanogaster Cu14/3e-118	7e-40	4e-95	3e-118	2e-107
Cullin ^d	00512.1	H. sapiens Apc2/1e-32	1e-15	1e-30	1e-32	6e-28
Cullin ^d	02498.1	H. sapiens Cu1-3/e-126	2e-45	1e-115	2e-128	3e-120

TABLE 34. Sulfur regulatory proteins

^a Caenorhabditis elegans, Drosophila melanogaster, Mus musculus, or Homo sapiens.

^b Arabidopsis thaliana or Oryza sativa.

^c Values below e-10 are represented as -

^d All cullins are listed that might be involved in the sulfur-related SCF complex.

Farmer	EC	NCU		BLAST match			
Enzyme	EC no.	NCU no.	Best overall	S. cerevisiae	S. pombe	Animal ^a	Plant ^b
ATP sulfurylase (CYS-11)	2.7.7.4	01985.1	A. fumigatus ATP sulf./0.0	e-152	3e-95	5e-28	2e-28
Adenylyl sulfate kinase	2.7.1.25	0896.1	A. nidulans APS kin./3e-75	2e-74	1e-73	5e-60	3e-51
PAPS reductase (CYS-5)	1.8.99.4	02005.1	P. chrysogenum ParA/4e-92	5e-69	8e-71	c	_
Sulfite reductase (Cys-2, beta chain)	1.8.1.2	04077.1	S. cerevisiae Met10p/e-167	e-167	e-162	7e-36	5e-27
Sulfite reductase (CYS-4, alpha chain)	1.8.1.2	05238.1	S. pombe SPAC4C5.05c/0.0	0.0	0.0	_	2e-99
Serine acetyl-transferase	2.3.1.30	00536.1	A. nidulans CysA/0.0	4e-32	5e-145	_	_
Cysteine synthase	2.5.1.47	06452.1	A. nidulans CysB/5e-167	2e-50	1e-122	3e-47	6e-39
Cysteine synthase	2.5.1.47	02564.1	S. pombe SPAC3A12.17c/2e-92	2e-88	2e-92	1e-36	2e-72
Cysteine synthase	2.5.1.47	03788.1	S. coelicolor Cys.Syn./2e-15	5e-11	1e-13	4e-10	7e-12

TABLE 35. Generation of sulfide and cysteine

^b Arabidopsis thaliana or Oryza sativa.

^c Values below e-10 are represented as -...

rospora appears to encode three high-affinity methionine transporters for which individual specialized functions or patterns of expression are currently unknown. Low-similarity animal genes exist, but no homologues are found in higher plants (Table 33). On sulfur starvation in *Neurospora*, there is the coordinate induction of an extracellular protease along with methionine permease activity. In yeast, Ssy1p is involved in sensing external amino acids and couples amino acid availability to transcriptional regulation of transporters (426). A homologue to Ssy1p cannot be identified in the *Neurospora* genome.

Generation of sulfide and cysteine. Analysis of the genomic data has allowed for the identification of all genes encoding the proteins necessary for the assimilation of sulfur as sulfate, sulfite, or sulfide (Fig. 7; Table 35). These sulfur assimilation enzymes confer on Neurospora the ability to synthesize cysteine from a variety of sulfur compounds. The findings from the genomic data are in agreement with biochemical evidence for the pathway (612), as follows: (i) ATP sulfurylase (cys-11) produces adenosine-5'-phosphosulfate (APS), (ii) adenylyl sulfate kinase produces 3'-phosphoadenosine-5'-phoshosulfate (PAPS), (iii) PAPS reductase (cys-5) generates sulfite, (iv) sulfite reductase (cys-4 alpha subunit; and cys-2 beta subunit) converts the sulfite into sulfide, and finally, (v) cysteine synthase uses the sulfide and O-acetyl serine (provided by serine acetyltransferase) for the synthesis of cysteine. The last step in the pathway provided an interesting finding in that there are three putative cysteine synthase homologues (NCU02564.1, NCU03788.1, and NCU06452.1) that each contain an identical or close match to the cysteine synthase motif (557), including a lysine residue which binds to pyridoxal 5'-phosphate. Each putative cysteine synthase has its most similar match to a different organism (i.e., A. nidulans, S. pombe, and Streptomyces coelicolor [Table 35]). The presence of two isoforms of cysteine synthase (cytoplasmic and mitochondrial) would be predicted. The presence of multiple cysteine synthases is known to occur in higher plants; rice has four genes encoding cysteine synthases that encode functionally distinct isoforms (557).

In relation to internal sulfur cycling, two key genes identified from the *Neurospora* genomic data suggest similarities to animal sulfur metabolism. The degradation of cysteine appears to occur by conversion to 3-sulfino-L-alanine catalyzed by cysteine dioxygenase. Significant similarity was observed only for mammalian cysteine dioxygenase (443, 602); no matches were observed for yeasts or higher plants (Table 32). The subsequent conversion to 3-sulfino-pyruvate is probably carried out by aspartate transaminase. Finally, 3-sulfino-pyruvate yields sulfite and pyruvate in an apparent nonenzymatic reaction. Sulfite can be used as in Fig. 7 or in a reaction catalyzed by sulfite oxidase to generate sulfate. Sulfite oxidase is regarded as a terminal enzyme in the degradation pathway of sulfur amino acids and can eliminate endogenously produced sulfite or detoxify exogenously added sulfite. Sulfite oxidase is present in *Neurospora* (Table 32), with the highest homology to *H. sapiens* and *C. elegans* and less similarity to higher plants. The presence of sulfite oxidase has only been recently established in higher plants (e.g., *A. thaliana* [216]). No matches were observed with *S. cerevisiae* or *S. pombe* for the putative *Neurospora* sulfite oxidases.

Homocysteine and methionine metabolism. Analysis of the Neurospora genomic data reveals homologues that cover every needed step for the interconversion of methionine and cysteine and for the SAM cycle (Table 36). Cystathionine is generated by cystathionine gamma-synthase (NCU08117.1; NCU05093.1; and *met-7*, NCU02430.1) from cysteine (derived as in Fig. 7) and O-acetyl homoserine (derived from aspartate-\beta-semialdehyde by homoserine dehydrogenase and homoserine O-acetyl transferase, *met-5*). Cystathionine β -lyase (*met-2*) then cleaves cystathionine to generate homocysteine. Methionine is generated by methionine synthase, which utilizes homocysteine and 5-methyl tetrahydrofolate as a methyl donor. An unanticipated finding is the presence of three putative methionine synthases. The known met-8 locus is homologous to the typical eukaryotic cobalamin-independent methionine synthases (e.g., A. nidulans MetH/D). Two additional putative methionine synthases have the closest similarity to bacterial methionine synthases (e.g., Rhodopseudomonas palustris) and are also cobalamin independent. No significant matches to S. cerevisiae, S. pombe, animals, or higher plants were observed for these two proteins, and they may represent potential cases of horizontal gene transfer. Since met-8 results in a phenotype of sulfur auxotrophy under typical vegetative growth, the role(s) of the bacterium-like putative methionine synthases cannot be defined at present. The 5-methyl tetrahydrofolate is supplied for methionine synthesis by 5,10-methylene tetrahydrofolate reductase, which is encoded by two homologues, met-1 and a previously unknown gene (NCU09545.1). met-1 would appear

]	BLAST match			
EC no.	NCU no.	Best overall	S. cerevisiae	S. pombe	Animal ^a	Plant ^b
2.5.1.48	08117.1	S. pombe SPBC1504.09c/1e-42	1e-29	1e-42	c	_
2.5.1.48	05093.1	P. anserina CAC6069/2e-50	1e-43	2e-29	_	_
2.5.1.48	02430.1	P. anserina Pa5D0012/0.0	7e-92	1e-131	3e-16	1e-15
1.1.1.3	03935.1	S. pombe SPBC776.03/1e-86	4e-85	1e-86	_	1e-70
2.3.1.31	07001.1	A. nidulans MetE/0.0	2e-142	9e-147	_	_
4.4.1.8	07987.1	B. fuckeliana meC/0.0	2e-97	3e-81	3e-74	1e-100
2.1.1.13	06512.1	A. nidulans MetH/D/0.0	0.0	0.0	1e-95	0.0
2.1.1.13	08434.1	R. palustris ZP8831/8e-44	_	_	_	_
2.1.1.13	10020.1	R. palustris ZP8831/8e-44	_	_	_	_
2.5.1.49	01652.1	A. nidulans CysA/0.0	e-141	2e-148	9e-43	1e-39
6.3.2.17	00892.1	S. cerevisiae Met7p/4e-104	4e-104	4e-98	1e-75	3e-60
6.3.2.17	01337.1	S. cerevisiae Fo13p/5e-65	5e-65	3e-64	2e-29	4e-27
2.1.2.1	02274.1	C. albicans SHM2/0.0	0.0	0.0	4e-151	1e-153
2.1.2.1	05805.1	S. pombe Shm2/3e-168	3e-164	3e-168	1e-153	3e-167
4.2.1.22	07690.1	S. pombe MTHFR2/4e-174	3e-168	4e-174	1e-135	4e-138
4.2.1.22	09545.1	S. cerevisiae Met12p/8e-143	8e-143	2e-142	3e-83	9e-94
2.5.1.6	06512.1	S. pombe Sam1/1e-155	e-145	1e-155	2e-141	4e-118
2.5.1	03032.1	R. sphaeroides btaA/2e-18	_	_	_	_
3.3.1.1	07930.1	S. cerevisiae Sah1p/0.0	0.0	0.0	1e-80	5e-142
4.2.1.22	08216.1	M. grisea CBS1/0.0	1e-118	2e-47	5e-107	5e-49
4.4.1.1	09230.1	A. chrysogenum MecB/3e-178	2e-129	5e-44	4e-96	1e-65
	EC no. 2.5.1.48 2.5.1.48 1.1.1.3 2.3.1.31 4.4.1.8 2.1.1.13 2.1.2.1 4.2.1.22 4.2.1.21 4.2.1.22 4.2.1.21 4.2.1.21 4.2.1.22 4.2.1.21 4.2.1.21 4.2.1.21 4.2.1.22 4.2.1.21 4.2.1.21 4.2.1.22 4.2.1.21 4.2.1.22 4.2.1.21 4.2.1.22 4.2.1.21 4.2.1.22 4.2.1.21 4.2.1.22 4.2.1.21 4	EC no. NCU no. 2.5.1.48 08117.1 2.5.1.48 05093.1 2.5.1.48 02430.1 1.1.1.3 03935.1 2.3.1.31 07001.1 4.4.1.8 07987.1 2.1.1.13 06512.1 2.1.1.13 00892.1 6.3.2.17 01892.1 6.3.2.17 00892.1 4.2.1.22 07690.1 4.2.1.22 07690.1 4.2.1.22 09545.1 2.5.1.6 06512.1 2.5.1.7 03032.1 3.3.1.1 07930.1 4.2.1.22 08216.1 4.2.1.22 08216.1	EC no. NCU no. Best overall 2.5.1.48 08117.1 S. pombe SPBC1504.09c/1e-42 2.5.1.48 05093.1 P. anserina CAC6069/2e-50 2.5.1.48 02430.1 P. anserina Pa5D0012/0.0 1.1.1.3 03935.1 S. pombe SPBC776.03/1e-86 2.3.1.31 07001.1 A. nidulans MetE/0.0 4.4.1.8 07987.1 B. fuckeliana meC/0.0 2.1.1.13 06512.1 A. nidulans MetH/D/0.0 2.1.1.13 08434.1 R. palustris ZP8831/8e-44 2.5.1.49 01652.1 A. nidulans CysA/0.0 6.3.2.17 00892.1 S. cerevisiae Fo13p/5e-65 2.1.2.1 02274.1 C. albicans SHM2/0.0 2.1.2.1 02805.1 S. pombe Shm2/3e-168 4.2.1.22 07690.1 S. pombe MTHFR2/4e-174 4.2.1.22 09545.1 S. cerevisiae Met12p/8e-143 2.5.1.6 06512.1 S. pombe Sam1/1e-155 2.5.1.6 06512.1 S. pombe Sam1/1e-155 2.5.1.6 06512.1 S. cerevisiae Sah1p/0.0 4.2.1.22 08216.1 <td< td=""><td>$\begin{array}{c c c c c c c c c c c c c c c c c c c$</td><td>$\begin{array}{c c c c c c c c c c c c c c c c c c c$</td><td>$\begin{array}{c c c c c c c c c c c c c c c c c c c$</td></td<>	$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	$ \begin{array}{c c c c c c c c c c c c c c c c c c c $

TABLE 36. Homocysteine and methionine metabolism

^b Arabidopsis thaliana or Oryza sativa.

^c Values below e-10 are represented as -.

to function during typical vegetative growth, while a role for the other version of tetrahydrofolate reductase awaits determination. The balance of folate metabolism appears typical and complete (Table 36) (612).

SAM synthetase (*eth-1*) generates SAM from methionine and ATP. Subsequently, SAM is used for a wide variety of methyltransferase reactions (e.g., DNA methylation). After donating a methyl group, SAM is converted into *S*-adenosylhomocysteine. A typical array of eukaryotic methyltransferases, about twice the number found in yeasts, are represented in the *Neurospora* genome. An interesting side note is that a biosynthetically rare SAM-dependent 3-amino-3-carboxypropyltransferase of the MAPEG (membrane-associated proteins in eicosanoid and glutathionine metabolism) class is present and shows similarity to a *Rhodobacter sphaeroides* protein with no significant hits to eukaryotic proteins (Table 36) (425). The SAM cycle can be completed by conversion of S-adenosylhomocysteine back to homocysteine (adenosylhomocysteinase), followed by regeneration of methionine by methionine synthase. Additionally, homocysteine can converted to cystathionine by β -cystathionine synthase. Cystathionine γ -lyase (*cys-16*) then converts the cystathionine into cysteine, thus completing the conversion of methionine into cysteine.

Additional aspects of sulfur metabolism. Glutathione metabolism has not been studied in detail in *Neurospora*. Glutathione *S*-transferases (GSTs) represent an important group of enzymes involved primarily in detoxification reactions (197). Three GSTs are of particular interest and do not show homology to yeast GSTs (Table 37). A putative microsomal GST is closest to animal and plant GSTs of that class, while a putative mitochondrial GST (NCU06494.1) is most similar to a human GST. The third predicted protein (NCU02888.1) is similar only to fungal GSTs (e.g., *Gibberella fujikuroi*).

TABLE	37.	Additional	aspects	of	sulfur	metabolism
TTDLL	<i></i>	1 Iddittonut	aspects	01	ounui	metuoonom

	EC	NCU		BLAST match			
Enzyme	EC no.	NCU no.	Best overall	S. cerevisiae	S. pombe	Animal ^a	Plant ^b
Glutathione S-transferase (microsomal)	2.5.1.18	01320.1	O. sativa Gst/3e-15	c	_	2e-12	3e-15
Glutathione S-transferase (mito.)	2.5.1.18	06494.1	H. sapiens GST13/5e-13	_		5e-13	
Glutathionine S-transferase	2.5.1.18	02888.1	G. fujikuroi GST/2e-55	_	_	_	_
MET-10 (unknown function)		09311.1	S. pombe SPAPB18E9.01/5e-73	9e-61	5e-73	4e-54	4e-43
Thioredoxin reductase (CYS-9)	1.6.4.5	08352.1	P. chrysogenum TrxB/e-132	e-120	e-113		e-103
Thioredoxin		00598.1	S. pombe SPC57.08c/1e-16	1e-15	1e-16	8e-14	3e-13
Thioredoxin		06556.1	P. anserina Trx2/1e-27	1e-19	6e-21	1e-16	8e-113
Halotolerance PAPS phosphatase (CYS-1?)	3.1.3.7	04069.1	S. pombe SPCC1753.04/1e-83	2e-62	1e-83		4e-59
Sulfide dehydrogenase	1.8.5	07112.1	A. oryzae BAC55902/3e-175		7e-102	1e-78	
Cysteine desulfurase	4.4.1	04636.1	C. albicans NFS1/0.0		5e-180	2e-177	1e-157
Methionine sulfoxide	1.8.4.6	100029.1	S. pombe SPAC30.09c/4e-41	6e-38	4e-41	2e-27	1e-29

^a Caenorhabditis elegans, Drosophila melanogaster, Mus musculus, or Homo sapiens.

^b Arabidopsis thaliana or Oryza sativa.

^c Values below e-10 are represented as -...

NCU no	Clanad gana		BLAS	score		
NCO IIO.	Cloned gene	Best	S. cerevisiae	S. pombe	Animal ^a	Plant ^b
Regulatory genes						
09068.1	nit-2	<i>C. lindemuthianum</i> AAN65464.1; 1.00e-139	Gat1p; 9.00e-14	AAC35593.1; 1.00e-13	Mm GATA-2; 3.00e-08	None
08294.1	nit-4	T. inflatum CAB71797.1; 0.00	Tea1p; 3.00e-17	SPCC757.04; 4.00e-30	None	None
04158.1	nmr	G. fujikuroi CAA75863.1; 1.00e-154	None	None	Hs XP 293629.1; 8.00e-06	None
07669.1	pco-1	A. nidulans uaY; 0.00	Ppr1p; 5e-21	SPBC530.05; 4e-09	None	None
Structural genes						
05298.1	nit-3/nitrate reductase	M. anisopliae CAA04554.1; 0.00	Yml125cp; 3.00e-28	SPCC970.03; 4e-24	Dm CG7280-PA; 1.00e-42	At nitrate reductase; 1.00e-156
04720.1	nit-6/nitrite reductase	P. nodorum CAA08856.1; 0.00	None	None	Ag XP 306456.1; 1.00e-166	None
07205.1	Nitrate permease	A. fumigatus CAD28427.1; 7.00e-77	None	None	None	NP_172754.1; 3.00e-20
01816.1	alc/allantoicase	S. pombe NP_594495.1; 1.00e-104	Dal2p; 3.00e-94	Best match	Mm allantoicase; 1.00e-44	None
03350.1	<i>xdh</i> /xanthine dehy- drogenase	A. nidulans Xdh; 0.00	None	None	Hs NP_000370.1; 0.00	At NP_195215.2; 0.00

TABLE 38. Nitrogen assimilation and regulation

^a Caenorhabditis elegans (Ce), Drosophila melanogaster (Dm), Anopheles gambiae (Ag), Mus musculus (Mm), or Homo sapiens (Hs).

^b Arabidopsis thaliana (At) or Oryza sativa (Os).

A number of additional genes that are involved in routine sulfur metabolic activities and that do not show unusual homology patterns are also included in Table 37.

Components of the regulatory machinery for sulfur metabolism. The complex regulatory network involved in the control of sulfur metabolism provides a fascinating model for understanding how fungi (and other organisms) adjust to a constantly changing nutrient environment in order to maintain sustaining levels of key elements (e.g., carbon, nitrogen, phosphorous, and sulfur) (586). An important aspect of the sulfur regulatory system of Neurospora is the role of the F-box protein sulfur controller-2 (scon-2) (440). F-box proteins assemble with Skp-1p, Cdc53p, and Rbx1p to form a complex known as the SCF (Skp1p/Cullin/F-box) (171). SCF complexes act as E3 ubiquitin ligases to target proteins for ubiqitin-mediated proteolysis by the proteasome (see "Proteasome" below). The SCF complex is represented in the Neurospora genome by SCON3 (741) as the Skp1p homolog, SCON2 as the F-box protein, NCU06224.1 as the Rbx1p (ring box) homologue, and four potential cullin components (Table 34). Additional F-box proteins, probably unrelated to sulfur metabolism, are also present in the *Neurospora* genome. Extensive searches did not reveal homologues of known sulfur regulatory proteins found in other organisms (e.g., the sac1, sac2, and sac3 genes of Chlamydomonas reinhardtii [182]). Major unanswered regulatory questions remain with regard to the identity of the sulfur sensor and subsequent signal transduction pathway. Functional genomic studies will provide an important future means of identifying other regulatory proteins in the sulfur control system.

Nitrogen Metabolism

A basic understanding of nitrogen metabolism and its regulation exists for *Neurospora* and includes the identification and characterization of many of the genes that participate in this system (Table 38). Neurospora utilizes ammonia and glutamine as preferred nitrogen sources but is also capable of using many different secondary nitrogen sources (505). The use of various alternative nitrogen sources, e.g., nitrate, purines, amides, amino acids, and proteins, requires the expression of specific sets of structural genes which encode enzymes for catabolism of a particular nitrogenous source. A globally acting regulatory gene, nit-2, encodes a DNA binding protein, a member of the family of GATA transcription factors (522) (see "Chromatin assembly and gene regulation" above). NIT-2 acts in a positive fashion to activate the expression of structural genes of diverse nitrogen pathways under conditions of nitrogen source limitation. A major negatively acting regulatory protein, NMR (for "nitrogen metabolite repression") mediates nitrogen repression and acts by directly binding to NIT-2 and inhibiting its function (590). Activation of the genes of specific pathways requires inducers and is mediated by minor control genes. A pathway-specific gene, nit-4, codes for a positively acting protein with a binuclear zinc finger DNA binding motif (257, 892) (see also "Chromatin assembly and gene regulation" II). NIT-4 mediates the induction of enzymes specific for assimilation of inorganic nitrate. The regulatory genes (*nit-2*, *nmr*, and *nit-4*) and the structural genes encoding nitrate reductase (nit-3), nitrite reductase (nit-6), and allantoicase (alc) were cloned by complementation of mutants (258, 259, 261, 460). These genes have been extensively characterized and demonstrated to have homologues in A. nidulans and other filamentous fungi.

Several new genes involved in nitrogen metabolism have recently been identified by analysis of the *Neurospora* genome sequence (Table 38). The gene encoding a nitrate permease (*nit-10*), which had never been revealed by conventional genetics despite extensive work in this pathway, was identified in the *Neurospora* genome sequence and its function was demonRegulatory

RPT6

		TAB	LE 39. Neurospora proteasome compo	onents	
Role	Subunit	Neurospora gene	Alias(es)	Symbol	Gene name
Catalytic	alpha-1	EAA28671, NCU10061.1	C7	pca-1	Proteasome catalytic alpha-1
Catalytic	alpha-2	EAA28122, NCU06764.1	Y7	pca-2	Proteasome catalytic alpha-2
Catalytic	alpha-3	EAA29550, NCU05942.1	Y13	pca-3	Proteasome catalytic alpha-3
Catalytic	alpha-4	EAA28095, NCU06440.1	PRE6	pca-4	Proteasome catalytic alpha-4
Catalytic	alpha-5	EAA32830, NCU05295.1	PUP2	pca-5	Proteasome catalytic alpha-5
Catalytic	alpha-6	EAA31656, NCU06712.1	PRE5	pca-6	Proteasome catalytic alpha-6
Catalytic	alpha-7	EAA35851, NCU02493.1	C1	pca-7	Proteasome catalytic alpha-7
Catalytic	beta-1	EAA28906, NCU09290.1	PRE3	pcb-1	Proteasome catalytic beta-1
Catalytic	beta-2	EAA34801, NCU08605.1	PUP1	pcb-2	Proteasome catalytic beta-2
Catalytic	beta-3	EAA35245, NCU03304.1	PUP3	pcb-3	Proteasome catalytic beta-3
Catalytic	beta-4	EAA31484, NCU01368.1	PRE1 (C11)	pcb-4	Proteasome catalytic beta-4
Catalytic	beta-5	EAA29213, NCU09309.1	PRE2	pcb-5	Proteasome catalytic beta-5
Catalytic	beta-6	EAA34540, NCU09366.1	C5	pcb-6	Proteasome catalytic beta-6
Catalytic	beta-7	EAA28757, NCU07365.1	PRE4	pcb-7	Proteasome catalytic beta-7
Regulatory	RPN1	EAA32735, NCU07721.1	Non-ATPase subunit	rpn-1	Regulatory particle, non-ATPase-like-1
Regulatory	RPN2	EAA28800, NCU09450.1	Non-ATPase subunit	rpn-2	Regulatory particle, non-ATPase-like-2
Regulatory	RPN3	EAA30401, NCU02224.1	Non-ATPase subunit	rpn-3	Regulatory particle, non-ATPase-like-3
Regulatory	RPN4	EAA26772, NCU01640.1	Non-ATPase subunit	rpn-4	Regulatory particle, non-ATPase-like-4
Regulatory	RPN5	EAA36187, NCU02650.1	Non-ATPase subunit	rpn-5	Regulatory particle, non-ATPase-like-5
Regulatory	RPN6	EAA27271, NCU01596.1	Non-ATPase subunit 4	rpn-6	Regulatory particle, non-ATPase-like-6
Regulatory	RPN7	EAA28375, NCU03972.1	Non-ATPase subunit	rpn-7	Regulatory particle, non-ATPase-like-7
Regulatory	RPN8	EAA27014, NCU01547.1	Non-ATPase subunit	rpn-8	Regulatory particle, non-ATPase-like-8
Regulatory	RPN9	EAA30559, NCU02374.1	Non-ATPase subunit 7	rpn-9	Regulatory particle, non-ATPase-like-9
Regulatory	RPN10	EAA36132, NCU02982.1	Non-ATPase subunit	rpn-10	Regulatory particle, non-ATPase-like-10
Regulatory	RPN11	EAA35130, NCU00823.1	Non-ATPase subunit 2, Pad1, Sks1	rpn-11	Regulatory particle, non-ATPase-like-11
Regulatory	RPN12	EAA28646, NCU10067.1	Non-ATPase subunit	rpn-12	Regulatory particle, non-ATPase-like-12
Regulatory	RPT1	EAA34894, NCU02840.1	AAA ATPase subunit 7, Cim5	rpt-1	Regulatory particle, ATPase-like-1
Regulatory	RPT2	EAA32354, NCU01224.1	AAA ATPase subunit	rpt-2	Regulatory particle, ATPase-like-2
Regulatory	RPT3	EAA30668, NCU02260.1	AAA ATPase subunit 6B, Ynt1	rpt-3	Regulatory particle, ATPase-like-3
Regulatory	RPT4	EAA29624, NCU07367.1	AAA ATPase subunit	rpt-4	Regulatory particle, ATPase-like-4
Regulatory	RPT5	EAA28255, NCU04414.1	AAA ATPase subunit 6A, Tbp1	rpt-5	Regulatory particle, ATPase-like-5

AAA ATPase subunit 8, Sug1, Cim3

strated by creation of a RIP mutant and parallel biochemical and molecular studies (272). In the purine catabolic pathway, xdh, which encodes xanthine dehydrogenase, was readily identified. A new gene, pco-1, which codes for a binuclear zinc DNA binding protein that regulates purine metabolism, was identified in the Neurospora genome sequence by its regions of homology to uaY, the corresponding factor of A. nidulans (T.-W. Liu and G. A. Marzluf, unpublished data). A pco-1 RIP mutant was created; analysis of phenotypes, in combination with DNA binding studies of the expressed protein, demonstrate that PCO-1 serves as a positive activator to induce expression of multiple enzymes of the purine catabolic pathway.

EAA34118, NCU05363.1

It appears that it should be possible to discover most, perhaps even all, of the genes involved in nitrogen metabolism or its regulation by careful analysis of the Neurospora genome sequence and subsequent functional studies. Significant questions in nitrogen metabolism remain. A postulated, but unknown, factor that senses glutamine as the initial step in nitrogen repression has yet to be identified. Similarly, other proteins in the signaling pathway for nitrogen repression that may converge on NIT-2 and NMR have yet to be identified. Additional factors that control specific metabolic pathways, as well as the way in which they interact with inducers and the globally acting NIT-2 protein, remain to be explored. Investigation of these areas and similar features will clearly be enhanced by the availability of the Neurospora genome sequence.

Proteasome

Regulatory particle, ATPase-like-6

rpt-6

The proteasome is one of the most complex oligomeric protein structures within the cell, and its function is the proteolysis of ubiquitin-tagged cellular proteins. Proteasomes are found within eukaryotic cells free in the cytoplasm, associated with the endoplasmic reticulum (ER), and in the nucleus. The structure and component polypeptide identities of the 26S proteasome and its 20S catalytic and 19S regulatory subunits have been studied (88). The subunits of both regulatory (285) and catalytic (327) components have been identified. The crystallographic structure of the S. cerevisiae proteasome has also been determined (303).

To date, none of the Neurospora genes encoding the subunits of the proteasome have been identified, although some proteins of the ubiquitin modification pathway have been isolated as regulatory genes in various pathways (e.g., "Sulfur metabolism", above). However, using the yeast polypeptide sequences as probes, all 14 polypeptides of the catalytic component and all 18 polypeptides of the regulatory component of the proteasome of Neurospora have now been identified (Table 39).

Lipids

Most work on Neurospora lipids has focused on understanding factors that control the synthesis and composition of the most common fatty acids and fatty acid-containing (acyl) lipids. These acyl lipids include both membrane lipids (phospholipids,

Engrand	EC no	NCU ao		BLA	ST match	
Enzyme	EC IIO.	NCU IIO.	Best overall	S. cerevisiae	Animal	Plant
Fatty acid synthesis						
Acetyl-CoA carboxylase	6.4.1.2	08535.1	0.0; E. nidulans	0.0		
Fatty acid synthase (alpha)	2.3.1.85	07308.1	0.0; E. nidulans	0.0		
Fatty acid synthase (beta)	2.3.1.86	07307.1	0.0; E. nidulans	0.0		
Fatty acid elongase	2.3.1.41	06694.1	4e-75; S. pombe	4e-73		
Fatty acid desaturation						
Fatty acid hydroxylase	1.14.15.	03492.1	1e-93; S. cerevisiae	Best match	3e-33; M. musculus	8e-35; O. sativa
Stearoyl-CoA desaturase	1.14.99.5	05259.1	0.0; Ajellomyces capsulatus	8e-121		
Oleate $\Delta 12$ desaturase	1.14.99.	02209.1	e-180; E. nidulans	5e-1		2e-80; S. oleracea
Oleate $\Delta 12$ desaturase	1.14.99.	09497.1	1e-78; A. parasiticus	3e-1		1e-68; S. oleracea
Fatty acid $\Delta 6$ desaturase	1.14.19.1	02408.1	0.0; P. anserina	2e-3		7e-54; T. aestivum
Fatty acid degradation						
Multifunctional beta-oxidation protein	1.1.1.35	08828.1	0.0; Y. lipolytica	0.0	e-141; H. sapiens	2e-52; A. thaliana
*	5.1.2.				· •	
	4.2.1.17					
Acyl group transfer						
Glycerol-3-phosphate acyltransferase	2.3.1.15	05985.1	e-150; S. pombe	e-132		
Lysophosphatidic acid acyltransferase	2.3.1	00168.1	2e-71; E. nidulans	1e-44	3e-24; M. musculus	
Diacylglycerol acyltransferase	2.3.1.20	02665.1	2e-89; M. ramanniana	1e-65	3e-68; H. sapiens	
Diacylglycerol acyltransferase	2.3.1.20	00035.1	3e-58; B. napus	2e-33	2e-51; D. melanogaster	Best match
Phospholipid acyltransferase	2.3.1	02416.1	0.0; P. anserina	e-137	ý 8	3e-52; A. thaliana
Sterol acyltransferase	2.3.1.43	04144.1	2e-69; S. pombe	7e-56		
Sterol acyltransferase	2.3.1.43	03991.1	4e-65; S. kluyveri	4e-79		
Serine palmitoyltransferase subunit 1	2.3.1.50	06870.1	0.0; E. nidulans	7e-85	4e-81; A. gambiae	3e-83; A. thaliana
Serine palmitoyltransferase subunit 2	2.3.1.50	00447.1	0.0; E. nidulans	e-157	e-139; M. musculus	e-131; S. tuberosom
Phospholipid synthesis						
Phosphatidylserine synthase	2.7.8.8	01141.1	e-107: S. cerevisiae	Best match	7e-66: D. melanogaster	
Phosphatidylserine synthase	2.7.8.8	02381.1	2e-76: T. aestivum	4e-59		Best match
Phosphatidylserine decarboxylase	4.1.1.65	01004.1	e-135: S. pombe	3e-96		7e-72: A. thaliana
Phosphatidylserine decarboxylase	4.1.1.65	03695.1	4e-94; S. cerevisiae	Best match	3e-52: H. sapiens	7e-54: A. thaliana
Phosphatidylserine decarboxylase	4.1.1.65	02302.1	8e-67: B. fungorum	3e-11	,	2e-21: A. thaliana
Phosphatidylethanolamine methyl-	2.1.1.17	08045.1	e-168: <i>S. pombe</i>	e-102		,
transferase						
Phosphatidyl-N-methylethanolamine	2.1.1.71	04699.1	8e-48; S. cerevisiae	Best match	3e-40; D. rerio	
methyltransferase						
Choline phosphotransferase	2.7.8.2	03223.1	8e-38; S. pombe	2e-32		1e-33; A. thaliana
Diacylglycerol choline/ethanolamine	2.7.8.1/2	01993.1	5e-70; S. pombe	3e-62		3e-45; B. napus
transferase						
CDP-alcohol transferase (probable	2.7.8	00135.1	2e-36; D. melanogaster	8e-29	3e-29; M. musculus	
cardiolipin synthase)						
Phosphatidylinositol synthase	2.7.8.11	09192.1	1e-49; S. pombe	2e-41	2e-42; R. norvegicus	
Choline kinase	2.7.1.32	03176.1	2e-58; S. pombe	2e-46		5e-28; P. sativum
Ethanolamine kinase	2.7.1.82	02726.1	2e-42; R. norvegicus		Best match	
Cholinephosphate cytidylyltransferase	2.7.7.15	03880.1	6e-80; S. pombe	1e-74		2e-59; P. sativum
Phosphoethanolamine cytidylyltransferase	2.7.7.14	04289.1	1e-63; D. melanogaster	3e-53	2e-60; R. norevgicus	
Inositol-3-phosphate synthase	5.5.1.4	06666.1	e-170; T. aestivum	e-291	e-169; H. sapiens	Best match
Phosphatidylcholine/-inositol exchange		02263.1	e-112; A. capsulatus	3e-80		
protein (sec14)		07220 1	7. 20. 6	Dest mot-1		
protein (sec14)		07520.1	10-29; S. cerevisiae	best match		

TABLE 40. Lipid metabolism

sphingolipids, and sterol esters) and storage lipids (triacylglycerols). *Neurospora* is an excellent model system for studying the roles of lipids: it synthesizes a broad range of lipids de novo, its lipid composition can be influenced by culture conditions, and many mutant strains have altered lipid metabolism (514, 612). Because of these traits, the biochemistry and genetics of *Neurospora* lipids are still among the best understood of any organism; in the fungal kingdom, more extensive studies have only been done for the yeast *S. cerevisiae* (128, 181). Characteristic of *Neurospora* is the accumulation of high levels of polyunsaturated fatty acids (linoleate and α -linolenate) and triacylglycerols under appropriate conditions (184, 485). The

de novo synthesis of these lipids is typical of plants and some other filamentous fungi (485, 563); *S. cerevisiae* does not synthesize detectable levels of polyunsaturates, and triacylglycerols are not as readily accumulated (181).

The major fatty acids of *Neurospora* are palmitate, stearate, oleate, linoleate, and α -linolenate; the relative proportion of these fatty acids in phospholipids and triacylglycerols is similar to that in many higher plants and is not typical of most fungi (289, 485, 825). The 16- and 18-carbon saturated fatty acids palmitate and stearate are synthesized from malonyl-CoA by a multifunctional fatty acid synthase, which consists of two subunits, α and β (219). In *S. cerevisiae*, the subunits are encoded on different chromosomes (694); however, in Neurospora, the genes are adjacent to each other on the same chromosome, oriented in opposite directions (Table 40), as in A. nidulans (108). The cel-1 mutant, which synthesizes only small amounts of fatty acids de novo due to impaired pantotheine binding activity of the α subunit, has been exploited extensively to study the effects of supplemental fatty acids on Neurospora physiology (612). The cel-2 mutants, obtained by RIP mutation of the β subunit, are blocked in fatty acid synthesis (288). S. cerevisiae deletion mutants with mutations of genes for either fatty acid synthase subunit require supplemental fatty acids for growth, and deletions of genes for acetyl-CoA carboxylase (which carries out the committed step in fatty acid biosynthesis, conversion of acetyl-CoA to malonyl-CoA) are lethal (181). A few predicted Neurospora proteins have homology to proteins carrying out individual enzymatic activities of fatty acid synthases (Table 40). Some of these proteins (for example, that encoded by NCU000563) are likely to be involved in mitochondrial β-oxidation or in the minor amount of fatty acid synthesis that takes place in mitochondria (528), which, similarly to bacterial cells, utilize individual enzymes to carry out fatty acid biosynthesis. Some seven other predicted proteins can be identified as probable polyketide synthases. Polyketides, common secondary metabolites in filamentous fungi, are synthesized by a pathway that resembles fatty acid biosynthesis (348). Although a polyketide synthesizes polyunsaturated fatty acids de novo in a few organisms (842), biochemical studies indicate that Neurospora, like higher plants and other fungi that synthesize polyunsaturated fatty acids, derives its polyunsaturates from stearate and oleate precursors utilizing a phospholipid substrate (41, 289).

A desaturase first converts stearoyl-CoA to oleoyl-CoA (41), as in other fungi and animals (563). Membrane-bound $\Delta 12$ and $\Delta 15$ desaturases carry out the final two consecutive desaturation steps, to linoleate and α -linolenate, respectively, after oleate has become incorporated into phosphatidylcholine (41), the major phospholipid in Neurospora and most other eukaryotes. The relative levels of α -linolenate, particularly in phospholipids, are affected by many factors, including growth temperature, developmental stage, and the circadian clock (612). Regulation of the desaturase genes is therefore of interest. Neurospora has at least three loci that control formation of oleate (612); however, a single candidate gene for a stearoyl-CoA desaturase has been identified based on homology, and in BLAST searches the sequence appears to be enriched in an EST library from sequences expressed during the evening phase of the circadian cycle (900). In common with the S. cerevisiae protein (588), the Neurospora enzyme contains a cytochrome b sequence. Membrane-bound desaturases have been best characterized in higher plants, most extensively in A. thaliana (60); to date, only a few fungal desaturases have been identified, and they form different polyunsaturates, generally in the $\Delta 6$ fatty acid series (including γ -linolenate). The highest homologies of candidate Neurospora proteins to membranebound desaturase sequences are to fungal $\Delta 12$ desaturases (which form linoleate) and to a putative fungal desaturase that itself has highest homology to a $\Delta 6$ sphingolipid desaturase (Table 40). In plants, $\Delta 6$, $\Delta 12$, and $\Delta 15$ desaturases and hydroxylases have high homology (107, 516); given that other fungal $\Delta 15$ desaturases have not been identified and that Neurospora does not synthesize detectable levels of $\Delta 6$ fatty acids, it is likely that either the putative $\Delta 6$ or one of the $\Delta 12$ desaturase sequences actually encodes a $\Delta 15$ desaturase, which would form α -linolenate. If, indeed, *Neurospora* does contain two $\Delta 12$ desaturase genes, this could account for the fact that known mutants are only partially blocked in the synthesis of polyunsaturated fatty acids (612). In *Neurospora*, longer-chain fatty acids are almost exclusively the 18- to 24-carbon 2-hydroxy fatty acids found in sphingolipids (1, 514); a sequence with homology to fatty acid elongases have also been identified (Table 40).

Lipid composition (both fatty acid composition and relative proportion of individual lipid classes) has many possible points of control. Membrane lipid composition is particularly important because it influences fluidity and related physical properties important to membrane function (563). Most acyl lipids can be synthesized through multiple routes, and modified by exchange of acyl groups (563). Degradative pathways are also important for control of lipid composition as well as for fatty acid utilization; many products of phospholipases and other lipases are also important signaling molecules (see "Calcium signaling" below). In addition to mitochondrial proteins probably involved in β -oxidation, *Neurospora* contains an inducible multifunctional peroxisomal β-oxidation protein (248) and many lipase candidates (see "Extracellular digestion" above). Most of the complex biochemical pathways involved in synthesis, remodeling, and turnover of membrane and storage lipids in Neurospora have not been well characterized. However, genes homologous to those encoding many enzymes involved in these processes in other organisms have been identified in the Neurospora genome database (Table 40). As expected, given the importance of acyl group transfer in these processes, the genome contains several candidate genes for acyltransferases, including glycerol-3-phosphate acyltransferase and lysophosphatidic acid acyltransferase, which carry out the two acylation steps leading to the synthesis of CDP-diacylglycerol and diacylglycerol (the precursors for the synthesis of phospholipids and triacylglycerols), genes representing two families of diacylglycerol acyltransferases, which form triacylglycerols, and genes for both subunits of serine palmitoyltransferase, which catalyzes the committed step for synthesis of sphingolipids.

The genome also includes candidate genes encoding a set of enzymes in the major pathway of phosphatidylcholine synthesis (two for phosphatidylserine synthase, three for phosphatidylserine decarboxylase [phosphatidylethanolamine synthase], and one each for phosphatidylethanolamine methyltransferase and phosphatidylmonomethylethanolamine methyltransferase). One of the phosphatidylserine synthase sequences includes a domain homologous to phospholipase D, and one of the phosphatidylserine decarboxylase sequences includes a domain homologous to protein kinase C. In addition, the genome includes candidate sequences for enzymes in the minor pathway of phosphatidylethanolamine and phosphatidylcholine synthesis via the CDP-alcohols (128, 181, 563). A similar pathway is used for synthesis of phosphatidylserine, phosphatidylinositol, and phosphatidylglycerol. A total of six CDPalcohol transferase gene homologues, the same number as in S. cerevisiae (128, 588), are identifiable. Although the CDP-

alcohol pathway is the major pathway in plants and animals (563), in *Neurospora* it is a minor pathway, except in mutants impaired in the methylation pathway such as the choline-requiring chol-1 and chol-2 mutants, which are blocked at the final steps leading to phosphatidylcholine formation (612). Another mutant, the inl mutant, requires inositol and is blocked in the formation of myo-inositol-l-phosphate, the rate-limiting step in the synthesis of both phosphatidylinositol and sphingolipids (612). All three mutants, the chol-1, chol-2, and inl mutants, have altered phospholipid composition and abnormal morphology at suboptimal levels of the growth supplement (612). In S. cerevisiae, regulation of the homologues of these genes and sec14, which encodes a phosphatidylcholine/phosphatidylinositol transfer protein, are key points for control of membrane lipid composition (128). Two genes in Neurospora have homology to sec14 (Table 40). Some enzymes and pathways in S. cerevisiae are specialized for cytoplasmic versus mitochondrial phospholipid synthesis (128, 181), consistent with a need for duplication of function in Neurospora as well. However, in Neurospora, three phosphatidylserine decarboxylase homologues are present, as in A. thaliana (60), while S. cerevisiae contains only two (128, 181). The reason for this is unclear, but one reason for apparent duplication of function may be a need for one enzyme that preferentially uses polyunsaturated fatty acids (822).

Over the past 15 years, the amenability of two model systems, *S. cerevisiae* and *A. thaliana*, to genetic and molecular studies has been exploited for systematic identification and characterization of genes involved in lipid metabolism, leading to a wealth of literature and a more complete understanding of the roles of lipids in these and other organisms. The diversity of *Neurospora* lipids and the well-studied effects of various developmental and environmental factors on their composition suggest an excellent opportunity for new insights from studies of this organism and productive future comparative genomics studies.

Protein Glycosylation, Secretion, and Endocytosis

N-linked protein glycosylation (dolichol) pathway. Most proteins present in the fungal cell wall are glycoproteins, and some of these are enzymes involved in structural modifications of the wall required for growth, morphogenesis, and nutrition. In addition, glycoproteins bear the most significant antigenic determinants of fungal cells (668). The biosynthesis of N-linked polysaccharides is rather complex and involves multiple steps. Thus, identification of the relevant components and determination of the overall degree of conservation of the pathway is an important advance in establishing the ability to structurally and functionally dissect the glycosylation process in *Neurospora*.

N glycosylation involves the assembly and transfer of oligosaccharide-P-P-dolichol, which is then processed and forms an N-glycosidic bond with one or more specific Asn residues of an acceptor protein emerging from the luminal surface of the membrane of the endoplasmic reticulum (ER). The commonality of glycoproteins throughout evolution suggests the presence of highly conserved components of the multistep process involved in glycoprotein biosynthesis. Hence, it is not surprising that the key elements of this pathway are present in *Neurospora*, even though some components of this pathway were previously not detected, most probably due to the technical shortcomings of heterologous probing (902).

Burda and Aebi (116) reviewed the dolichol pathway of N-linked glycosylation and suggested that N-linked glycosylation in eukaryotes and in archaea have a common evolutionary origin. The S. cerevisiae dolichol pathway was used as a basis for comparison while scanning the Neurospora genome for similar components. A very high degree of conservation was found in the machinery involved in the stepwise production of the core oligosaccharide Glc₃Man₉GlcNAc₂. However, the en bloc transfer of the core oligosccharide catalyzed by the ERresident enzyme N-oligosaccharyltransferase (OST) complex appears to be more streamlined in Neurospora (Fig. 8; Table 41). This is evident on the basis of the apparent absence of a good match to the S. cerevisiae Ost4p and Ost5p proteins in the Neurospora sequence database (homologues of these proteins are also not found in the *M. grisea* genome sequence [http: //www-genome.wi.mit.edu]). Interestingly, a similar degree and pattern of streamlining of this part of the pathway is also evident in the human OST complex. In S. pombe, there is an apparent homologue to Ost4p but not to Ost5p. Even though these differences may have evolutionary significance, it should be noted that Ost4p and Ost5p are very short proteins (36 and 86 amino acids, respectively), a feature that may impose limitations on the efficacy of the standard BLAST-based searches.

Apart from evolutionary implications, the structural similarity of the individual components, as well as the fact that the two major modules (oligosaccharide production and transfer complexes) are highly conserved, also has practical implications. For example, it is clear that much of the success in the harnessing of filamentous fungi for heterologous expression of proteins (383) has stemmed from the presence of proper posttranslational protein modification processes, including glycosylation.

Secretory and endocytic pathways. The high elongation rates and high secretory capacities observed in filamentous fungi pose special requirements on their endocytic/exocytotic pathways. Small GTPases of the Rab/Ypt and ARF branches of the Ras superfamily play central roles during secretory and endocytic trafficking. ARF subfamily proteins are involved in the formation of carrier vesicles from the donor organelle, while Rab proteins are required for vesicle targeting and fusion with the acceptor organelle (140, 187). Due to their high specificity for a given transport step, these proteins can be used as markers to monitor the different trafficking routes that are present in *Neurospora* and unicellular yeasts (Table 42) (304).

The S. cerevisiae secretory pathway is defined by the Rab proteins Ypt1p, Sec4p, and the paralogues Ypt31p/32p, while Ypt6p, Ypt7p, and the paralogues Ypt51p/52p/53p regulate the endocytic pathway (Table 42; Fig. 9). These proteins represent the minimal set of Rab-type GTPases present in all fungi examined. Deletion of any of the Rab genes of the secretory pathway is lethal in S. cerevisiae, C. albicans, and S. pombe, while mutations in the endocytic pathway result in viable S. cerevisiae and S. pombe mutants. In addition to this minimal set, Neurospora and other filamentous fungi possess Rab proteins that are not present in unicellular yeasts but do exist in mammals, implying that the more complex growth behavior also requires more complex organization of the ve-



Cytoplasm

FIG. 8. *Neurospora* dolichol pathway. The mechanism for sequential glycosylation of proteins in the ER is shown. *S. cerevisiae* homologues are presented in parentheses. GlcNAc, *N*-acetylglucosamine. Modified with permission from Markus Aebi (personal communication, 2003)

sicular transport system. Filamentous fungi are known for their high secretory capacity, and this may have required an expansion in the number of the secretion-related GTPases. In addition to an Ypt1p orthologue, *Neurospora* contains a Rab2related protein that is likely to function in the transport from endoplasmic reticulum to the Golgi apparatus. Surprisingly, in contrast to yeasts, deletion of *srgA* (the orthologue of Sec4p) was not lethal in *A. niger* (636). Also, *srgA* did not complement the *S. cerevisiae sec-4* defect, arguing for differences in the organization of the secretory pathway between filamentous fungi and yeasts that are not apparent at the genome level. The

A. niger srgA phenotype further suggested that two different secretory routes might exist: a major SrgA/Sec4 dependent route for the constitutive transport of cell wall proteins and a second inducible pathway that is less dependent on SrgA function.

Endocytosis provides a mechanism for plasma membrane proteins and lipids, and extracellular molecules, to be internalized by cells. It is generally regarded as an essential process of eukaryotic cells serving many functions, including recycling of membrane proteins and lipids, removal of membrane proteins and lipids for degradation, and the uptake of signal molecules. Endocytosis has been well characterized in *S. cerevisiae* (276)

TABLE 41. Neurospora oligosaccharyl transferase complex components

	NCU		BL	AST match		
Enzyme	NCU no.	Best overall	S. cerevisiae	S. pombe	Animal ^a	Plant ^b
Oligosaccharyltransferase subunit	02708.1	S. cerevisiae Stt3/0.00	0.00	4e-25	0.00; C. elegans	0.00
Dolichyl-diphosphooligosaccharide protein glycosyltransferase	02541.1	H. sapiens Rib1/3e-49	e-44	6e-45	3e-49; H. sapiens	2e-52
Dolichyl-diphosphooligosaccharide protein glycotransferase	00669.1	<i>S. pombe</i> 7e-55	9e-43	7e-55	3e-48; C. elegans	8e-46
Oligosaccharyl transferase delta subunit	03224.1	H. sapiens QPCT/2e-44	2e-47	2e-54	2e-44; H. sapiens	No match
Oligosaccharyl transferase gamma subunit	03995.1	R. norvegicus AIG2/6e-26	8e-23	No match	5e-25; H. sapiens	3e-7
Oligosaccharyl transferase epsilon subunit	09216.1	H. sapiens DAD1/5e-26	4e-12	2e-15	5e-26; H. sapiens	2e-13

^a Caenorhabditis elegans, Drosophila melanogaster, Mus musculus, or Homo sapiens.

^b Arabidopsis thaliana.

Neurospora ^b protein	Proposed transport role	S. cerevisiae	Mammals
Rab proteins			
Minimal set required in all fungi			
NCU08477.1	ER-GA, IGA, GA-ER	Ypt1p ^c	Rab1
NCU06404.1	LGA-PM	Sec4p ^c	Rab8
NCU01532.1	IGA, LGA–PM?	Ypt31/32p ^c	Rab11
NCU05234.1	IGA, LGA-LE, LE-LGA	Ypt6p ^c	Rab6
NCU03711.1	LE-V	$\dot{Ypt7p^{c}}$	Rab7
NCU00895.1, NCU06410.1	PM-EE, EE-LE	Ypt51/52/53p ^c	Rab5
Specific for S. cerevisiae or S. pombe			
1 1	?	Ypt10p	
	?	Ypt11p	
	EE-LE, LE-V	1 1	Rab7
Lost in <i>S. cerevisiae</i> , but present in <i>S. pombe</i> , filamentous fungi and animals			
NCU00889.1	EE-PM		Rab4
Present in filamentous fungi and animals, but not <i>S. cerevisiae</i>			
NCU01647.1	ER-GA, GA-ER		Rab2
NCU01453.1	PM-EE?, EE-PM?		Rab18?
NCU08271.1 (unusually long protein)	PM-EE, EE-LE		Rab5
ARE/ARE-like proteins			
Minimal set required in all fungi			
NCU08340 1	GA-FR IGA I GA PM	$Arf1/2n^{c}$	Arf1
NCU08989 1	LGA	$\operatorname{Arl1n}^{c}$	Arl1
NCU00333 1	LGA	$Arl3n^c$	Arl3
NCU00218 1	2	$Arf3n (=Arl2n)^c$	Arl2
NCU00381 1	FR-GA	Sar1n ^c	Sar1
NC000301.1	LK-0A	Sarp	5411
Present in filamentous fungi and animals, but not <i>S. cerevisiae</i>			
NCU07173.1	PM-EE		Arf6
NCU08618.1	?		Hypothetical protein

TABLE 42. Distribution and proposed role of transport-related small GTPases^a

^a Abbreviations: GA, Golgi apparatus; LGA, late Golgi apparatus; IGA, intra-Golgi transport; PM, plasma membrane; ER, endoplasmic reticulum; EE, early endosome; LE, late endosome; V, vacuole.

^b Orthologues are also present in *M. grisea* and *A. fumigatus*.

^c Orthologues are also present in S. pombe and C. albicans.

but is still considered controversial for filamentous fungi. Nevertheless, there is a growing body of evidence for endocytosis occurring in filamentous fungi, and particularly *Neurospora* (644). The *Neurospora* genome database was searched for homologues of 29 key proteins involved in endocytosis in *S. cerevisiae*. Each of these proteins was highly similar to one or more predicted proteins in the *Neurospora* genome (644). These



FIG. 9. Proteins of the secretory pathway. *Neurospora* predicted proteins homologous to those involved in the various steps of protein sorting to membranous organelles are shown. Abbreviations: EGA, early Golgi apparatus; LGA, late Golgi apparatus; PM, plasma membrane; V, vacuole; LE, late endosome, and EE, early endosome. NCU numbers for putative Rab proteins are indicated using boldface type.

NCU no	BI	AST match			
NCU IIO.	Best overall match in other organisms	S. cerevisiae	S. pombe	Animal ^a	Plant ^b
10073.1	7e-23 H. sapiens cervical SH3P7	2e-11	1e-16	7e-23	8e-04
07989.1	8e-54 A. thaliana clathrin coat assembly protein AP17	1e-34	2e-49	2e-44	8e-54
07171.1	1e-167 S. cerevisiae actin-like protein ACT2	1e-167	1e-166	1e-153	1e-129
01756.1	1e-158 S. pombe actin-like protein 3	1e-154	1e-158	1e-121	1e-124
02510.1	0.0 S. pombe clathrin heavy chain	0.0	0.0	0.0	0.0
04115.1	3e-24 S. pombe clathrin light chain	5e-09	3e-24	6e-06	4e-03
09808.1	0.0 S. cerevisiae Dnm1p	0.0	0.0	1e-157	1e-163
04100.1	0.0 A. nidulans VpsA	0.0	0.0	1e-149	1e-158
01255.1	1e-143 S. pombe dynamin-related protein	1e-137	1e-143	5e-47	1e-51
06347.1	1e-116 A. nidulans SAGA	1e-34	9e-63	8e-20	3e-06
04783.1	1e-32 H. sapiens ENTH domain of epsin	6e-31	2e-30	1e-32	7e-26
03298.1	1e-162 S. pombe synaptojanin-like protein	1e-146	1e-162	6e-77	4e-36
03792.1	4e-44 C. elegans synaptojanin UNC-26A	6e-39	2e-43	4e-44	2e-38
00896.1	1e-125 S. cerevisiae RSD1 (SAC1)	1e-125	1e-109	1e-90	2e-65
01330.1	3e-75 H. sapiens KIAA0966 protein	4e-51	2e-51	3e-75	2e-43
07438.1	2e-61 A. thaliana putative protein	3e-36	2e-34	9e-10	2e-61
02111.1	0.0 A. nidulans myosin 1 myoA	0.0	0.0	0.0	1e-135
06397.1	4e-29 S. pombe profilin	2e-22	4e-29	2e-14	7e-13
06171.1	7e-71 S. cerevisiae Pan1p	7e-71	3e-53	6e-16	3e-35
06777.1	2e-08 S. cerevisiae Pep12 p	2e-08	4e-05	2e-04	2e-07
06397.1	5e-29 S. pombe profilin	2e-22	5e-29	3e-14	No hits
01069.1	4e-96 S. pombe Hob3	6e-70	4e-96	2e-15	No hits
04637.1	2e-89 S. pombe RVS167 homologue	5e-83	2e-89	8e-10	8e-07
02978.1	3e-17 S. cerevisiae Sla1p	3e-17	9e-16	7e-06	6e-14
01956.1	0.0 Y. lipolytica SLA2-like protein	5e-75	4e-95	3e-42	9e-14
04119.1	4e-38 S. pombe t-SNARE complex subunit	3e-26	4e-38	2e-22	3e-19
06192.1	0.0 A. nidulans VpsB	1e-122	1e-148	1e-148	1e-117
06410.1	2e-61 S. cerevisiae Ypt51p	2e-61	6e-46	7e-51	2e-46
00895.1	5e-68 S. pombe Ypt5 protein	3e-52	5e-68	1e-61	2e-55
01523.1	2e-83 D. melanogaster Rab11	No match	1e-75	2e-83	1e-77

TABLE 43. Proteins involved in endocytosis in Neurospora

^a Bos taurus, Caenorhabditis elegans, Drosophila melanogaster, Homo sapiens, Mus musculus, or Rattus norvegieus.

^b Arabidopsis thaliana.

results strongly suggest that *Neurospora* possesses the complex protein machinery required to conduct endocytosis and that this machinery is well conserved in both budding yeast and filamentous fungi. This hypothesis is supported by experiments that have shown that the membrane-selective markers of endocytosis, FM4-64 and FM1-43, are internalized by *Neurospora* hyphae in an energy-dependent manner (239, 336, 643, 795). Sequences of the 29 *Neurospora* endocytic proteins were then used to search for homologues in budding and fission yeast,



FIG. 10. *Neurospora* endocytosis proteins. The percentage of *Neurospora* proteins with the greatest homology to proteins in *S. cerevisiae*, *S. pombe*, animals, and plants is shown.

animal, and *A. thaliana* genomes (Table 43). The highest homology was found to endocytic proteins in *S. pombe* (Table 43; Fig. 10). In a comparison with just animals and plants, 68% of the proteins showed higher homology to animal homologues, while 32% were better matches to plant homologues.

The analysis of the different coat complexes and adaptors required for the generation of vesicles from various membrane compartments revealed significant differences in the vesicular transport system in Neurospora and S. cerevisiae. All components that constitute the core COPI, COPII, and clathrin complexes are highly conserved in Neurospora, but, interestingly, the Neurospora genome encodes an additional pair of proteins corresponding to the large subunit of the AP-1 adaptor complex (NCU01992.1, γ -subunit; NCU04404.1, β -subunit) that are not found in S. cerevisiae but are present in mammals (89, 90, 420). The functional significance of this duplication is unknown. Clathrin-coated vesicles and pits, which are commonly visualized with the electron microscope in animal and plant cells, are often indicative of clathrin-mediated endocytosis. However, there is no ultrastructural evidence for their occurrence in fungal cells, including those of N. crassa (644). Nevertheless, both heavy- and light-chain clathrin and a clathrin coat assembly protein are present in Neurospora (Table 43). Clathrin-coated vesicles are also involved in other parts of the vesicle trafficking network of eukaryotic cells. Whether clathrin-mediated endocytosis occurs in N. crassa or other filamentous fungi is, as yet, unclear.

An indirect route through the late endosomal compartment and back to the Golgi apparatus functions to recycle plasma membrane components in yeast. The high growth rate of Neurospora places a much higher demand on the recycling of material toward the growing tip (643), and several proteins that function in this recycling step have been identified. Homologues of mammalian Rab4p and ARF6 are probably acting during the transport from early endosomes to the plasma membrane. Interestingly, ARF6 function is thought to be linked to Rac (which is absent from yeasts) in vertebrate cells (587). Sequence analysis of NCU001453.1 showed that this protein is distantly related to Rab18p proteins, which are involved in the recycling to the plasma membrane, but also has significant homology to Sec4p. Therefore, this protein may act in the late secretory pathway, perhaps in parallel with Sec4. Finally, NCU08271.1 and NCU08618.1 encode unusually long proteins with homology to Rab5p and a hypothetical but highly conserved ARF that is found in mammalian cells.

ENVIRONMENTAL SENSING

Filamentous fungi are able to grow in more diverse environments than yeasts and presumably are confronted with a wider variety of environmental stimuli. Therefore, it is expected that this group of organisms should possess an extensive array of sensing and signaling capabilities. Analysis of genes implicated in signal transduction and stress responses demonstrates that *Neurospora* possesses classes of sensing molecules not found in the two sequenced yeasts. The expansion of upstream signaling proteins is often coupled with a conserved core of downstream components, suggesting the presence of extensive networking and/or new signaling interactions not found in *S. cerevisiae* and *S. pombe*.

Major Signal Transduction Pathways

Two-component regulatory systems. Two-component signal transduction systems are minimally composed of two proteins: a sensor histidine kinase and a response regulator (sometimes called a receiver) (600). The histidine kinase is autophosphorylated in an ATP-dependent manner on a conserved histidine residue, and this autophosphorylation activity is regulated by an environmental signal sensed by the kinase. Stimuli include osmolarity, nutrient levels, oxygen levels, cellular redox status, and light. Phosphate transfer occurs from the histidine kinase to a conserved aspartate residue in the response regulator. This differential phosphorylation of the response regulator results in an altered cellular response such as activation of enzyme activity, modulation of transcription, or altered protein-protein interactions. Two-component signal transduction pathways are extensively used to mediate prokaryotic signaling events. In recent years it has been discovered that these signaling systems are also found in eukaryotes, including plants, yeasts, filamentous fungi, and slime molds (679, 758, 857). Their presence has not yet been demonstrated in mammals, and since some two-component systems are involved in virulence responses in fungi, they present attractive antifungal targets.

Histidine kinase (HK) and response regulator (RR) domains can be easily recognized in proteins by using sequence alignments (600). The HK domain contains several conserved sequence elements, including the H box, which is the site of autophosphorylation, and the N, G_1 , F, and G_2 boxes, which mediate binding of ATP, as shown by the recent structures of two bacterial HK domains (79, 80, 793). As with most signal transduction proteins, the HK and RR domains are modular; that is, they can be combined in various arrangements with each other and with other protein domains to yield the desired signaling protein.

A more complex version of the two-component pathway is the phosphorelay. In a phosphorelay system, the sensor is typically a "hybrid" HK, containing both HK and RR domains. In this pathway, there are two layers of His-to-Asp phosphotransfer. The sensor autophosphorylates in response to a signal, and the phosphate is subsequently transferred to the aspartate of its own response regulator domain. Transfer of the phosphate from the sensor aspartate to the histidine of a separate Hpt (histidine phosphotransfer domain) protein follows. The phosphate is then ultimately transferred to another aspartate in a separate response regulator, resulting in a His-Asp-His-Asp phosphorelay. It has been speculated that the additional phosphotransfer steps allow for more complex regulation of signaling events in a cell, since there are more opportunities to modulate phosphorylation activity. In addition, multiple HK inputs can be channeled into common downstream signaling proteins (Hpt and RR). This scenario is reminiscent of multiple receptor inputs into a single HK in bacterial chemotaxis.

Interestingly, although the hybrid kinases comprise only a small fraction of the sensor kinases found in prokaryotes, they are the major form found in eukaryotes. Experimental confirmation of the phosphorelay system has been obtained for some of these systems, but it is only inferred in others. The bestcharacterized example of a eukaryotic phosphorelay is the SLN1-YPD1-SSK1 system that regulates the HOG1 mitogenactivated protein kinase (MAPK) cascade in S. cerevisiae (624, 626). Sln1p is a transmembrane hybrid HK whose phosphorylation is regulated in response to changes in extracellular osmolarity, although the actual signal sensed remains unknown. Under conditions of low osmolarity, the Sln1p HK autophosphorylates on its histidine residue and transfers this phosphate to its RR domain. The phosphate is subsequently transferred to the histidine of the Hpt called Ypd1p. Finally, the phosphate is transferred to the aspartate of an RR domain of Ssk1p. It is the phosphorylated form of Ssk1p that is unable to activate the MAPKKKs Ssk2/22p. Ssk2/22p are a pair of redundant MAPKKKs that activate the MAPKK, Pbs2p, which ultimately regulates the Hog1p MAPK. Under conditions of increased osmolarity, Sln1p is not phosphorylated; hence, there is no phospho transfer to Ypd1p and Ssk1p. The unphosphorylated form of Ssk1p is able to activate Ssk2/22 and ultimately Hog1p, which then modulates transcription to allow glycerol synthesis in the cell. Sln1p also mediates a phosphorelay to a second response regulator in the cell, Skn7p. Skn7p regulation by Sln1p results in control of expression from of a lacZ reporter construct that contains an Mcm1p binding site in its promoter (473, 891). The physiological relevance of this has vet to be determined. However, the Skn7p response regulator is able to mediate responses to oxidative stress independent of its role in the phosphorelay with Sln1p (435, 637).



FIG. 11. Domain organization of *Neurospora* HKs. Abbreviations: An, *A. nidulans;* Af, *A. fumigatus;* Ca, *C. albicans;* Sc, *S. cerevisiae;* Gc, *Glomerella cingulata.* Note that assignment of the total number of PAS/PAC domains in the relevant proteins is somewhat subjective, since it depends on the threshold values used during BLAST analyses.

The genome sequence of *Neurospora* reveals that it has 11 putative HK containing proteins, all of them hybrid kinases (Fig. 11). Only one of these (NCU04615.1) is predicted to span the membrane and appears to be most similar to Sln1p from S. cerevisiae and TcsB from A. nidulans. Two of the HKs have been previously characterized. Nik-1/Os-1 (NCU02815.1) has been shown to play a role in hyphal development (13, 695). Deletion mutants exhibit multiple morphological defects including aberrant hyphal development and reduced conidiation. These defects become exacerbated when cells are grown in media of high osmolarity; this may be the result of a defective cell wall structure. Nik-1/Os-1 has a homologue, Cos-1, in the opportunistic pathogen C. albicans, where it plays a role in hyphal development under various conditions on solid surfaces (14). Cos-1 also plays a role in virulence (704). Nik-2 (NCU01833.1) is a hybrid HK that also contains a PAS domain, which suggests that it may be involved in redox or light sensing (783) (see also "Photobiology and circadian rhythms" below). However, nik-2 deletion mutants exhibit no obvious phenotype when grown under a variety of conditions (L. Alex and M. Simon, unpublished results).

Interestingly, two of the HK proteins appear to be light sensors. NCU05790.1 and NCU04834.1 contain domains com-

mon to plant phytochromes and bacteriophytochromes, respectively (see "Photobiology and circadian rhythms" below). Deletion of these genes and their respective phenotypes is currently under investigation. Although there is no evidence that *Neurospora* growth and development are sensitive to red light, the presence of these two genes suggests that this may need to be readdressed.

There is also an HK that is similar to Fos-1 from *A. fumigatus* and TcsA from *A. nidulans*. These HKs play a role in conidiation in their respective species, but the phenotypes are slightly different (627, 834). The remaining 7 of the 11 *Neurospora* HKs have some similarity to other known HK sequences; however, the function of these HKs remains to be elucidated.

A comparison of HK numbers among a variety of fungi shows that the filamentous fungi have a larger complement of these proteins than any of the sequenced yeasts or *C. albicans*. *S. cerevisiae* has only 1 HK, Sln1p; *S. pombe* has 3 HKs, Mak1, Mak2, and Mak3; *C. albicans* has 3 HKs, Chk1, CaNik1/Cos-1; and CaSln1; and *Neurospora* has 11 HKs (269). Sln1p homologues appear in *S. cerevisiae*, *C. albicans*, and *Neurospora* but are absent from *S. pombe*. The three HKs from *S. pombe* do not have good homologues in *Neurospora*, *S. cerevisiae*, or *C. albicans*.

The downstream components of two-component pathways are also present in *Neurospora*. There is one Hpt protein (NCU01489.1) that is most similar to *S. pombe* Spy1/*S. cerevisiae* Ypd1p. Two response regulators are present; NCU01895.1 and NCU02413.1. These are most similar to *S. pombe* Mcs4 (*S. cerevisiae* Ssk1p) and *S. pombe* Prr1 (*S. cerevisiae* Skn7p), respectively. The conservation of a few downstream signaling elements, i.e., the Hpt and RR proteins, suggests that these proteins may act to integrate multiple signaling inputs from the many HKs to evoke the proper cellular response. Alternatively, some of the HKs may not act through phosphorelays. These various possibilities await testing.

Heterotrimeric G proteins. Seven-transmembrane helix, Gprotein-coupled receptors (GPCRs) are used to detect diverse environmental stimuli in eukaryotes (198). GPCRs regulate the activity of second messengers through their interaction with heterotrimeric G proteins (562). A heterotrimeric G protein consists of a G α subunit, which binds and hydrolyzes GTP, and a tightly associated G β and G γ subunit. Ligand binding to the GPCR leads to exchange of bound GDP for GTP on the G α subunit and to dissociation of the heterotrimer into G α -GTP and G $\beta\gamma$ units. G α -GTP and G $\beta\gamma$ can both interact with effectors to generate changes in cellular physiology and development.

Prior to the completion of the genome sequence, the total number of G-protein subunits in a filamentous fungal species was not known. This analysis (Table 44). showed that *Neurospora* possesses more heterotrimeric G α subunits than do budding and fission yeasts (three versus two) but the same number of G β and G γ genes (one each) (Fig. 12). Previous work has demonstrated that these subunits are important for hyphal growth, conidiation, female fertility, and stress responses in *Neurospora* (37, 369, 406, 873).

It has recently been shown that protein "mimics" for G β (Gpb1p and Gpb2p) and G γ (Gpg1p) subunits in *S. cerevisiae* are coupled to the Gpa2p G α protein and regulate filamentous growth in this organism (314). The two G β mimics each contain seven Kelch repeats and may assume a tertiary structure similar to the β -propeller of G β proteins. However, analysis of the *Neurospora* genome sequence did not reveal any proteins with seven Kelch repeats or with high similarity to either the G β or G γ mimics.

Analysis of the genome sequence shows that Neurospora possesses at least 10 predicted seven-transmembrane helix proteins that are potential GPCRs (Fig. 12). These proteins fall into five families: microbial opsins, pheromone receptors, glucose sensors, nitrogen sensors, and a novel class, not previously identified in fungi. The first class consists of an opsin (nop-1) and an opsin-related protein (orp-1) that were identified during prior EST projects (78); this family has three members in S. cerevisiae (Hsp30p, Yro2p and Mrh1p) (78, 297, 592, 861). In the second family are two predicted pheromone receptors, similar to S. cerevisiae α -factor and a-factor pheromone receptors (reviewed in reference 199). The expression patterns of the putative Neurospora pheromone receptor genes have been published (622). The third class of GPCRs is represented by a protein similar to putative glucose sensors in both S. cerevisiae and S. pombe (847, 866), while the fourth group consists of two proteins similar to the Stm1 protein from S. pombe, implicated in nitrogen sensing (149).

The fifth class contains three related GPCR-like genes. This group is not present in the genomes of *S. cerevisiae* or *S. pombe*. The encoded proteins are most similar to known and predicted GPCRs from the protists *Dictyostelium discoideum* (423) and *Polysphodylium pallidum* (TasA) and also to predicted proteins in *C. elegans* and *A. thaliana* (388); similarity is highest in the predicted seven transmembrane helices. Functions for this group of proteins have been elucidated only in *D. discoideum*, where four related cAR receptors sense cyclic AMP (cAMP) levels during multicellular development and sporulation (reviewed in reference 33).

In both fungi and higher organisms, phosducin and Regulator of G-Protein Signaling (RGS) proteins regulate G-protein signaling pathways by modulating the activity of $G\beta\gamma$ dimers and the GTPase activity of $G\alpha$ subunits, respectively (402, 734). *S. cerevisiae* possesses two phosducin and two RGS protein genes, while *Neurospora* contains two phosducin genes and one RGS genes (Fig. 12).

Ras-like GTPases. The superfamily of monomeric small GTPases (20 to 40 kDa) function as molecular switches that control a wide variety of cellular processes including signal transduction, cell polarity, the cytoskeleton, and the identity and dynamics of membranous compartments (reviewed in references 654 and 773).

A total of five Ras/Ras-related small GTPases are present in the *Neurospora* genome (Table 44). *Neurospora* is similar to *S. cerevisiae* in that it has two Ras proteins (Fig. 12), in contrast to the one in *S. pombe*. The two *S. cerevisiae* Ras proteins regulate adenylyl cyclase activity (791), while the sole *S. pombe* Ras, Ras1, regulates the activity of a Cdc42 guanine exchange factor to control morphogenesis, as well as a MAPK cascade that regulates mating in this organism (595). The *Neurospora* RAS-1 (18) and RAS-2/SMCO-7 (397) proteins have already been defined. Mutation of *ras-2/smco-7* leads to severe defects in hyphal growth and conidiation (397). The function of *Neurospora* RAS-1 is unknown.

The three *Neurospora* Ras-related proteins include one that is most similar to Rheb, which is evolutionarily conserved and controls arginine uptake in *S. cerevisiae* and *S. pombe* (reviewed in reference 649). The second, the previously identified KREV-1 protein (367), is similar to mammalian Rap (which can antagonize Ras signaling [reviewed in reference 649]) and *S. cerevisiae* Rsr1p (which controls bud site selection and cell polarity [598]). *Neurospora krev-1* null mutants are phenotypically indistinguishable from the wild type; however, analysis of mutationally activated alleles suggests involvement of *krev-1* in the sexual cycle (367). The third Ras-related *Neurospora* protein was not characterized previously but is most similar to a predicted Ras protein from the yeast *Yarrowia lipolytica*.

cAMP signaling. As in the two sequenced yeasts, *Neurospora* contains one gene encoding adenylyl cyclase, cr-1 (427), one cyclase-associated protein gene, cap, and one protein kinase A (PKA) regulatory subunit gene (*mcb*) (Table 44; Fig. 12) (113). In contrast, the genome sequence predicts two distinct PKA-cat subunits in *Neurospora*. One is most similar to PkaC from *Collectorichum trifolii* but is also related to the three *S. cerevisiae* Tpk PKA-cat genes (790). Mutation of the *Neurospora* gene causes a phenotype similar to that of the adenylyl cyclase mutant cr-1 (M. Plamann, unpublished data). The second protein is most closely related to Pka-cat from *M. grisea* and is also

					1		
Ductories along	Neurospora crassa	NCU		BL	AST match		
	protein	no.	Best overall	S. cerevisiae	S. pombe	Animal ^a	Plant ^b
Heterotrimeric Ga	GNA-1	06493.1	Cryphonectria parasitica Cpg-1; 0.00	Gpa2p; 1e-71	Gpa1; 5e-70	Gαi-3; e-102	GPA1; 1e-48
	GNA-2	06729.1	Magnaporthe grisea MAGC; 1e-137	Gpa2p; 3e-46	Gpa1; 1e-59	G-α65A; 3e-63	GPA1; 2e-34
	GNA-3	05206.1	M. grisea MAGA; 1e-178	Gpa2p; e-102	Gpa2; 4e-85	G-αi-3; e-102	GPA1; 1e-48
Heterotrimeric G _β	GNB-1	00440.1	M. grisea GB; 1e-164	Ste4p; 5e-51	Git5; 1e-60	GPB-1; e-108	GB; 2e-73
Heterotrimeric Gy	GNG-1	00041.1	Botrytis cinerea CNS01BJB ($G\gamma$); 9e-20	None	None	None	None
Pheromone receptor (GPCR)	PRE-1	00138.1	Sordaria macrospora Pre-1; 1e-138	Step3; 0.063	None	None	None
e e	PRE-2	05758.1	S. macrospora Pre2; 0.00	None	None	None	None
Putative cAMP GPCR	GPR-1	00786.1	Dictyostelium discoideum CrlA; 3e-15	None	None	None	GPCR; 0.002
	GPR-2	04626.1	Arabidopsis thaliana GPCR CAA72145.1;	None	None	None	GPCR; CAA72145.1; 2e-09
	GPR-3	09427 1	D discoilerum AA062367· 3e-07	None	None	None	GPCR CAA72145 1.
							1e-04
Putative glucose sensor GPCR	GPR-4	06312.1	S. cerevisiae Gpr1p; 3e-14	Gpr1p; 3e-14	None	None	None
Stm1-like GPCR	GPR-5	00300.1	S. cerevisiae YOL092w; 1e-29	YOL092w; 1e-29	Stm1; 1e-18	NP 493686.1; 3e-13	AT4g36850; 2e-14
	GPR-6	09195.1	S. cerevisiae YBR147w; 4e-32	YBR147w; 4e-32	Stm1; 8e-14	Y43H11AL2; 2e-14	NP_568009.1; 9e-10
Microbial opsin	NOP-1	10055.1	Leptosphaeria maculans opsin; 2e-75	Yro2p; 1e-08	SPCC31H12-02e; 9e-07	None	None
Opsin-related protein	ORP-1	01735.1	Coriolus versicolor Hsp30; 5e-31	Yro2p; 8e-23	SPCC31H12.02c; 8e-12	None	None
Phosducin		00441.1	C. parasitica Bdm-1; 2e-57	None	None	None	None
		00617.1	S. cerevisiae Plp2p; 2e-42	Plp2p; 2e-42	SPBC2A9; 1e-34	IAP-associated factor	F18O22.30; 9e-12
						VIAF1; 8e-17	
Regulator of G-protein signaling		08319.1	A. nidulans FlbA; e-139	Sst2p; 1e-14	C22F3.12c; 9e-27	RGS16; 9e-05	None
Adenylyl cyclase	CR-1	08377.1	M. grisea MAC1; 0.00	Cyr1p; 3e-160	Adenylyl cyclase; e-111	Scribbled; 1e-27	F3M18.12; 1e-21
Cyclase-associated protein		08008.1	Candida albicans CAP; 2e-67	Srv2p/CAP; 2e-65	CAP; 1e-66	CAP2; 1e-59	Cap1; 1e-36
PKA regulatory subunit	MCB	01166.1	Colletotricum lagenarium Rpk-1; e-142	Bcy1p; 2e-67	Cgs1; 2e-58	PKA regulatory, type II α ;	None
PKA catalytic subunit		06240.1	Colletotricum trifolii pkaC; e-164	Tpk1p; e-129	KapB; e-110	3e-55 PKA C-beta; 4e-93	ATPK19; 2e-58
•		00682.1	M. grisea PKA-ct; e-142	Tpk1p; 5e-74	KapB; 1e-67	PKX1; 1e-81	ATPK19; 2e-58
High-affinity cyclic nucleotide		00478.1	C. albicans high-affinity cAMP phospho-	Pde2p; 4e-23	None	XP319716.1; 3e-25	None
phosphodiesterase			diesterase; 2e-45				:
Low-attinity cyclic nucleotide phosphodiesterase		00237.1	S. pombe Pde1; 3e-16	Pde1p; 4e-15	Pde1; 3e-16	None	None
Ras	RAS-1 PAS-2/SMCO-7	08823.1	Sclerotinia sclerotiorum hypothetical Ras; 3e-63	Ras1p; 1e-24 Ras2n: 7e-38	Ras1; 1e-26 C A A 7 7 300 1· 4e-30	rRas2; 9e-28 BAB7790 1: 7e-47	BAB78669.1; 1e-06 Bic1: 9e-19
Ras-related	KREV-1	02167.1	H. sapiens Raichu404X; 6e-49	Rsr1p; 3e-47	CAA27399.1; 5e-33	Raichu404X; 6e-49	Ran1; 7e-18

TABLE 44. Heterotrimeric G-protein signaling, cAMP metabolism, and Ras proteins

^a Caenorhabditis elegans, Drosophila melanogaster, Anopheles gambiae, Mus musculus, or Homo sapiens. ^b Arabidopsis thaliana or Oryza sativa.

MICROBIOL. MOL. BIOL. REV.



FIG. 12. Known and predicted heterotrimeric G-protein, Ras, cAMP, and PAK/MAPK signaling pathways in *Neurospora*. The number of images for each signaling protein class in the cartoon represents the number of *Neurospora* predicted gene products in each group. Arrows depict interactions that are supported by evidence from other systems but have not yet been demonstrated in *Neurospora*. The MAPK cascade(s) may also receive input from two-component signaling pathways (Fig. 11). Various Rho GTPase superfamily members (Fig. 18) may regulate certain signaling events downstream of Ras and upstream of PAK proteins. Abbreviations: GPCR, G-protein-coupled receptor; RGS, regulator of G-protein signaling; PAK, p21-activated kinase; GCK, germinal-center kinase; AC, adenylyl cyclase; CAP, cyclase-associated protein; PKA-R, regulatory subunit of cAMP-dependent protein kinase; PKA-C, catalytic subunit of cAMP-dependent protein kinase; MAPKK, and MAPKKK, mitogen-activated protein kinase, kinase kinase, and kinase kinase, respectively.

similar to Uka1 from *Ustilago maydis* (209) but is less similar to *S. cerevisiae* PKA-cat subunits than the first. Two predicted cAMP phosphodiesterase genes are most similar to the high-and low-affinity forms, respectively, found in *S. cerevisiae* (491, 682).

cAMP accumulates in the extracellular medium of wild-type *Neurospora* cultures (371, 724), and it is has recently been shown that loss of the heterotrimeric G α gene, *gna-1*, blocks the ability of strains lacking a functional adenylyl cyclase or *gna-3* G α gene to respond to cAMP supplementation (370, 405). These data, coupled with the presence of GPCRs similar to slime mold cAMP receptors, suggest that cAMP or a related molecule may serve as an environmental signal and GPCR ligand in *Neurospora*. The existence of such a pathway has not previously been demonstrated in any fungal system.

PAKs and GCKs. (i) PAKs. p21-activated kinases (PAKs) are serine/threonine protein kinases that can be activated by binding to GTP-bound Rho-like GTPases of the Cdc42/Rac family. PAKs have been found in most eukaryotes, where they

regulate a wide variety of processes including modulation of MAPK pathways, cytoskeletal dynamics, cell cycle progression, and apoptosis (92). PAKs have a common arrangement of protein domains, a CRIB domain at the N terminus and the serine/threonine kinase domain at the C terminus. The CRIB domain is able to bind to the kinase domain and inhibit its activity. On binding of a GTP-bound Rho-like GTPase to the CRIB domain, inhibition of kinase activity is relieved (reviewed in references 92 and 177).

The most well-characterized member of the PAK family in fungi is Ste20p from *S. cerevisiae*. Ste20p activity is important for the response to pheromone, osmostress, filamentation, and polarized growth. Of these, the best understood is the pheromone response. On pheromone binding to its GPCR, the heterotrimeric G-protein is activated. This, in turn, results in recruitment of the Ste20p-Cdc42p complex to the mating MAPK module and allows phosphorylation of the MAPKKK Ste11p. *S. cerevisiae* also has a second PAK, Cla4p, that is involved in budding and cytokinesis (66, 176). In addition to

			-	-	
NCU	Kinase			Best BLAST match ^a	
no.	type	S. cerevisiae	Animal	Other fungi	Plant
03894.1 00406.1	РАК	Ste20p ^b ; 1e-109 Cla4p; 9e-99	Pak1 (Hs); 1e-108 Pak1 (Hs); 2e-92	Um; 1e-107 CHM1 ^b (Mg); 1e-153	Putative S/T kinase (At); 1e-55 Putative S/T kinase (Hv); 1e-47
00772.1 04096.1	GCK	Sps1p; 5e-65 Sps1p; 1e-67	Severin kinase ^b (Dd); 1e-109 Severin kinase ^b (Dd); 1e-102	Ste20-like kinase (Um); 6e-81 Ste20-like kinase (Um); 1e-101	Putative S/T kinase (Os); 9e-74 Putative protein kinase (At); 5e-82

TABLE 45. p21-Activated and germinal center kinases

^a Hs, Homo sapiens; Um, Ustilago maydis; Mg, Magnaportha grisea; At, Arabidopsis thaliana; Hv, Hordeum vulgare; Os, Oryza sativa; Dd, Dictyostelium discoideum. ^b Best overall hit.

the CRIB and kinase domains present in all PAKs, Cla4p also has a PH (plekstrin homology) domain adjacent to the CRIB domain. PH domains have been implicated in protein-protein interactions, binding to phosphatidylinositol-4,5-bisphosphate, and membrane association (467).

The genome sequence of *Neurospora* indicates that there are two PAKs, one a Ste20p homolog and one a Cla4p homolog (Table 45; Fig. 12). Many of the components that are known to interact with Ste20p and Cla4p in *S. cerevisiae* are present in *Neurospora* (Ste50p, Cdc42p, and Ste11p). Therefore, it can be expected that these two PAKs will control similar responses in *Neurospora* but that there will definitely be differences, given the difference in growth processes for these two organisms. For example, NcCla4p may play a role in conidiophore production by analogy to budding yeast. The physical characterization of the PAKs and their functions awaits further investigation.

(ii) GCKs. A second group of kinases that are similar to PAKs are the germinal center kinases (GCKs). The domains and their organization in GCKs differ from those in PAKs, in that the serine/threonine kinase domain is present at the N terminus of the protein and this is followed by a sequence that is not conserved with PAKs or between many GCKs themselves (444, 719). There are no CRIB or PH domains present. It is thought that the C-terminal sequences act as autoinhibitory to the kinase activity and that binding of other components to the C-terminal sequences probably relieves this inhibition. GCKs are responsible for mediating some stress responses (reviewed in reference 444). In yeast, there appears to be two GCK-like proteins, Sps1p and Kic1p. Sps1p is a developmentally regulated kinase that is necessary for later events in sporulation (255). Kic1p is a kinase whose activity is dependent on the yeast centrin-like protein Cdc31p and is necessary for cell wall integrity (764). *Neurospora* has two predicted GCK-like proteins (Table 45; Fig. 12) that show the highest homology to severin kinase from *D. discoideum*. The *D. discoideum* severin kinase phosphorylates the actin binding protein severin, leading to rapid rearrangement of the actin cytoskeleton during amoeboid movement (215). The role of these two GCK-like proteins in *Neurospora* is currently unknown, but the roles of similar proteins in other eukaryotic microbes suggest involvement in cytoskeletal dynamics and/or ascospore development.

MAPKs. MAPK pathways consist of three serine/threonine protein kinases (MAPKKK, MAPKK, and MAPK) that act sequentially, culminating in phosphorylation of target proteins that regulate transcription, the cell cycle, or other cellular processes (138, 604, 625). MAPK modules are regulated by a wide variety of signaling proteins in other fungi, including GPCRs, PAKs, histidine phosphorelays, and Cdc42p (138, 604, 625).

A total of nine MAPKKK/MAPKK/MAPK protein-encoding genes were found in the *Neurospora* genome sequence (Table 46; Fig. 12). The similarity scores demonstrate that these nine proteins could form three pathways corresponding to those for pheromone response/filamentation, osmosensing/

TABLE	46.	MAPKs
-------	-----	-------

Ductoin along	Neurospora	NCU			BLAST match		
Protein class	protein	no.	Best overall	S. cerevisiae	S. pombe	Animal ^a	Plant ^b
MAPKKK (osmo- sensing/stress?)		03071.1	S. pombe Win1/Wak1/Wis4; 0.00	Ssk2p; 1e-128	Win1/Wak1/Wis4; 0.00	PRO0412; 9e-54	NPK1-related protein kinase3; 2e-49
МАРКК		00587.1	S. cerevisiae Pbs2p; 1e-95	Pbs2p; 1e-95	Wis1/Sty2; 6e-92	Drosophila Dsor1; 2e-60	MKK2; 6e-44
MAPK	OS-2	07024.1	<i>M. grisea</i> osmotic stress MAPK; 0.00	Hog1p; 1e-165	Sty1; 1e-170	EAA00104.1; 1e-99	MPK3; 2e-91
MAPKKK (phero- mone response?)	NRC-1	06182.1	A. nidulans MAPKKK; 1e-179	Ste11p; 1e-82	NP595714.1; 5e-84	CAD38973.1; 1e-58	NPK1-related protein kinase2; 1e-69
MAPKK		04612.1	Glomerella cingulata EMK1; e-165	Ste7p; 6e-66	Byr1; 4e-80	EAA01212.2; 1e-74	MKK6; 2e-50
MAPK	MAK-2	02393.1	Podospora anserina CAD60723.1; 0.00	Fus3p; 2e-120	Spk1; 1e-133	CAA77753.1; e-108	MPK4; 3e-95
MAPKKK (cell integrity?)		02234.1	P. anserina AAL77223.1; 0.00	Bck1p; 4e-80	Mkh1; 1e-100	MEKK 3; 2e-49	NPK1-related protein kinase3; 3e-59
MAPKK		06419.1	M. grisea Mkk1; 1e-176	Mkk1p; 3e-85	Skh1; 4e-79	EAA01212.2; 3e-50	MKK2; 8e-43
MAPK		09842.1	C. lagenarium AAL50116.1; 0.00	Slt2p; 1e-136	Spm1; 1e-140	BMK1 kinase; 6e-85	MPK6; 9e-90

^a Caenorhabditis elegans, Drosophila melanogaster, Anopheles gambiae, Mus musculus, or Homo sapiens.

^b Arabidopsis thaliana or Oryza sativa.



FIG. 13. Calcium signaling proteins in *Neurospora*. The numbers of each gene in a particular class are in parentheses. An asterisk indicates that the location in the plasma membrane and/or intracellular calcium store membrane has not been determined. CPC, Ca^{2+} -permeable channel; CT, Ca^{2+} -transporter (Ca^{2+} -ATPases, cation-ATPases, Ca^{2+}/H^+ exchangers, and Ca^{2+}/Na^+ exchangers); CaM, calmodulin; reg, regulated.

stress, and cell integrity pathways in S. pombe and S. cerevisiae (reviewed in reference 625). Two of the MAPKs and one MAPKKK had been previously identified in Neurospora. The MAK-2 MAPK is related to the class of proteins represented by S. cerevisiae Fus3p and Kss1p, required for the pheromone response and filamentous/invasive growth pathways, respectively (494). Mutations in *mak-2* result in inappropriate conidiation, female sterility, and loss of hyphal fusion (D. J. Ebbole and N. L. Glass, unpublished data). The OS-2 MAPK, related to the MAPK Hog1p from S. cerevisiae, regulates osmosensitivity and resistance to phenylpyrrole fungicides in Neurospora (897). NRC-1 is a MAPKKK that is most similar to Pneumocystis carinii mekk and is also related to S. cerevisiae Ste11p, which participates in multiple MAPK pathways. Similar to mak-2 mutants, nrc-1 mutants show inappropriate conidiation, female sterility, and inability to undergo hyphal fusion (431); N. L. Glass, unpublished data). The *Neurospora* genome also contains a protein similar to Sho1p, which operates upstream of the osmosensing MAPK pathway in S. cerevisiae (reviewed in reference 625).

The third *Neurospora* MAPK, similar to *Magnaporthe grisea* Mps1 (864) and *S. cerevisiae* Slt2p/Mpk1p (462), had not been identified prior to completion of the genome sequence. The existence of this sequence argues for the presence of a cell integrity-type MAPK pathway in *Neurospora*.

In contrast to *S. cerevisiae* and *S. pombe, Neurospora* does not possess additional MAPK components outside the three conserved modules found in all three species. This finding, coupled with the diversification of GPCRs and histidine kinases, suggests greater integration of signals from multiple upstream sensory proteins in *Neurospora* than in yeasts. Alternatively, these nine basic parts may be used to create multiple MAPK modules, leading to even more signaling diversity and cross talk than observed in yeasts. The latter hypothesis is consistent with the observation that loss of proteins similar to those involved in yeast pheromone response/filamentation (MAK-2 and NRC-1) leads to multiple phenotypic defects in *Neurospora*.

Calcium signaling. Calcium plays a central role as an intracellular signal in eukaryotic cells (70, 676), yet little is known about Ca²⁺ signaling in filamentous fungi compared to animals and plants. Much evidence, particularly from pharmacological studies, has indicated that Ca²⁺ signaling is involved in regulating numerous processes in filamentous fungi, including secretion, cytoskeletal organization, hyphal tip growth, hyphal branching, sporulation, infection structure differentiation, and circadian clocks (267, 723). However, information about the main proteins involved in any one Ca2+-mediated signal response pathway is lacking for Neurospora or, indeed, any other filamentous fungus. Initial analysis of the Neurospora genome indicated that more than 25 proteins are likely to be involved in Ca^{2+} signaling (269). This has now been updated with the identification of 22 additional Ca2+-signaling proteins in Neurospora (Table 47) (http://fungalcell.org/fdf/). These results indicate that Neurospora possesses a complex Ca²⁺-signaling machinery and that Ca^{2+} signaling is a significant component of its signal transduction network (Fig. 13).

Significantly, more Ca²⁺-signaling proteins are present in *Neurospora* than in *S. cerevisiae* (Table 47) (http://fungalcell.org/fdf; M. Bencina, B. J. Bowman, O. Yarden, and N. D. Read, unpublished data). Three new Ca²⁺ channel proteins, which have close homologues to the three Ca²⁺ channel proteins (Mid1p, Cch1p, and Yvc1p [238, 545, 583]) in *S. cerevisiae*, have been identified in *Neurospora*. Eight P-type Ca²⁺-ATPases have been identified in *Neurospora*, of which four are new (64). All of these have close homologues to the four P-type Ca²⁺-ATPases in *S. cerevisiae* (135). Another P-type cation-ATPase has also been identified in *Neurospora*. It is homologous to an *S. cerevisiae* hypothetical ORF of unknown function. Six recognizable Ca²⁺/H⁺ exchangers are present in

NOU	NT		BLAST m	atch			
NCU no.	Name	Type of protein	Best overall	S. cerevisiae	S. pombe	Animal	Plant
02762.1		Ca ²⁺ -permeable channel	0; Aspergillus nidulans (AAL37946)	0.0	0.0	0.0	None
06703.1		Ca ²⁺ -permeable channel	A. nidulans (Mid1)	9e-27	1e-48	None	None
07605.1		Ca ²⁺ -permeable channel	6e-77; Saccharomyces cerevisiae (Yvc1p)	6e-77	None	None	None
03305.1	NCA1	Ca ²⁺ -ATPase	0; Caenorabditis elegans (SCA-1)	None	None	0.0	0.0
04736.1	NCA2	Ca ²⁺ -ATPase	0; Schizosaccharomyces pombe (NP 593890)	0.0	0.0	e-120	e-155
05154.1	NCA3	Ca ²⁺ -ATPase	0; S. pombe (SPAPB2B4)	0.0	0.0	e-170	e-165
03292.1	PMR1	Ca ²⁺ -ATPase	0; Aspergillus niger (PmrA)	0.0	0.0	0.0	e-105
08147.1	PH-7	Ca ²⁺ -ATPase	0; S. pombe (SPBC839)	0.0	0.0	3e-98	1e-94
04898.1		Ca ²⁺ -ATPase	0; S. cerevisiae (SCE9871)	0.0	0.0	0.0	0.0
03818.1		Ca ²⁺ -ATPase	0; S. cerevisiae (Neo1p)	0.0	0.0	0.0	0.0
07966.1		Cation-ATPase	0; S. pombe (Cta3p)	0.0	0.0	e-100	e-102
01437.1		Cation-ATPase	0; S. pombe (NP 587882)	0.0	0.0	0.0	9e-77
07075.1	CAX	Ca^{2+}/H^+ exchanger	1e-80; S. cerevisiae (Mnr1p)	1e-80	2e-73	None	2e-58
00916.1		Ca ²⁺ /H ⁺ exchanger	1e-43; S. cerevisiae (Vex1p)	1e-43	8e-41	None	1e-32
00795.1		Ca ²⁺ /H ⁺ exchanger	3e-43; S. cerevisiae (Vex1p)	3e-43	1e-38	None	4e-30
06366.1		Ca ²⁺ /H ⁺ exchanger	1e-43; S. cerevisiae (Mnrp)	1e-43	7e-41	None	6e-43
07711.1		Ca ²⁺ /H ⁺ exchanger	2e-64; S. cerevisiae (Hum1p)	2e-64	2e-36	None	3e-50
05360.1		Ca^{2+}/H^+ exchanger	e-128; S. cerevisiae (Rpd3p)	e-128	4e-80	None	2e-11
02826.1		Ca ²⁺ /Na ⁺ exchanger	2e-23; S. pombe (SPAC3A12)	9e-08	2e-23	2e-16	3e-12
08490.1		Ca ²⁺ /Na ⁺ exchanger	6e-07; Anopheles gambiae (EAA01911)	None	None	6e-05	None
01266.1		Phospholipase C	0; Magnaporthe grisea (MPLC1)	5e-87	1e-61	1e-64	5e-28
06245.1		Phospholipase C	6e-53; Botryotinia fuckeliana (BCPLC1)	1e-47	1e-47	1e-43	2e-23
09655.1		Phospholipase C	2e-38; S. cerevisiae (YSCL9606)	2e-38	3e-24	6e-26	3e-22
02175.1		Phospholipase C	1e-30; S. cerevisiae (Plc1p)	1e-30	1e-30	7e-28	2e-13
04120.1	CaM^a	Calmodulin	3e-52; synthetic construct (CAD79597)	4.8e-36	1.1e-45	8e-47	8e-57
05225.1		Ca ²⁺ and/or CaM binding protein	2e-90; Solanum tuberosum (CAB52797)	1e-79	1e-48	None	2e-90
02115.1		Ca ²⁺ and/or CaM binding protein	2e-7; Homo sapiens (HERC2)	None	3.2e-05	2e-7	7e-5
01564.1		Ca ²⁺ and/or CaM binding protein	8e-73; S. cerevisiae (YNL083W)	8e-73	2.5e-55	9e-38	3e-45
03804.1	CNA-1	Calcineurin catalytic subunit	0; A. nidulans (cna ⁺)	e-165	0.0	0.0	0.0
03833.1	CNB-1	Calcineurin regulatory subunit	1e-73; B. cinerea (CNS018VZ)	4e-49	7.9e-52	3e-47	5e-18
09265.1		Calnexin	e-168; A. niger (ClxA)	2e-40	e-100	9e-86	7e-76
06948.1		Ca ²⁺ and/or CaM binding protein	2e-40; synthetic construct (AF084415)	1.3e-35	9.8e-38	None	1e-41
04379.1		Ca ²⁺ and/or CaM binding protein	5e-91; M. grisea (MgNCS1)	1e-59	2e-66	2e-64	2e-13
02738.1		Ca ²⁺ and/or CaM binding protein	1e-37; Mus musculus (AK008610)	4e-20	5.4e-9	1e-37	2e-26
09871.1		Ca ²⁺ and/or CaM binding protein	9e-27; B. cinerea (CNS01CAX)	3e-24	9.5e-24	3e-23	5e-21
01241.1		Ca ²⁺ and/or CaM binding protein	e-133; H. sapiens (SLC25A12)	e-100	5.3e-32	e-133	4e-29
06347.1		Ca ²⁺ and/or CaM binding protein	e-125; A. nidulans (SagA)	3e-43	5e-41	7e-23	None
06617.1		Ca ²⁺ and/or CaM binding protein	5e-68; B. cinerea (CNS019V4)	3.6e-24	3e-32	2e-27	1e-27
03750.1		Ca ²⁺ and/or CaM binding protein	2e-53; B. cinerea (CNS01AXV)	1.8e-11	2.5e-13	3e-5	2e-05
08980.1	NDE-1	Ca ²⁺ and/or CaM binding protein	e-174; Yarrowia lipolytica (NDH2)	e-124	e-134	None	5e-92
02283.1		Ca ²⁺ and/or CaM binding protein	0; Glomerella cingulata (cmk)	9e-61	e-105	7e-67	3e-52
09123.1		Ca ²⁺ and/or CaM binding protein	e-158; A. nidulans (CmkA)	2e-94	1.4e-63	4e-68	3e-49
02814.1		Ca ²⁺ and/or CaM binding protein	6e-71; S. cerevisae (SCYDL101C)	6e-71	3e-53	2e-70	1e-50
09212.1		Ca ²⁺ and/or CaM binding protein	e-134; S. pombe 9SPCC13220	7e-97	e-134	1e-57	1e-42
06650.1		Ca ²⁺ and/or CaM binding protein	1e-32; Helicosporium sp. (BAB70714)	None	None	None	None
02411.1		Ca ²⁺ and/or CaM binding protein	0; Podospora anserina (CAD60740)	7.4e-20	1.9e-23	1e-13	2e-25
06177.1		Ca ²⁺ and/or CaM binding protein	e-112; <i>A. nidulans</i> Ca ²⁺ /CaM-dependent protein kinase C (AAD38851)	4.6e-42	1.5e-55	2e-67	4e-50
04265.1		Ca2+ and/or CaM binding protein	3e-71; <i>Bacillus megaterium</i> β -fructosidase (FruA)	1.3e-35	1e-36	None	1e-33

TABLE 47.	Calcium	signaling	proteins	in	Neurospora

^a CaM, calmodulin.

Neurospora, of which only one (CAX [501]) was previously known. With one exception, all of these proteins have the same homologue in *S. cerevisiae* (Vax1p/Hum1p [532]). The exception is homologous to a hypothetical ORF of unknown function in *S. cerevisiae*. Our analysis also revealed two novel putative Ca^{2+}/Na^+ exchangers in both *Neurospora* and *S. cerevisiae*. It is interesting that animals possess Ca^{2+}/Na^+ exchangers but not Ca^{2+}/H^+ exchangers, plants contain Ca^{2+}/H^+ exchangers but not Ca^{2+}/Na^+ exchangers, while fungi possess both. As with *S. cerevisiae*, only one calmodulin is present in *Neurospora* (125, 518).

Calcium signaling in animal and plant cells normally involves Ca^{2+} release from internal stores (71, 676). This is commonly mediated by the second messengers inositol-1,4,5-trisphosphate (InsP₃) or cADP ribose or by Ca^{2+} -induced Ca^{2+} re-

lease, although other second messengers (e.g., sphingolipids and NAADP) also exist (95).

Four novel phospholipase C- δ subtype (PLC- δ) proteins (which synthesize InsP₃) have been identified in *Neurospora*, in contrast to one in *S. cerevisiae* (22). Furthermore, InsP₃ is present within *Neurospora* hyphae (447), and physiological evidence suggests that it plays a role in Ca²⁺ signaling (170, 737, 738). In spite of this, *Neurospora* (and *S. cerevisiae*) lack recognizable InsP₃ receptors. Whether InsP₃ signaling in *Neurospora* involves InsP₃ receptors that differ from those found in animal cells remains to be determined. Neither ADP-ribosyl cyclase (which synthesizes cADP ribose or NAADP) nor ryanodine receptor proteins (which are also key components of Ca²⁺ release mechanisms in animal cells) are recognizable in *Neurospora*. Furthermore, no homologues of either sphingo-



FIG. 14. Calcium signaling proteins. The percentage of *Neurospora* proteins with the greatest homology to proteins in *S. cerevisiae*, *S. pombe*, animals, and plants is shown.

sine kinase, which synthesizes the second-messenger sphingosine-1-phosphate, or the sarcoplasmic reticulum Ca^{2+} release channel, SCaMPER, which is a possible target of sphingolipids (70), could be indentified in this fungus. These observations raise the intriguing question of whether other, perhaps novel second-messenger systems responsible for Ca^{2+} release from internal stores remain to be discovered in filamentous fungi.

The reason for the diversification of PLC- isoforms is unclear. Besides $InsP_3$, PLC also synthesizes diacylglycerol, a second messenger which regulates protein kinase C (PKC). Two PKCs have been identified in *Neurospora*, but neither possesses a C2 domain with Ca²⁺ binding sites (689).

Another Ca^{2+} -signaling protein which has not been identified in the *Neurospora* genome is the external calcium-sensing receptor described in animal cells (109).

In terms of downstream elements involved in Ca^{2+} signaling, 23 $Ca^{2+}/calmodulin-regulated proteins have been found. Very$ few of these proteins have been analyzed with respect to thebiological responses that they regulate in*Neurospora*. One of $these proteins is the <math>Ca^{2+}/calmodulin-dependent$ Ser/Thr phosphatase calcineurin, which is composed of a catalytic subunit, calcineurin A (CNA), and a regulatory subunit, calcineurin B (CNB) (337, 430). One function of *Neurospora* calcineurin is regulation of hyphal tip growth and branching (631). Another downstream element is a $Ca^{2+}/calmodulin$ dependent protein kinase (CAMK-1) that is involved inregulating growth and the circadian clock (875). A third $downstream element is a <math>Ca^{2+}$ -dependent neutral trehalase (TREB), which is responsible for trehalose mobilization at the onset of conidial germination (188).

The Neurospora Ca²⁺-signaling proteins (Table 47) showed a markedly greater homology to Ca²⁺-signaling proteins in both *S. cerevisiae* and *S. pombe* than to those in animals or plants (Fig. 14). In comparison with just animals and plants, 58% of the *Neurospora* proteins exhibited greater similarity to animal homologues whereas 42% showed higher homology to plant proteins.

Protein phosphatases. Protein phosphorylation and dephosphorylation are essential elements of signal transduction pathways in eukaryotic cells. Protein phosphatases (PPs) are classified as Ser/Thr, Tyr, dual-specificity, or His PPs on the basis

of substrate specificity. More recently, a two-family gene classification has been proposed (164, 202). The two gene families, designated PPP and PPM, are defined by distinct amino acid sequences and three-dimensional atomic structures. The PPP family includes the signature phosphatases PP1, PP2A, PP2B, and PP5 (also known as PPT), while the PPM family comprises the Mg²⁺-dependent protein phosphatases, which include PP2C and pyruvate-dehydrogenase phosphatase.

PP activity in Neurospora was first reported by Tellez de Inon and Torres (787), and more recently, the biochemical and genetic analyses of Neurospora phosphatases (337, 430, 631, 766, 767, 881-884, 893, 896) has been complemented, in part, with the isolation of several phosphatase-encoding genes and their functional analysis (56, 768, 885) (Table 48). Within the PP1 and PP2A subfamilies, several novel PPs have been identified that show less than 65% sequence identity to PP1 and PP2A. These novel-type phosphatases also possess properties indicating that they perform cellular functions distinct from PP1 and PP2A (164) and are also present in Neurospora (Table 48). In addition to the structural features of the catalytic subunits, it is important not to underestimate the involvement of PP regulatory subunits in phosphatase specificity and function (883). Stark (754) has listed some 25 S. cerevisiae proteins associated with phosphatase function and suggests that there are still considerably more PP regulatory polypeptides to be identified. Although in some instances, structural conservation between such polypeptides in various organisms may assist in identifying some of these polypeptides in filamentous fungi (196), the structural diversity of PP regulatory proteins is likely to be much greater than that of the catalytic counterparts. In accordance, only a few of the more highly conserved regulatory PP subunits have likely structural homologues in *Neurospora*, while the presence of proteins considered to be associated with PPs in other organisms cannot readily be detected on the basis of structural similarity searches.

Mammalian signaling proteins not found in *Neurospora*. Although the SH2 domain (reviewed in reference 821) is present in some proteins, *Neurospora* does not contain recognizable tyrosine kinases, including c-Src (reviewed in references 308 and 821) and nuclear hormone receptors (reviewed in reference 645). In addition, *Neurospora* lacks proteins with scores less than 2e-10 to the SH2- or SH3-containing Grb2, Shc, and Sck (556) tyrosine kinase adaptor proteins. Raf kinases (reviewed in reference 614) are apparently absent from the genome, as are β -arrestin homologues (reviewed in reference 308). There are no good matches to the *S. cerevisiae* Ste5p MAPK scaffold (218) or to mammalian scaffold proteins, such as MP-1 (683) or JIP-1 (879). There is no GTPase protein homologous to Gh (558), which regulates PLC- δ activity in mammals.

Photobiology and Circadian Rhythms

Neurospora is equipped to respond to, and anticipate predictible changes in, a variety of environmental stimuli. Aside from the obvious role of nutrition, the most prominent and pertinent environmental factors are ambient light and temperature, both of which affect the circadian system of the organism. Acutely, light is known to affect the expression of a large number of genes influencing carotenogenesis, conidiation,

	NCU		BLAST n	natch		
Enzyme	no.	Best overall	S. cerevisiae	S. pombe	Animal ^a	Plant ^b
PPP1: type 1 serine/threonine-specific protein phosphatase	00043.1	A. nidulans BimG/e-178	e-166	e-171	e-161; M. musculus	e-148
PPH1: type 2A serine/threonine-specific protein phosphatase, catalytic subunit	06630.1	H. sapiens P2AA/e-142	e-138	e-141	e-149; H. sapiens	e-142
CNA: serine/threonine protein phospha- tase 2B catalytic subunit (calmodulin- dependent calcineurin A subunit)	03804.1	A. nidulans CnaA/0.00	e-167	0.00	0.00; M. musculus	9e-61
Protein phosphatase 2C	00434.1	S. pombe p2c1/4e-70	2e-65	4e-70	2e-22; H. sapiens	7e-19
PPE-like serine/threonine-specific pro- tein phosphatase	03436.1	S. pombe ppe1/e-112	e-108	e-112	e-101; H. sapiens	e-91
PPG-like serine/threonine-specific pro- tein phosphatase	06563.1	S. pombe YD44/e-120	e-110	e-120	2e-95; H. sapiens	4e-97
PPT (type 5) serine/threonine-specific protein phosphatase	01433.1	R. norvegicus PPP5/e-136	e-111	e-55	e-136; M. musculus	5e-57
Protein phosphatase X	08301.1	H. sapiens PPP4/e-41	4e-38	2e-38	e-41; H. sapiens	e-41
PZL1: type Z serine/threonine protein phosphatase	07489.1	S. cerevisiae Ppz2p/e-141	e-141	e-116	e-120; H. sapiens	e-119
Pyruvate dehydrogenase phosphatase, catalytic subunit 1 and similar to type 2C protein phosphatase	01767.1	H. sapiens PDP1/3e-44	2e-12	5e-13	3e-44	2e-9
RGB1: type 2A phosphatase regulatory B subunit	09377.1	S. pombe 2aba/e-178	e-149	e-178	e-132; M. musculus	e-141
Putative type 2A protein phosphatase B56 regulatory subunit	03786.1	S. pombe e-150	e-148	e-150	e-142; H. sapiens	e-114
Putative Type 2A phosphatase-associ- ated protein	08268.1	S. pombe putative protein/4e-39	5e-28	4e-39	2e-21; H. sapiens	e-24
CNB1: calcineurin regulatory subunit	03833.1	S. pombe, probably calcineurin B subunit/5e-54	e-51	5e-54	3e-47	5e-18
Putative phosphatase-associated protein	10018.1	S. cerevisiae Sit4p-associated protein/3e-82	3e-82	2e-42	4e-35; H. sapiens	2e-21
Putative phosphatase-interacting protein	08779.1	S. cerevisiae Gip2p/2e-10	2e-10	6e-9	6e-8; H. sapiens	No hits

TABLE 48. N. crassa Ser/Thr phosphoprotein phosphatas

^b Arabidopsis thaliana.

perithecial neck bending, and directionality in ascospore shooting (184, 477). A major, but less immediate, effect of light derives from its central role in phasing the circadian clock, which, even in the absence of light, can influence the expression of several percent of the genome. In published work on *Neurospora* light sensing extending back over 100 years (38, 184), mention is made only of its ability to sense blue light; no red-light responses are known. An unexpected aspect of the *Neurospora* genome is the presence of novel genes whose products reflect environmental sensing of light stimuli.

Perhaps largely reflecting the fact that circadian rhythmicity has been intensively studied in Neurospora for the past 15 to 20 years (61, 206, 484), there were no notable additions to the list of proteins involved in the core regulatory feedback loop. Known central clock components (Fig. 15; Table 49) that function in the core circadian oscillator in continuous darkness include FRQ (29, 30), WC-1 and WC-2 (173), and the kinases that act on them in a circadian context, including casein kinases 1 (291) and 2 (874) and calmodulin-regulated kinase (875). It is perhaps interesting that WC-1 retains the dark function as a positive element in the circadian feedback loop, similar to its mammalian counterpart BMAL1, but also has acquired a lightbased activity as the blue light photoreceptor, as described below. There were no unidentified duplicates of any core clock genes found, nor were there sequence homologues to the Drosophila or vertebrate period or CLOCK genes, although a gene similar to Drosophila vrille is found; the strong sequence homology between Neurospora WC-1 and the vertebrate clock protein BMAL1 has already been noted (461). Likewise, the PAS protein VVD, which acts to link circadian output with light input/output, was already identified (328, 697, 733). In *Drosophila*, the *shaggy* gene, encoding a GSK-3 kinase homologue, plays an important role in regulating nuclear movements of the key negative element TIM (503). *Neurospora* does contain a GSK-3 homologue, whose significance in the circadian feedback loops has yet to be appraised.

In spite of intensive study for two decades (478), and in contrast to the analysis of clock components, the genome sequence revealed a surprising number of homologues to novel light-sensing genes (Fig. 16; Table 49). DNA photolyase (see "Genome defense, DNA repair, and recombination" above), previously characterized as a photoresponsive molecule, plays no role in light responses beyond DNA repair (730). Known elements in light sensing include the circadian blue-light photoreceptor WC-1 (42, 256, 323), WC-2 (478), and VVD (328, 698); however, no homologues of plant photoresponse proteins such as phytochromes or cryptochromes have ever been identified, despite intense genetic screening. Nonetheless, they are present, including clear phytochrome and cryptochrome homologues and a gene whose product, like VVD, contains a single PAS/LOV-type domain of the type associated with light sensing. There is also a homologue of the Aspergillus velvet gene, implicated in regulation of both red- and blue-light responses (537).

Perhaps the biggest surprise to emerge from the genome analysis is the appearance of a cryptochrome, which had pre-



FIG. 15. Known molecular components in the coupled feedback loops of the *Neurospora* circadian system. The WC-1 and WC-2 proteins form a White Collar Complex (WCC) that activates *frq* gene expression and also *clock-controlled gene* (*ccg*, output) and *vvd* expression in the dark. The WCC also mediates light-induced transcription from *frq*, *ccg* genes, *vvd*, and *wc-1* (gold arrows). VVD expression is strongly light induced, and VVD in turn is a photoreceptor that mediates light adaptation responses, transiently turning down the WCC activity. In the circadian cycle in the dark, *frq* mRNA is translated to make FRQ proteins which dimerize and play two roles: (i) FRQ feeds back into the nucleus to rapidly block the activity of the WCC in driving *frq* transcription, and (ii) FRQ acts to promote the synthesis of new WC-1 and *wc-2* mRNA, thus making more WCC, which is held inactive by FRQ. Phosphorylation of FRQ by several kinases, including casein kinases 1 and 2 and CAMK-1, triggers its turnover mediated by an interaction with the ubiquitin ligase encoded by *fwd-1;* the kinetics of phosphorylation-mediated turnover is a major determinant of period length in the clock. When FRQ is degraded in the proteasome, the pool of WCC is released to reinitiate the cycle. See the text for details. Adapted from reference 461

viously been extensively sought by other molecular methods, along with two phytochromes. At first glance, the appearance of two phytochromes is unexpected, given that these molecules in plants are chiefly known in the context of red/far-red sensing, and to date, no red-light photobiology has been described in *Neurospora*. However, since *Arabidopsis* phytochromes have recently been shown to associate with cryptochromes and to play a role in blue-light sensing and signaling (194), this aspect of photobiology would be a likely candidate for the *Neurospora* phytochromes. The same might be said for the *velvet* homologue.

The Neurospora genome contains nearly a dozen proteins containing PAS/PAC domains (compared to four in *S. cerevisiae* and five in *S. pombe*). PAS and PAC proteins as a class are associated with light, oxygen, and voltage sensing and are often associated with protein-protein interaction domains, such as those in the heterodimers that act as positive elements in fungal and vertebrate circadian systems (783). In some cases, the PAS domains are juxtaposed to HK domains. These domain associations are characteristic of bacterial and plant two-component regulatory systems and have been associated with light responses in the cyanobacterium *Synechococcus* (385) (see also Two-component regulatory systems above).

Heat Shock and Stress Responses

All classes of the major heat shock-induced proteins are encoded in the genome of *Neurospora* (Table 50). These stress proteins, which are ubiquitously and abundantly synthesized in response to supraoptimal temperature and other protein-denaturing stresses, function as chaperones that guide and stabilize the conformation of other proteins. Most heat shock proteins (Hsps) are also synthesized constitutively, and many are essential.

DnaK/Hsp70, the first Hsp to be characterized, is strongly conserved among many organisms. It has an amino-terminal ATPase domain and a carboxyl-terminal peptide binding domain. The release of peptides by Hsp70, allowing them to refold, is linked to ATP hydrolysis (278). Homologues of the chief cytosolic classes of Hsp70 (*S. cerevisiae* nomenclature) are present in *Neurospora*: the major heat-induced Ssap, which was previously sequenced in *Neurospora* (400), the cold-inducible Ssbp, and Ssz1p. Whereas *S. cerevisiae* has multiple Ssap and Ssbp proteins, *Neurospora* has only one of each. Ssap is required for protein translocation into mitochondria and the ER (191), and each of the three cytosolic Hsp70s contributes to mRNA translation (350, 356). *Neurospora* also has organelle-localized Hsp70s that assist in protein translocation

Protein	Neurospora	Function	NCU			BLAST match		
class	protein		no.	Best overall	S. cerevisiae	S. pombe	Animal ^a	Plant^{b}
Clock-associated genes	FRQ WC-1	N. crassa central clock component N. crassa blue-light photoreceptor, central oscil- lator component, 2 PAS domains, 1 LOV domain GATA zinc fuxer domain	02265.1 02356.1	Sordaria fimicola; e 0.0 Podospora anserina; e 0.0	None Ymr1369; e-06	None SPCC1902.01; 5e-05	None Drosophila; e-14	None NP851210.1; 4e-18
	WC-2	Forms compares with WC-1, central oscillator forms component and blue-light signaling, 1 PAS component and blue-light signaling, 1 PAS domain, GATA zinc finger domain	00902.1	Fusarium solani; e-138	Ymr136W; 6e-08	None	None	AC026875.23; 3e-07
	CKA	N. crassa casein kinase II catalytic subunit	03124.1	Candida albicans; e-148	YorR061W	SPAC23C11 CKII;	P28020; e-105	AF370308; e-112
	CKB1	N. crusta in 1100 N. crusta castin kinase II regulatory subunit involved in FRO nboschorolation	05485.1	S. pombe; 4e-61	Yg109W CKB1;	SPBC2G5.02C CKII: 4e-61	P28021; 3e-46	AF239816; 4e-41
	CKB2	N. crasse case kinese II regulatory subunit involved in FRO phoenhordation	02754.1	S. pombe; 2e-77	Y1r039C RIC1; 8e-4	SPAC1851.04e;	XP317865.1; 2e-60	NP181996.1; 1e-58
	HHP1	N. crasse case in kinase I involved in FRQ	00685.1	S. pombe; e-150	NP015120.1; e-125	NP595760.1; e-150	BAB0347 3.1; e^{-135}	BAB92346.1; e-130
	CAMK	Purospiroryation N. creases protein kinase involved in FRQ	09123.1	Aspergillus nidulans; 10-165	Yoi016C CMK2;	SPACUNK12.02c;	NP065130.1; 2e-69	NP192383.1; 8e-49
	SHAGGY/GSK-3 homologue	prosprospration Drosophild protein kinase Shaggy involved in phosphorylation and nuclear movement of the central socillator commonent TIM	04185.1	Colletotrichum gloeo- sporioides; e-157	Ymi139W Rim11; e-104	SPAC1687.15; e-148	NP571465; e-130	AF43225.1; e-119
	FWD-1	E3 ubiquitin ligase involed in targeting FRQ for proteasomal turnover	04540.1	Homo sapiens; e-47	None	SPAC30.05; e-40	Homo sapiens; e-47	None
Other light-	VVD	N: crassa blue-light sensing, PAS/LOV domain	03967.1	Aspergillus nidulans;	None	None	None	BAA84780.1; 3e-19
201128 601100	PHY-1	Putative phytochrome red/far-red-light sensing	04834.1	Pseudomonas putida; 8e-61	Ylu206W; 3e-09	SPAC27E2.09; 2e-27	None	P55004; 4e-33
	PHY-2	Putative phytochrome red/far-red-light sensing	05790.1	Agrobacterium tume- faciens: 2e-33	Yi1147C SLN1; 8e-9	SPAC1834.08;	None	P55004; 2e-21
	CRY homolog	Putative blue-light sensing, central oscillator component in mammals, light entrainment of the clock in <i>Drosophila</i>	00582.1	Trichodesmium eryth- raeum; 4e-69	Yoi386w PHR1; 4e-26	None	None	NP58461.2; 2e-49
	VELVET homologue	Red- and blue-light sensing in Aspergillus	01731.1	Aspergillus nidulans; 5e-40	None	None	None	None
	NOP-1	Neurospora homologue of bacteriorhodopsin	10055.1	Leptosphaeria maculans onsin: 2e-75	Yro2p; e-08	SPCC31H12.02c; 9e-07	None	None
	ORP-1	Neurospora opsin-related protein, lacks conserved lysine residue	01735.1	Coriolus versicolor Hsp30; 5e-31	Yro2p; 8e-23	SPCC31H12.02c; 8e-12	None	None
Other PAS proteins	Hypothetical protein	4 PAS domains, sensory box histidine kinase/ resnonse reculator	03164.1	Shewanella oneidensis; 6e-31	Yi1147C SLN1; 2e-10	SPAC27E2.09; 1e-20	None	None
	Predicted protein	3 PAS domains	06390.1	Caenorhabditis elegans; e1.4	None	None	None	None
PAS/PAC do-	Hypothetical protein	PAS/PAC domain 1e-88, sensory transduction histiding binase	02057.1	Magnetococcus sp. strain MC-1- 4e-75	Yi1147C SLN1; 4e-18	SPAC183 4.08; 6e-57	None	None
	Hypothetical protein	PAS/PAC domain 3e-59, two-component hybrid	00939.1	Anabaena sp. strain PCC 7120: 4e-23	Yill47C SLN1; 6e-11	SPAC27E2.09;	None	None
	Hypothetical protein	PaS/Pac duration and a second particle two-compo- part histiciane timese Foc.1	07221.1	Aspergillus fumigatus;	Yi1147C SLN1; T_{2-17}	SPAC27E2.09;	None	None
	Hypothetical protein	PAS/PAC domain 3-68, related to two-compo- nent histidine kinase <i>chk_I</i>	01833.1	Glomerella cingulata; 0.0	Ylu206W; e-10	SPAC1834.08; 8e-79	None	None
	Hypothetical protein	PAS/PAC domain 56-42, probable serine/threo- nine protein kinase, related to plant NPH-1 protein	07268.1	Caulobacter crescentus; 2e-07	None	None	None	T013535; e-07

TABLE 49. Known and predicted N. crassa proteins associated with circadian clocks and photobiology

^a Caenorhabditis elegans, Rattus norvegicus, Xenopus laevis, Drosophila melanogaster, Anopheles gambiae, Mus musculus, or Homo sapiens. ^b Arabidopsis thaliana, Zea mays, Ipomoea nil, or Oryza sativa.

BORKOVICH ET AL.

70



FIG. 16. Real and potential *Neurospora* photoreceptors. The approximate sizes and locations of pertinent protein functional domains are shown for this series of proteins having known or plausible roles in photobiology. WC-1 and WC-2 work together as the White Collar Complex they are known to comprise a photoreceptor that appears to be the circadian photoreceptor and a major blue light photoreceptor in *Neurospora*. VVD is also a blue light photoreceptor that is responsible for modulating the WCC and contributing to photoadaptation. NOP-1 binds retinal and undergoes a photocycle, but the associated photobiology has not been elucidated (see also Fig. 12). Likewise, CRY, PHY-1, and PHY-2 all show strong sequence homology to known photoreceptors from other organisms (Table 49; Fig. 11) but do not yet have any demonstrated role in photobiology.

and protein folding within the organelle: Ssc1 in mitochondria (763) and Kar2 (663) and Lhs1 (681) in the ER. *Neurospora* has five other Hsp70-related proteins that do not have known counterparts in fungi and plants but whose closest homologues are a group of related proteins in vertebrates. The mitochondrial Ssq1p of *S. cerevisiae* (763) is absent from *Neurospora*.

Hsp110 of mammals is a distant relative of Hsp70 that is strongly induced by heat shock (463). Its homologue in *Neurospora*, HSP88, was identified by its ability to bind to the major small Hsp (sHsp [see below]) of *Neurospora* (619), HSP30, an affinity also described for HSP110 and the mammalian sHsp (844), suggesting that these two proteins may be cochaperones. The homologous Sse proteins of *S. cerevisiae* are reported to interact physically and functionally with Hsp90 (480).

Hsp90 is a major, conserved heat shock-induced protein that has important functions in signal transduction during normal growth by directly regulating protein kinases and transcription factors. Hsp90 was shown to negatively regulate cell cycle progression in S. pombe by its interaction with the Wee1 tyrosine kinase (15). cDNA for the single Hsp90 gene of Neurospora was previously sequenced (666), and, like S. cerevisiae, Neurospora lacks Grp94, the ER-localized Hsp90 of animal cells. Hsp90 has an amino-terminal ATPase domain to which the inhibitor geldanamycin binds, thereby blocking Hsp90 homodimerization and binding of the p23 cochaperone (630). p23, which couples ATPase activity with substrate release by Hsp90 (890), is present in *Neurospora* but is poorly conserved across species. Neurospora also has a homologue of Aha1, which binds to the middle region of Hsp90 and enhances its ATPase activity (486). Assembly of Hsp70 and Hsp90 into a superchaperone folding complex is mediated by the cochaperone Hop/Sti1, which is an adaptor protein that interacts with both chaperones through separate tetratricopeptide repeat (TPR) domains (2). Sti1 is strongly conserved in Neurospora. A less strongly conserved homologue of Sti1, Cns1, which binds to Hsp90 (201), is also present in the *Neurospora* genome.

Protein folding by mitochondrial GroEL/Hsp60 has been carefully analyzed. It has been shown that the *Neurospora* HSP60, like other Hsp60s, assembles into two stacked rings of seven monomers each (358). Unfolded proteins bind at hydrophobic apical domains of the Hsp60 structure, refold in the more hydrophilic interior, and are released as ATP is hydrolyzed (865). Its cochaperone GroES/Hsp10 (488), present in *Neurospora*, is moderately conserved across species.

Although a Clp-related Hsp was initially ignored, due to its absence from *D. melanogaster*, where the heat shock response was first characterized, subsequent experiments with *S. cerevisiae* showed that Clp/Hsp104p is important for the acquisition of induced thermotolerance (675). Hsp104p is an ATP-dependent chaperone that, when assembled into hexamers, possesses the unique ability to disentangle protein aggregates after they have formed rather than merely preventing their formation (601). Vertebrates do have a moderately conserved counterpart to Hsp104p (609), and it is strongly conserved in *Neurospora*, as is the mitochondrially localized Clp, Hsp78, which helps turn over unassembled mitochondrial proteins in *S. cerevisiae* (468).

A relatively nonconserved but ubiquitous group of Hsps share a conserved region with α -crystallin of the vertebrate eye lens (363). These small Hsps (sHsps) assemble into large oligomeric particles that function as non-ATP-dependent chaperones and are thought to be the first line of defense against stress (212). The sHsps are nonessential proteins that are induced chiefly in response to stress and during development. There are three sHsps in the *Neurospora* genome, of which the most highly conserved, HSP30, has been characterized (621). RIP mutagenesis indicated that HSP30 helps *Neurospora* survive extended exposure to high temperature coupled with carbohydrate deprivation (620), conditions that would promote
TABLE	50.	Heat	shock	and	stress	proteins
-------	-----	------	-------	-----	--------	----------

D. () ()		NOU	В	LAST match			
Protein family	Name/reature	NCU no.	Best overall	S. cerevisiae	S. pombe	Animal ^a	Plant ^b
Hsp70	Hsp70	09602.1	Paracoccidiodes brasiliensis; 0.00	0.00	0.00	0.00	0.00
1	Grp78/Kar2:ER	03982.1	Aspergillus amawori; 0.00	0.00	0.00	0.00	0.00
	Ssb	02075.1	Aspergillus nidulans; 0.00	0.00	0.00	e-177	e-175
	Ssc1:Mt	08693.1	S. cerevisiae; 0.00	0.00	e-178	e-166	e-145
	Ssz1	00692.1	<i>S. cerevisiae</i> ; e-104	e-104	e-102	8e-54	3e-49
	Lhs1:ER	09485.1	Anopheles gambiae; 4e-73	2e-43	3e-65	1e-64	1e-52
	KIAA0417	00573.1	Mus musculus; /e-26	1e-05	2e-06	/e-26	4e-05
	KIAA041/ Uses 12b	01499.1	Homo sapiens; 1e-23	1e-06	2e-04	1e-23	2e-04
	пара120 КГА А0/17	04390.1	H. sapiens, 5e-21 H. sapiens: 6e 10	36-00	46-04	5e-21	/e-04
	KIAA0417 KIAA0417	03288 1	M. musculus: 7e-05			7e-05	
Hsn110	HSP88	05269.1	S pombe: 0.00	e-168	0.00	e-136	e-132
Hsp90	Hsp90	04142.1	Podospora anserina: 0.00	0.00	0.00	0.00	0 102
Hsp90 associated	p23/Wos2	01792.1	S. pombe: 8e-20	2e-16	8e-20	2e-11	1e-05
r-	Aha1	04087.1	<i>S. pombe</i> ; 2e-71	3e-51	2e-71	6e-37	2e-19
	Hop/Sti1	00714.1	<i>S. pombe</i> ; e-138	e-129	e-138	3e-95	2e-91
	Cns1	06340.1	<i>S. pombe</i> ; 5e-26	7e-25	5e-26	1e-16	2e-20
Hsp60	Hsp60:Mt	01589.1	Coriolus immitis; 0.00	0.00	0.00	e-162	e-173
Hsp10	Hsp10:Mt	04334.1	<i>S. pombe</i> ; 4e-28	6e-27	4e-28	2e-17	1e-23
Clp	Hsp104	00104.1	P. brasiliensis; 0.00	0.00	0.00	8e-46	0.00
	Hsp78:Mt	02630.1	Leptosphaeria maculans; 0.00	0.00	5e-38	0.00	
sHsp	HSP30	09364.1	A. nidulans; 7e-27	0.69	2e-05		4e-04
	[mito]	07232.1	A. nidulans; 8e-23	1.3	6e-04		2e-04
		09420.1	A. nidulans; 8e-17	• • • •	0.20		0.41
Hsp40	Ydj1	07414.1	<i>S. pombe</i> ; e-107	2e-98	3e-107	2e-85	9e-70
	Ydj1	00465.1	<i>S. pombe</i> ; 93-67	3e-49	9e-67	3e-56	2e-49
	Mdj1:Mt	05196.1	S. pombe; /e-/9	36-55	/e-/9	8e-4/	1e-60
	Zuoi SaitED	03009.1	S. pombe; 3e-81	3e-60	3e-81	2e-41	1e-26
	SCJ1:EK	8.1	<i>5. pombe</i> ; 1e-68	46-00	16-08	20-55	36-39
	Sis1	03732.1	<i>S. pombe</i> ; 4e-61	3e-31	4e-61	2e-44	3e-56
	Sec63:ER	00169.1	<i>S. pombe</i> ; 4e-61	1e-41	4e-61	3e-44	2e-27
	[TPR domains]	00170.1	<i>S. pombe</i> ; 3e-80	9e-12	3e-80	2e-69	3e-37
	Mandelate racemase	07064.1	Burkholderia fungorum; e-150	3e-11	1e-29	e-121	5e-21
		02432.1	P. anserina; e-129	9e-43	1e-20	2e-40	4e-36
	Djp1	06052.1	<i>P. anserina</i> ; e-169	2e-35	2e-50	1e-18	1e-26
	Hlj1:ER	03335.1	<i>S. pombe</i> ; 9e-29	4e-19	9e-29	2e-19	9e-13
	[TPR domain]	02424.1	P. anserina; e-161	2e-11	5e-16	1e-43	2e-10
		05/10.1	Rhodospirillum rubrum; 9e-12	2e-07	Se-07	2e-11	4e-11
		05199.1	Salmonella enterica; 1e-11 Plagmo dium falsingmum 0o 12	3e-08	8e-11	5e-11	5e-11
		04505.1	Chlorobium tanidum: 20.11	10-09	3e-08	40-10 50.11	9e-11 60.08
		04145.1	P falcingrum: 1e 10	4e-07	20.05	20.06	50 08
GrnF	Mge1·Mt	01516.1	S cerevisiae: 2e-41	2e-41	1e-39	20-00 3e-25	1e-23
CIPL	Fes1	04172.1	S cerevisiae: 4e-17	4e-17	5e-14	0.25	2e-06
	SIs1:ER	00968.1	Yarrowia lipolytica: 4e-11	5e-08		4e-06	6.2
Bag-1		01221.1	<i>S. pombe</i> : 0.015		0.015	0.081	
Cyclophilin	Cvp40/Cpr6	03853.1	M. musculus; 1e-81	4e-76	1e-72	1e-81	2e-71
- J I	CvpC/Cvp1	00578.1	Aspergillus niger; 5e-58	9e-28	2e-51	8e-53	2e-51
	Cyp20/Cpr1:Mt/Cyt	00726.1	Fusarium sporotrichioides; 4e-70	3e-47	2e-48	9e-47	6e-44
	CypB:ER	01200.1	Aspergillus niger; 1e-65	2e-48	2e-60	2e-57	5e-50
	[U-snRNP-assoc]	02614.1	<i>Echinococcus multilocularis</i> ;	3e-43	8e-47	6e-49	6e-45
	KIAA0073 [WD40] [nucleus]	09819.1	<i>H. sapiens</i> ; e-153	8e-27	e-134	e-153	e-142
	[U-Box]	00181.1	Drosophila melanogaster; 2e-83	4e-16	2e-30	2e-83	5e-39
	[nucleus]	08514.1	Arabidopsis thaliana; 2e-47	3e-15	2e-44	5e-45	2e-47
	Ppil4 [RNA-Bind] [nucleus]	07179.1	<i>S. pombe</i> ; 6e-99	1e-10	6e-99	2e-85	1e-82
FKBP	FKBP13:Mt/Cyt	04140.1	P. anserina; 2e-29	4e-23	6e-24	3e-26	4e-26
	FKBP22:ER	02455.1	P. anserina; 1e-36	2e-11	2e-07	2e-16	1e-12
	Fpr4:Nucleus	03241.1	S. cerevisiae; 6e-36	6e-36	4e-17	1e-26	3e-20
	Fpr1	04371.1	<i>S. pombe</i> ; 4e-17	7e-13	4e-17	8e-13	1e-11
Hsp30	ORP-1	01735.1	Coriolus versicolor; 5e-31	7e-23	8e-12		5.1
	NOP-1	10055.1	L. maculans; 2e-75	1e-08	9e-07	0 00	0 1-
HSF	HSF	08512.1	H. sapiens; 2e-22	7e-16	1e-19	2e-22	3e-15
		08480.1	S. cerevisiae; 3e-04	3e-04	0.018	0.029	0.006

^a Caenorhabditis elegans, Drosophila melanogaster, Mus musculus, or Homo sapiens. ^b Arabidopsis thaliana or Oryza sativa.

oxidative stress. The hsp30 RIP mutants were defective in glucose-phosphorylating activity and in mitochondrial protein import (617, 618). The next most highly conserved sHsp of Neurospora, which, like HSP30, is induced by heat shock, has the properties of an imported mitochondrial protein with a targeting presequence (MitoProt [152]) that was, unfortunately, omitted by automated gene prediction programs. There are also sHsps of plants and D. melanogaster with mitochondrial targeting presequences that are imported into mitochondria (539). Conditions under which the least highly conserved sHsp of Neurospora is expressed are not known. Despite their sequence divergence, these three sHsps of Neurospora are most similar to one another in the N-terminal domain that is predicted to form an amphipathic helix (HelixWheel, www.site .uottawa.ca/~turcotte/resources/HelixWheel/ [M. Turcotte, 1996]), a predicted structure that is also conserved.

The largest class of Hsps in Neurospora is the DnaJ/Hsp40 proteins, which number 18 in the genome; 16 Hsp40s have been identified in the S. cerevisiae genome (871). Despite the similar number of Hsp40 proteins in Neurospora and S. cerevisiae, many of them are unrelated to one another. The Hsp40 class is defined minimally by containing a "J domain" through which the proteins interact with Hsp70. Hsp40s function as independent chaperones, as well as being cochaperones for Hsp70s, whose ATPase activity they enhance (23). Several Hsp40s have a domain structure throughout that resembles that of the prokaryotic DnaJ, including a central cysteine-rich zinc finger domain and a characteristic C-terminal domain, as well as the N-terminal J domain (871). There are four of these Hsp40s in Neurospora that correspond most closely to the S. cerevisiae proteins Ydj1p (two), Scj1p, and Mdj1p. Like Ydj1p, the Neurospora Ydj1p homologues terminate in a CAAX motif that signals isoprenylation and that is required for Ydj1p association with nuclear membranes (126). Scj1p and Mdj1p are located in the ER lumen and mitochondrial matrix, respectively, where they cooperate with organellar Hsp70s in protein folding (665, 736). Neurospora also has a homologue of Sis1p, an essential cytosolic protein in S. cerevisiae, which has the conserved J and carboxyl domains (871).

The other conserved Hsp40s of Neurospora have only the J domain. These include homologues of ER membrane-localized Sec63, which interacts with luminal Hsp70 in protein import (671); Zuo1, which binds Ssb Hsp70 on ribosomes (872); and cytosolic Djp1, which is required for protein import into peroxisomes (335). Neurospora has a less strongly conserved counterpart of the ER membrane-localized Hlj1p (316). Other Hsp40-related proteins that contain the J domain do not have obvious counterparts in S. cerevisiae. Two of these proteins contain TPR domains, and one is in the mandelate racemase enzyme family. Of the unidentified Hsp40s of Neurospora, two show moderate similarity and five show little similarity to hypothetical proteins of other species. Hsp40s of S. cerevisiae that are absent from the Neurospora genome include microsomal Jem1p and mitochondrial Mdj2p and Jac1p, which interacts with the minor mitochondrial Hsp70 Ssq1p (836). Some of the unique Hsp40s of Neurospora may interact with Hsp70 proteins for which no orthologue is evident in other organisms.

Prokaryotic GrpE is a nucleotide exchange factor for DnaK, and in *S. cerevisiae* mitochondria the GrpE homologue, Mge1p, forms a complex with mitochondrial Hsp70/Ssc1p and Hsp40/ Mdj1p (93). This complex is essential for the import of proteins into mitochondria and for their folding. Neurospora has a conserved homologue of Mge1p; however, much less strongly conserved in Neurospora and other species is an ER-localized protein, Sls1p, which was shown in S. cerevisiae to act as a nucleotide exchange factor for Kar2p, the ER Hsp70 (391). Neurospora also has a counterpart to S. cerevisiae Fes1p, a cytosolic homologue of Sls1p that interacts with Ssa1p. Fes1p is associated with polysomes, and it facilitates translation together with Ssa1p and Ydj1p (390). Bag-1 is another type of protein shown to be an Hsp70 cochaperone in mammalian cells and to act as its nucleotide exchange factor (344). Bag-domain proteins are poorly conserved overall, but they were identified in Neurospora, S. cerevisiae, and S. pombe by the presence of conserved residues essential for Hsp70 interaction (746, 774). The S. cerevisiae Bag-domain protein Snl1p, which bears little sequence similarity to the N. crassa protein, was shown to function as a cochaperone of Hsp70 (746).

Immunophilins are cis-trans peptidyl-prolyl isomerases that are expressed at higher levels during heat shock and that assist in protein folding; they are present in the cytosol and in organelles. In Neurospora, one gene encodes both the cytosolic and the mitochondrial cyclophilins, which bind cyclosporin A (801), and, similarly, one gene encodes the cytosolic and the mitochondrial FK506 binding proteins, FKBPs (802). Like other organisms, Neurospora also has a cyclophilin and an FKBP that localize to the ER (745), as well as a nuclear FKBP. Three cyclophilins of Neurospora are predicted to be nuclear (PredictNLS [165]); one contains an RNA binding domain, one contains a WD40 repeat, while the third contains a U-box (ring finger) domain. In addition to acting as independent chaperones, a subset of immunophilins contain a TPR domain through which they bind to Hsp90 and Hsp70 as part of the superchaperone complex (582). Neurospora has one TPR-containing 40-kDa cyclophilin compared with two in S. cerevisiae: Cpr6p and Cpr7p, which bind to Hsp90 (205). Similar to S. cerevisiae, none of the four Neurospora FKBPs has a TPR domain comparable to that of mammalian FKBP51 and FKBP52 that would indicate involvement with the Hsp90/Hsp70 superchaperone complex.

Neurospora has two seven-transmembrane helix proteins that are homologous to the plasma membrane-localized Hsp30p of *S. cerevisiae*. *S. cerevisiae* Hsp30p is a non- α -crystallin-related protein that protects the membrane ATPase from high-temperature stress (592). The more strongly conserved homologue in *Neurospora* (ORP-1; see also Tables 44 and 49) may be functionally similar to Hsp30p. The less strongly conserved homologue is an opsin (NOP-1), which has been characterized and shown to bind retinal (see also "Major signal transduction pathways" above) (Tables 44 and 49) (78).

The heat shock transcription factor (HSF) is central to the mechanisms used by cells to dramatically increase the transcription of Hsp genes. HSF becomes activated by the accumulation of unfolded proteins, resulting from heat shock or other stresses (803), which diverts the Hsp70 and Hsp90 chaperones from their inhibitory binding to HSF (352). Plants have multiple HSFs (21 for *A. thaliana* [571]), while vertebrates have four and *S. cerevisiae* has only one (780). *Neurospora* has two HSFs, one of that is moderately conserved and one that is divergent.



co2

succinyl-CoA

The assembly of HSF into homotrimers is mediated by its coiled-coil domains (748). However, most stress-induced HSFs are constrained from trimerization, until activated, by intramolecular coiled-coil formation (852). The HSF of *S. cerevisiae*, on the other hand, forms trimers and binds to DNA in the absence of stress (748), presumably due to a lack of strong intramolecular interactions. The conserved HSF of *Neurospora*, like the stress-induced HSF1 of mice, has three regions with high propensity for coiled-coil formation (490), suggesting that the *Neurospora* HSF would require activation for homotrimer formation. The more divergent HSF of *Neurospora* possesses an HSF-like DNA binding domain, but it has no predicted coiled-coils.

succinate

GROWTH AND REPRODUCTION

Cell Wall

The fungal cell wall protects the organism from an osmotically and, in the case of pathogens, an immunologically hostile environment. The cell wall not only serves a supportive function but also plays a dynamic role in all aspects of fungal physiology. For filamentous fungi, including *Neurospora*, growth and cell wall assembly occur only at each hyphal apex (119, 324, 656, 668, 799, 851, 863). This is in contrast to yeasts, where extension occurs at bud tips, followed by intercalary growth (reviewed in references 424, 479, and 493). *Neurospora* walls are composed of 7 to 10% chitin (a polymer of *N*-acetylglucosamine [GlcNAc]), 25% (1,3)β-linked glucans, 35% other glucans, and 10% proteins (47, 668). Each carbohydrate polymer is synthesized de novo at hyphal tips. The synthase responsible for production of each polymer is transported in an inactive form to hyphal tips in vesicles that fuse with the apical plasma membrane and then begin extruding each carbohydrate polymer through the membrane (372). The wall is assembled exterior to the membrane by processes not completely understood.

Glucan synthases. Although *S. cerevisiae* cell wall structure and assembly have served as the general models for fungal cell wall assembly, recent results, including the analysis of the *Neurospora* genome, have underscored some fundamental differences. Most notable is the observation that *Neurospora* lacks not only $(1,6)\beta$ -linked glucans but also all of the enzymatic machinery required for $(1,6)\beta$ -glucan synthesis encoded in the *S. cerevisiae* genome (see below).

(i) (1,3) β -Glucan synthesis. The synthesis of (1,3) β -linked glucan is catalyzed by an enzyme complex composed of at least two proteins, the (1,3) β -glucan synthase catalytic subunit encoded by a single gene in *Neurospora* (FKS) and a regulatory subunit, Rho1 (Fig. 17; Table 51) (35, 398, 576, 661). In *S. cerevisiae*, there are two and possibly three genes, FKS1, FKS2, (and FKS3), each encoding a catalytic subunit of (1,3) β -glucan synthase (510). *Neurospora* has only one FKS gene, which encodes a protein of 1955 amino acids. Correspondingly, the *Neurospora rho-1* gene encodes a protein of 195 amino acids. The substrate for (1,3) β -glucan synthase activity is UDP-glucose, which is synthesized from glucose-6-phosphate using two enzymes, phosphoglucomutase and a uridylyl transferase (Fig. 17; Table 51).

There is evidence that in situ, $\beta(1,3)$ -linked glucans branch through (1,6) β -linkages from the main (1,3) β -glucan chain (243). In addition, it is likely that (1,3) β -glucans are crosslinked to chitin (243). Unfortunately, the enzymes responsible

			TABLE	51. Glucan synth	ases		
Enzyme	EC no	NCI 1 no			BLAST	r match	
слауше	EC IIO.	INCO HO.	Best overall	S. cerevisiae	S. pombe	Animal	Plant
(1,3)β-Glucan synthase	2.4.1.34	06871.1	C. immitis FKS/0.0	FKS2; 0.0	NP 588501; 0.0		A. thaliana NP_196804/2e-88
GTPase Rho		08683.1	S. pombe RHO2/5e-81	RHO2; 2.5e-62	RHO2; 2e-81	M. musculus NP_080570/1e-60	I
GTPase Rho		01484.1	A. fumigatus Rho1/1e-32	RHO1; 1.6e-30	RHO1; 2e-32	M. musculus Arha2/1e-28	
Phosphoglucomutase	5.4.2.2	10058.1	A. oryzae PgmA/0.0	PGM2; 2.1e-184	NP_596153; 0.0	X. laevis AAH43876/1e-160	B. napus CAB60109/1e-155
UTP-glucose-1-phosphate uridylyltransferase	2.7.7.9	02797.1	S. pombe NP_588132/0.0	UGP1; 3.4e-170	NP_588132; 0.0	M. musculus AAH26626/1e-138	M. acuminata MWUGPA/1e-128
Killer toxin resistance protein 5		02349.1	P. anserina CAD60785/0.0	KRE5; 1.1e-42	GPT1; 0.0	A. gambiae EAA08752/0.0	A. thaliana NP_177278/0.0
Mannosyl-oligosaccharide glucosidase	3.2.1.106	03657.1	S. pombe NP_594106/0.0	CWH41; 2.2e-88	NP_594106; 0.0	M. musculus Ges1/1e-109	A. thaliana NP_176916/1e-106
Glucan 1,4-α-glucosidase homologue	3.2.1	04203.1	S. pombe NP_593490/0.0	ROT2; 3.6e-195	NP_593490; 0.0	M. musculus BAC27099/0.0	A. thaliana NP_201189/0.0
Glucan 1,4-α-glucosidase homologue	3.2.1	04674.1	A. nidulans AgdB/0.0	ROT2; 5.9e-68	NP_593996; 1e-163	R. norvegicus SI/1e-87	P. pinaster XYL1/1e-142
Calnexin precursor	None	09265.1	A. niger ClxA/1e-168	CNE1; 2.2e-40	CAL1; 1e-135	H. sapiens CANX/9e-86	H. tuberosus CAA84491/1e-80
Killer toxin resistance protein 1?	None	05229.1		WSC4; 1.8e-10	NP_588031; 3e-14		
Ketol isomerase	2.6.1.16	07366.1	S. castellii GFA1/0.0	GFA1; 3.2e-236	NP_596011; 0.0	M. musculus Gfpt1/0.0	A. thaliana NP_189051/0.0
Glucosamine-phosphate-N-acetyltransferase	2.3.1.4	01902.1	C. albicans GNA1/1e-35	GNA1; 7.2e-26	GNA1; 2e-20	C. elegans Gna-1/6e-24	A. thaliana NP_197081/1e-24
Phosphoacetylglucosamine mutase	5.4.2.3	07458.1	C. albicans AGM1/1e-126	PCM1; 7.2e-81	NP_592933; 1e-116	D. melanogaster NP_648588/1e-120	A. thaliana BAB9465/1e-102
UDP-N-acetylglucosamine pyrophosphorylase	2.7.7.23	02109.1	C. albicans UAP1/1e-134	UAP1; 5.2e-110	NP_596832; 7e-91	D. rerio AAH44137/1e-118	A. thaliana NP_181047/7e-97
Phosphoglucose isomerase	5.3.1.9	07281.1	A. oryzae PgiA/0.0	PGI1; 1.4e-196	PGI1; 0.0	M. cephalus Pgi-1/1e-173	
Isocitrate lyase	4.1.3.1	04230.1	M. grisea Icl1/0.0	ICL1; 1.2e-165	NP_595067; 1e-90		I. batatas ICL/1e-166
Malate synthase	4.1.3.2	10007.1	A. nidulans ACUE/0.0	MLS2; 6.7e-181		C. elegans Gei-7/1e-150	G. hirstum CAA36546/1e-143

for branch synthesis or for chitin-glucan cross-linking have not

GENOME ANALYSIS OF NEUROSPORA

75

been identified with certainty in any fungal organism. (ii) $(1,6)\beta$ -Glucan synthesis. $(1,6)\beta$ -Linked glucans play a pivotal role in cell wall assembly in yeast, where they interconnect all other wall components into a lattice (424, 479). Genetic analysis of mutants defective in $(1,6)\beta$ -glucan synthesis has revealed the involvement of a number of gene products that are localized throughout the secretory pathway and at the cell surface (94, 110, 536, 720, 721, 739). In sharp contrast, $(1,6)\beta$ -linked glucans are not found in a number of filamentous fungi, including Neurospora and A. fumigatus (243). In addition, analysis of the Neurospora genome has indicated the presence of homologues of only 4 of the 10 genes involved in $(1,6)\beta$ -glucan synthesis. For example, all of the S. cerevisiae proteins that function in the ER, Cwh41p, Rot2p, Kre5p, and Cne1p, are present in Neurospora. Kre11p, which appears to be involved in the transport of nascent $(1,6)\beta$ -glucans from the ER to the Golgi, is present only in S. cerevisiae and not in Neurospora (or in A. fumigatus). The Golgi genes, KRE6 and SKN1, are present in S. cerevisiae, A. nidulans, and A. fumigatus but not Neurospora. None of yeast cell surface genes, KRE9, KNH1, and KRE1 are present in Neurospora.

These results indicate that cell wall synthesis may be fundamentally different in filamentous fungi and *S. cerevisiae* and that caution should be exercised when using budding yeast as a model for cell wall assembly in filamentous fungi.

Chitin substrate synthesis-the Leloir pathway. The substrate for each chitin synthase isozyme is UDP-N-acetylglucosamine (UDP-GlcNAc), which is synthesized from fructose-6-phosphate by the Leloir pathway (Fig. 17; Table 51) (466, 529). The first pathway-specific enzyme is a ketol-isomerase that synthesizes glucosamine-6-phosphate (GlcN-6-P) and glutamate from glutamine and fructose-6-phosphate. The Neurospora enzyme has been studied in detail and is feedback inhibited by the end product of the pathway, UDP-GlcNAc (221). The second pathway-specific enzyme (the acetyltransferase) forms N-acetylglucosamine-6-phosphate and CoA from GlcN-6-P and acetyl-CoA. The third enzyme, the mutase, forms Nacetylglucosamine-1-phosphate (GlcNAc-1-P) from GlcNAc-6-P. The final enzyme, the pyrophosphorylase, catalyzes the formation of UDP-GlcNAc and PP_i from UTP and GlcNAc-1-P. Neurospora has only a single gene for each enzyme of the pathway, similar to the situation in budding yeast and other filamentous fungi.

Cell wall precursors. An overview of the metabolic machinery to synthesize the precursors for the cell wall polymer synthases is shown in Fig. 17. As indicated, the substrates for $(1,3)\beta$ -glucan and chitin synthase are synthesized from hexosephosphate precursors. Interestingly, fungi, including *Neurospora*, grow on a number of suboptimal carbon sources, including lipids. Under these conditions, *Neurospora* utilizes the glyoxylate shunt (428) to form the sugar substrates required for carbohydrate wall assembly. The enzymes of the shunt—isocitrate lyase and malate synthase—are similar to those from *S. cerevisiae* and are present in single copy in *Neurospora*. The genes encoding each of these enzymes are presented in Table 51.

Chitin synthases. In their pioneering work, Galzer and Brown (284) utilized *Neurospora* when first describing cell-free chitin synthase activity. *Neurospora* has since been the organ-

	NCU		BLAST m	atch		
Enzyme	no.	Best overall	S. cerevisiae	S. pombe	Animal ^a	Plant ^b
Chitin-UDP acetylglucosaminyl transferase	03611.1	Emericella nidulans ChsB/0.00	e-151	e-160	e-22; D. melanogaster	No hits
Chitin-UDP acetylglucosaminyl transferase	05239.1	E. nidulans ChsA/0.00	0.00	0.00	5e-17; C. elegans	No hits
Chitin-UDP acetylglucosaminyl transferase	04251.1	Exophiala dermatitidis CHS2/0.00	0.00	0.00	5e-15; C. elegans	No hits
Chitin-UDP acetylglucosaminyl transferase	09324.1	Magnaporthe grisea CHS4/0.00	0.00	e-13	5e-16; D. melanogaster	No hits
Chitin-UDP acetylglucosaminyl transferase	04350.1	Ustilago maydis CHS6/0.00	e-125	5e-10	4e-20; D. melanogaster	e-08 ^c
Chitin-UDP acetylglucosaminyl transferase	04352.1 ^d	U. maydis CHS6/0.00	e-122	6e-11	2e-21; D. melanogaster/ C. elegans	3e-04
Chitin-UDP acetylglucosaminyl transferase	05268.1	A. fumigatus CHSD/0.0	2e-13	e-05	e-10: C. elegans	No hits
Chitin synthase associated	07435.1	S. pombe CHS5/e-64	2e-63	e-64	2e-10; C. elegans	No hits
Chitin synthase associated	04511.1	S. pombe Hypo./e-140	2e-65 ^e (Bud7p)	e-140	No hits	No hits
Chitin synthase associated	05720.1	S. cerevisiae chs7p/e-72	e-72	No hits	No hits	No hits

TABLE 52. Neurospora chitin synthase components

^a Caenorhabditis elegans, Drosophila melanogaster, Mus musculus, or Homo sapiens.

^b Arabidopsis thaliana.

^c Based on the myosin component of the protein.

^d Based on manual annotation, this predicted polypeptide spans contig 3.225 nucleotides 69201 to 72540 and encodes a 1,112-amino-acid polypeptide (in contrast to the 143-amino-acid prediction in the *Neurospora* database).

^e Even though chs6p was used to identify NCU04511.1, when the latter was used as a query on the yeast database, bud7p was the closest match.

ism of choice in many of the significant phases of progress made in the research of chitin synthesis, including the description of chitosomes and subsequent localization of chitin synthase in vesicular organelles (735), the cloning of a first chitin synthase from a filamentous fungus (878), and the use of partial chitin synthase gene sequences as a phylogenetic tool (127).

As has been determined for a variety of fungi, *Neurospora* has multiple chitin synthase-encoding genes, some of which have been functionally analyzed (74–76, 878). Based on the analyses performed to date, it appears that there is at least some redundancy in chitin synthase-encoding genes in *Neurospora*. Such redundancies have also been demonstrated in other filamentous fungi. Based on the full *Neurospora* genome sequence, there are three additional chitin synthase-encoding genes (other than the four mentioned above), including NCU04350.1, which has an apparent myosin motor-like domain (Table 52) (263). This suggests that at least in some instances there may be a direct association between a chitin synthase and cytoskeletal elements. Similarly, seven chitin synthases have been identified in *A. fumigatus*.

In S. cerevisiae, several other genes have been associated with chitin biosynthesis (although their products probably do not catalyze the reaction involving the polymerization of the UDP-GlcNAc substrate into chitin). Interestingly, the S. cerevisiae Sbe2p and Sbe22p proteins (677), which have been implicated in trafficking of chitin synthases (and perhaps other components of the cell wall biosynthetic machinery), do not have apparent homologues in Neurospora. However, Chs5p, Chs6p, and Chs7p, most probably involved in the budding yeast Chs3p transport and/or activation (129, 678, 797, 901), have apparent homologues in Neurospora (Table 52). Both the similarities and the differences between the S. cerevisiae and the Neurospora chitin synthase-related machinery are intriguing. Similar catalytic components are present in both (even though there are more such components in the filamentous fungus, perhaps due to its morphological complexity). In contrast, a more pronounced divergence is apparent in at least one of the additional components involved in regulation of chitin synthase activity. This, again, could be linked with the significant

differences in the morphology of the different species (which may require different modes of regulation). Another possibility is that these components play a secondary role in *S. cerevisiae* chitin synthesis, even though they have been associated with the cell wall biosynthetic process. The notion that these proteins are involved in other functions is supported by experimental evidence (678). Interestingly, higher eukaryotes that produce chitin (e.g., *C. elegans* and *D. melanogaster*) appear to contain fewer members of the chitin synthase gene family (Table 52). Whether chitin biosynthesis in these (and perhaps similar) organisms has been streamlined or whether additional, as yet unidentified, components have altered the way in which chitin production is regulated is yet to be determined.

As expected, the fact that plants and mammals lack chitin is also evident from the absence of structural chitin synthase homologues. The unique composition of the fungal cell wall has been a favored target in the continuous search for antifungal compounds (279, 293, 325). Even though direct and indirect inhibitors of chitin synthesis have been identified, their use as commercial antifungals has been limited; this may be due to the combination of their pharmacological properties and the potential difficulties in inhibiting a process that can apparently be performed by several enzymes in a compensatory manner. However, the recent commercial introduction of glucan synthase inhibitors is a clear indication that cell wall components are valid and potentially rewarding targets for the development of antifungal compounds (see also "Relationship to animal and plant pathogens" below).

Hyphal Morphogenesis

Determining and maintaining cell shape is a fundamental prerequisite for proper development of any organism. The defining characteristic of filamentous fungi is the development of hyphae, tip-growing cellular elements that undergo regular branching, exhibit high developmental versatility, and respond to a myriad of signals during their invasive and exploratory growth within the natural environment. Hyphal compartments are frequently multinucleate, and movement of organelles between compartments is facilitated by the existence of incom-

S. cerevisiae GO ^b term	Orthologues found in <i>Neurospora</i> (homology over >60% of protein length)	Domains conserved (e<-10)	No matches found
Establishment of polarity	ABP1 ⁺⁺ , ACT1 ⁺⁺ , BCK1 [*] , BEM1°, BN11 ⁺⁺ , BUD6 ⁺⁺ , CDC3 ^{**} , CDC10 ^{**} , CDC11 ^{**} , CDC12 ^{**} , CDC24°, CDC42°, CKA1 [*] , CKA2 [*] , CKB1 [*] , CKB2 [*] , CLA4 [*] , EXO70°, EXO84°, MLC1, PWP2, RGA1°, RGA2°, RHO1°, RHO2°, RHO3°, RHO4°, ROM1°, ROM2°, SEC3°°, SEC5°°, SEC6°°, SEC8°°, SEC10°°, SEC15°°, MYO2 ⁺⁺ , TPM1 ⁺⁺ , TPM2 ⁺⁺	BEM2°, BEM3°, BOI1, BOI2, CDC43, MSB1, SHS1**, SPA2°,	BEM4°, GIC1°, GIC2°, PEA2°, LAS1, MSB2, SPH1°, SLG1°, ZDS1°, ZDS2°
Exocyst	EXO70°°, EXO84°°, SEC3°°, SEC5°°, SEC6°°, SEC8°°, SEC10°°, SEC15°°		
Polarisome	BNI1 ⁺⁺ , BUD6 ⁺⁺	SPA2°	PEA2°, SPH1°
Bud site selection	AXL1, BUD6, BUD7, BUD23, BUD31, BUD32, RSR1°, STE20*	BUD10, BUD2°, BUD4, BUD5°, BUD13, BUD14, BUD20, RAX1, RAX2	BUD3, BUD8, BUD9, BUD16, BUD17, BUD19, BUD22, BUD25, BUD26, BUD27, BUD28, BUD29, BUD30, HKR1, YOR300W
Axial budding	AXL1, CDC3**, CDC10**, CDC11**, CDC12**, ERV14°°, MYO1 ⁺⁺ , PFY1 ⁺⁺ , RSR1°	BUD10, BNR1 ⁺⁺ , BUD2°, BUD4, BUD5°, ELM1*, GIN4°, KCC4°, PAN1 ⁺⁺	BUD3, GIC1°, GIC2°
Invasive growth	CDC24°, CDC42°, RGA1°, RGA2°	BEM3°, DIA3, DIA4, RIM20, SPT3 ⁺ , STE12 ⁺	DFG16, DIA1, DIA2, DIG1 ⁺ , DIG2 ⁺ , FLO8 ⁺ , MUC1 ⁺ , NRG1 ⁺ , NRG2 ⁺ , RIM8, RIM21, RXT2
Pseudohyphal growth	BCY1*, BMH1, BMH2, CDC24°, CDC42°, CDC55*, DFG5, GPA2*, KSS1*, MEP2, PGU1, PLC1*, RAS2°, RGA1°, RGA2°, SHO1*, STE7*, STE11*, STE20*, TPK1*, TPK2*, TPK3*	BEM3°, BUD5°, DFG10, DIA3, DIA4, ELM1*, FKH1 ⁺ , FKH2 ⁺ , PHD1 ⁺ , SOK2 ⁺ , SPA2°, STE12 ⁺	ASH1 ⁺ , BUD8, DIA1, ECM23, FLO8 ⁺ , GPR1 [*] , HMS1 ⁺ , HMS2 ⁺ , MSS11 ⁺ , MUC1 ⁺ , PAM1, TEC1 ⁺ , SPH1°, SRO9°

TABLE 53. Classification of proteins important for cell polarity development^a

*a**, signaling components; **, septation machinery; °, Rho- and Ras-type GTPases modules and interacting proteins; °°, secretory pathway;⁺, transcriptional regulators;⁺⁺, actin cytoskeleton.

^b GO, gene ontology.

plete cross walls. Over the last few years, evidence has accumulated that, similar to cells of higher eukaryotes, the microtubule cytoskeleton provides the structural basis for the longdistance vectorial transport of secretory vesicles toward the growing hyphal tip (464, 701). Secretory vesicles then fuse with the tip to deliver membrane and materials required for continuous cell wall synthesis. This is thought to be coordinated by the Spitzenkörper, a fungal-specific organelle assemblage which is localized in the hyphal apex and serves as a vesicle supply center (49, 283). Through its microtubule-dependent movement and positioning within the hyphal apex, this organelle complex determines the shape and growth directionality of the hypha (48, 656). New hyphal tips are generated in subapical regions by branching, which requires some ill-defined signal(s) to establish the site of the new bud emergence and to regulate the spacing of branch points along the hypha.

Generation of hyphal polarity. On a molecular level, the best-described example of polar growth is the budding yeast *S. cerevisiae*. Many genes required for cellular development have been identified and arranged into functional hirarchies in this organism (203, 632, 633). This interaction map can be used as a starting point for the analysis of similarities and differences that control the different morphologies of filamentous fungi and unicellular yeasts. In *S. cerevisiae*, polarized growth is mediated by a series of steps, including the action of cortical landmark proteins, Rho- and Ras-type GTPases, that polarize the actin cytoskeleton and direct the motor-driven transport of secretory vesicles and cell wall components to the site of growth (566). One key component for the establishment of cortical landmarks that determine the future bud site is the

small GTPase Rsr1p, which acts in combination with information from the previous division site. Additional factors, such as the septins, several other Bud proteins, and the exocyst component Sec3p, act in combination with the DNA content of the cell and the available growth conditions to further specify an axial or bipolar budding pattern (45, 235). Following bud site establishment, Cdc42p-dependent organization of the actin cytoskeleton and the recruitment of the budding machinery occur, finally leading to actin-dependent targeted secretion and cell wall formation at the site of polarization (summarized in reference 236).

(i) Proteins important for cell polarity development. To determine the conservation of components that constitute the morphogenetic network leading to polarized growth in S. cerevisiae, all proteins that are associated with polarity-related gene ontology (GO) terms at the Saccharomyces Genome Database were compared with the Neurospora genome (Table 53). Most proteins that represent the core machinery of cellular morphogenesis, such as components of cAMP and MAPK signaling pathways, Ras- and Rho-type GTPases, proteins that are necessary for the coordinated polarization and organization of the actin cytoskeleton and the secretory pathway, and proteins that constitute the septation machinery, are present in the Neurospora genome. Little is known about their function in Neuropora or other filamentous fungi, and experiments are needed to clarify their role in filamentous growth. However, in addition to this highly conserved core machinery that governs cellular polarity, other genes that modulate the different morphologies of S. cerevisiae (e.g., establishment of different budding patterns or pseudohyphal/invasive growth) are either absent from or highly diverged in *Neurospora*. In contrast to the true hyphae of filamentous fungi, the pseudohyphae produced by yeast under nitrogen starvation conditions are the result of unipolar budding that leads to chains of uninucleate elongated cells with no apparent communication between cellular compartments (540). Although the two signaling pathways that regulate pseudohyphal differentiation and invasive growth in *S. cerevisiae*—the MAPK and cAMP modules—are conserved among yeast and filamentous fungi, the key transcription factors Flo8p, Muc1p, and Tec1p, as well as other transcriptional regulators that are implicated in the switch from budding to pseudohyphal growth (Dig1p, Dig2p, Hms1p, Hms2p, Nrg1p, Nrg2p, and Mss11p), are not detectable in the *Neurospora* genome. Thus, different sets of transcription factors appear to regulate true hyphal versus pseudohyphal development.

In its budding mode, S. cerevisiae can exist in different forms, each with a specific cell morphology and cell division pattern. The typical yeast cell is ellipsoid, and haploid cells bud in an axial pattern with the bud formed next to the preceding site of cytokinesis. However, in diploid yeast cells, daughter cells bud 180° from their birth site in a bipolar manner. Interestingly, most BUD proteins that are specifically involved in the generation of these different budding patterns are not conserved in Neurospora (Table 53), suggesting that novel (and to date unknown) mechanisms have been developed in filamentous fungi to define the sites of the new hyphal tip emergence and to regulate spacing of branch points along the hypha. Also of interest is that the genes encoding several proteins that bind to Rho-type GTPases in budding yeast (BEM4, GIC1, GIC2, NIP100, and ZDS2) or act as downstream effectors of Rho proteins (PEA1, SLG1, SPH1, WSC1, WSC2, and WSC3) either are not found in the *Neurospora* genome or are highly diverged.

(ii) Rho-type GTPases as key regulators of polarity. Rhotype GTPases are molecular switches that cycle between an active (GTP bound) and an inactive form (GDP bound). Transition between these two forms is achieved through GTPaseactivating proteins (GAPs), leading to the inactive form, and GDP-GTP exchange factors (GEFs), that activate the small G protein. Originally, Rho proteins were described as key regulators of the actin cytoskeleton, but now it has been shown that they influence an amazing variety of cellular processes that are crucial for coordinated morphogenesis (for reviews, see references 224 and 820). Therefore, a comparison of the *Neurospora* set of these master regulators of polarity with their budding yeast orthologues is required to elucidate similarities and differences that may contribute to the different morphologies in the two organisms.

A phylogenetic analysis shows that *Neurospora* Rho proteins and available fungal sequences of Rho1, Rho2, Rho3, and CDC42 fall into distinct subgroups (Fig. 18; Table 54), and experimental data from *S. cerevisiae* and other filamentous fungi suggest that there is at least a central set of conserved functions for these proteins. *S. cerevisiae* Rho1p regulates the organization of the actin cytoskeleton and is also required to maintain cell wall stability. This is achieved through two independent mechanisms. Rho1p activates cell wall synthesis through the activation of the MAPK pathway, which monitors cell wall integrity, and it is also directly required to stimulate glucan synthase activity, which catalyzes the synthesis of the main structural component of the yeast cell wall. Similarly,



FIG. 18. Phylogenetic tree of *Neurospora* Rho proteins. Rho proteins were manually annotated and analyzed using Clustal W. The numbers adjacent to nodes indicate the percentages of 1,000 additional bootstrap trials in which the indicated protein groups were found.

Rho1 was also found to be part of the glucan synthase complex that localizes to zones of active growth at the hyphal apex in A. fumigatus (57). A role for cell wall integrity was also suggested for Rho1 of Ashbya gossipii, since deletion mutants show reduced filamentous growth and a high rate of lysis (848). Deletion of the A. gossipii gene encoding Bem2 (which was suggested to act as a Rho1-specific GAP in this fungus) resulted in aberrant switching from isotropic to polar growth during germination and maintenance of polarity during hyphal elongation (849). Areas of lost polarity showed a uniform actin distribution, suggesting that hyperactivity of Rho1 may result in depolarization of the actin cytoskeleton. Rho3p is important for coordinated polarization of the actin cytoskeleton and the secretory apparatus in S. cerevisiae. Trichoderma reseei rho3 can complement the corresponding yeast mutant, and overexpression of this protein can also suppress several late-acting secretory pathway mutations, suggesting conserved functions (829). Surprisingly, deletion of T. reseei rho3 shows no morphological defects, while deletion of RHO3 is lethal in yeast. This example

Neurospora protein	(Drthologue(s) ^a	S. cerevisiae	Proposed role in	Interacting Rho
<i>Neurospora</i> protein	S. cerevisiae	S. pombe	conserved	filamentous fungi	S. cerevisiae
Rho proteins					
NCU01484.1/Rho1	Rho1p	Rho1; SpAC20H4.11C		Part of $\beta(1,3)$ -glucan synthase complex, regulation of actin organization and cell wall integrity	
NCU08683.1/Rho2	Rho2p	Rho2			
NCU0600.1/Rho3	Rho3p	SpAC23C4.08		Interaction with secretory pathway, main- tenance of polarity	
NCU03407.1/Rho4	Rho4p	SpAC16A10.04		I I I I	
NCU02160.1/Rac	1	1		Actin organization	
NCU06454.1/CDC42	Cdc42p	Cdc42		Essential for establishment and mainte- nance of polar growth	
NCU03346.1					
Regulators of Rho proteins RhoGAP proteins NCU02689.1 NCU02524.1 NCU00553.1	Lrg1p Rgd1p		Bem2/3		Cdc42p, Rho1p, Rho2p Cdc42p, Rho1p Rho3p Rho4p
NCU00196.1	nguip		Bag7/Sac7		Rho1p, Rho2p
NCU07688.1 NCU09537.1 NCU02915.1 NCU07622.1	Rga1p/2p Rgd2p				Cdc42p, Rho1p Cdc42p, Rho5p
RhoGEF proteins NCU06067.1 NCU00668.1 NCU02131.1 NCU06579.1 NCU02764.1	Cdc24p Rom1p/2p Tus1p				Cdc42p Rho1p Rho1p

TABLE 54.	Rho G	TPase	modules	in	Neurospora
-----------	-------	-------	---------	----	------------

^a Domain structure conserved and homology over >60% of protein length.

clearly illustrates that although it can be assumed that core functions of the Rho GTPases are conserved between yeast and filamentous fungi, a detailed analysis is necessary to clarify their exact roles during hyphal morphogenesis.

For the CDC42 module, several mutants are available in filamentous fungi. In A. gossipii, loss of CDC42 as well as CDC24 (acting as a GEF for Cdc42p) leads to apolar growth of the spore and inability to initiate polarized growth during germination (848). Experiments with dominant-negative and dominant-active mutants in Penicillium marneffei indicate that CDC42 is essential to maintain polarity during hyphal elongation in addition to establishing polarity during germination (101). No functional data are available for Rho2 and Rho4 in Neurospora or other filamentous fungi. In S. cerevisiae, these proteins appear to function in a partially redundant manner with Rho1p and Rho3p, respectively. While members of the Rho2 group are evolutionarily conserved, the available sequences constituting the Rho4 class are quite strongly diverged (Fig. 18), making it difficult to predict potential functions. Similarly, S. cerevisiae Rho5p is an orphan protein that cannot be correlated to any other available fungal Rho protein.

In addition to these five Rho proteins (Rho1 to Rho4 and CDC42), the *Neurospora* genome contains a clear ortholog of Rac that is conserved in filamentous fungi and higher eukaryotes but is missing from all sequenced yeasts (*S. cerevisiae*, *S. pombe*, and *C. albicans*). Interestingly, this type of Rho protein is especially important for the growth and development of polarity in neuronal cells and may play a similar role in the

filamentous fungi. Although the suggestion that the hyphal cell is a simpler version of a neuron may seem far-fetched, several examples from the area of motor proteins and organelle transport support this view (343, 421, 538, 700, 831). Interestingly, several potential Rac homologs are present in the dimorphic fungus Yarrowia lipoytica (357). The deletion of one Rac protein revealed that it is not an essential gene. Its loss does not impair actin organization in Y. lipoytica cells but does prevent the switch to filamentous growth, indicating an important function during hyphal morphogenesis. On the other hand, in P. marneffei, Rac colocalizes with actin at the tips of vegetative hyphal cells and at septa (102). Loss of Rac in this organism results in growth defects in both vegetative hyphal and conidiophore cell types, such that cells become depolarized and the actin cytoskeleton is severely disrupted. These data suggest that Rac proteins can play a crucial role in actin-dependent polarized growth and division. Finally, the Neurospora genome contains an unusual protein with a domain that has weak homology to Rho in filopodia (NCU03346.1). Orthologues of this unusual protein are also found in the genome sequences of M. grisea and A. fumigatus, implying that this protein has a function that is unique for filamentous fungi; however, no functional data are currently available.

A major complication in determining the functions of small G proteins is that the number of GAPs and GEFs that have been uncovered in the sequencing projects far outnumber the GTPases that they regulate (e.g., the *D. melanogaster* genome contains only six Rho proteins but about 20 GEFs and more

than 20 GAPs [5]). This is also true for *Neurospora*, where the GAP and GEF repertoires are both expanded relative to budding yeast (especially if *S. cerevisiae* paralogues are not considered in the comparison [Table 54]). The task for the future will be to elucidate the regulatory networks for all the Rho proteins and their regulators and to determine how the different pathways interact to generate a well-shaped hypha.

Cytoskeleton and motor proteins. Mechanochemical enzymes responsible for intracellular cytoskeletal transport can be grouped into three superfamilies: the microtubule-based kinesins and dyneins and the actin-associated myosins (341, 342, 413). Based on the sequence similarity of their ATPhydrolyzing motor domains, several families or classes within each superfamily can be defined that often correlate with conserved cellular functions. The genome of the unicellular yeast S. cerevisiae encodes six kinesins, one cytoplasmic dynein, and five myosins. In contrast, mammalian genomes encode more than 50 distinct kinesins, four functional classes of dyneins, and over 40 distinct myosins (31, 623). These numbers suggest that as eukaryotic cells increase in volume and morphological complexity, there is a corresponding increase in the complexity of cytoskeletal transport systems. This is borne out in our analysis of Neurospora, with an elaboration of certain cytoskeletal components reflecting the more complex morphology of Neurospora than of S. cerevisiae (Table 55).

(i) Structural components. The main structural components of the cytoskeleton are actin filaments, microtubules, and intermediate filaments. The Neurospora genome did not offer any major surprises at the level of the structural components of the cytoskeleton. All S. cerevisiae genes coding for the actin and microtubule cytoskeleton as well as actin binding and microtubule-associated proteins are strongly conserved in Neurospora (204, 434, 688). Ten actin-related proteins (Arps) have been found in S. cerevisiae (684), and Neurospora encodes all but Arp7p and Arp9p, which are involved in chromatin remodeling (120). The Arp2p-Arp3p complex and formins participate in two independent actin nucleation pathways (634). The components of the Arp2p-Arp3p complex are well conserved in Neurospora; however, Neurospora encodes only a single formin, while S. cerevisiae and S. pombe contain two and three, respectively. In addition to proteins required for actin and microtubule formation and function, an ortholog of S. cerevisiae Mdm1p was identified in the Neurospora genome. Mdm1p is important for mitochondrial and nuclear distribution in S. cerevisiae (512) and shows sequence similarity to vimentin and keratin, suggesting that an intermediate filament system is a universal component of eukaryotic cells.

(ii) Kinesins. The extremely high growth rate (>1 μ m/s) and highly polar form of *Neurospora* requires that a large amount of material for cell wall synthesis, as well as various organelles, be transported toward hyphal tips. Intracellular transport in budding yeast is exclusively myosin dependent. In contrast, filamentous fungi, similar to higher eukaryotes, utilize a combination of actin- and microtubule-based systems.

Examination of the *Neurospora* genome indicates that it encodes 10 distinct kinesins, of which 4 are likely to be involved in cytoplasmic transport (Table 55). Conventional kinesin purified from animal sources is the founding member of the kinesin superfamily and is involved in a wide spectrum of cytoplasmic transport processes (401, 831). A relative of ani-

mal conventional kinesin, Nkin, was first identified and characterized in Neurospora (756), but additional work has shown that members of this family are present in all filamentous fungi examined to date (464, 648, 755, 862). Nkin was proposed to be involved in the transport of secretory vesicles toward the growing tip in Neurospora (701, 702), but the analysis of orthologs in other filamentous fungi suggested that in addition to polarized secretion, conventional kinesin might be necessary for the organization of vacuoles and could affect microtubule dynamics (648, 757). These results argue for multiple functions of a single motor protein in filamentous fungi, similar to what has been described for animal kinesins. Mutational analysis of Nkin has identified regions of cargo association and regulation of the ATPase activity that are conserved between fungal and animal kinesins (421, 700). Interestingly, fungal conventional kinesins lack copurifying light chains (422), which, in addition to the C terminus of the motor protein itself, were proposed to function in cargo attachment and motor activation in animals (830), suggesting that this fungal motor could serve as a simplified model to study motor-cargo interaction and its regulation.

Additional *Neurospora* kinesins implicated in organelle transport include two of the unc104 family and one related to KIF21A. Both families are involved in the transport of a variety of cytoplasmic cargoes in metazoan systems (84, 502). Interestingly, members of the KIF21 group are restricted to filamentous fungi and higher eukaryotes and not present in unicellular yeasts, suggesting a potential role in long-range transport processes. KIF21 orthologs were also found in *U. maydis* and the thermophilic fungus *Thermomyces lanuginosus* (670, 846), suggesting that this family may be widely used for vesicular transport in filamentous fungi.

In S. cerevisiae, five kinesin motors are required to build up counteracting forces to organize the mitotic and meiotic spindle. The C-terminal motor Kar3p, the functionally redundant BimC family members Kip1p and Cin8p, and the proteins Kip2p and Kip3p, which have at least partly overlapping functions, function together with cytoplasmic dynein for spindle assembly and chromosome segregation (summarized in reference 338). In contrast to both budding and fission yeast, Neurospora is more streamlined and has a minimal set of only one mitotic kinesin per subfamily with no apparent overlap in function. The orthologues of NCU04581.1 and NCU00927.1 have been studied in A. nidulans (KlpA and BimC, respectively [573]), and the results suggest the existence of conserved functions between yeast, filamentous fungi, and higher eukaryotes for these proteins (for a detailed analysis of mitotic motors in filamentous fungi, see reference 12).

In addition to these minimal components, the *Neurospora* genome contains two other kinesins that are known to be involved in mitosis in metazoan cells (NCU05180.1 and NCU05028.1 [6 and 265]) are not found in the *S. cerevisiae* genome. A possible function for these motor proteins during fungal mitosis remains to be determined, but their existence suggests mechanistic similarities in spindle formation and function between animals and filamentous fungi that are not shared with unicellular yeasts. This is also reflected by the absence (Spc25p, Spc29p, Spc34p, Spc42p, and Ndc1p) or high divergence (Bbp1p, Cnm1p, Stu1p, Spc24p, Spc72p, Spc105p, and Spc110p) of components of the *S. cerevisiae* spindle pole body

		Orthologue(s)		
NCU no.	S. cerevisiae	S. pombe	Family/class	Proposed role(s)
Kinesins 09730.1	Smy1p	Klp3 (SPAC1834.07)	Conventional kinesin	Transport of secretory(?) vesicles, nuclear
06733.1	NF^a	NF SPAC144.14	Unc104	Vesicular transport
06832.1 04581.1	NF NF Kar3p	NF Pkl1/Klp1 (SPAC3A11.14C)	Kif21/chromokinesin C-terminal	Vesicular transport, DNA binding Dyanamics if spindle microtubules, counteracts BimC-like motors
06144.1	Kip3p	Klp2 (SPAC664.10) Klp5 (SPBC2F12.13)	Kip3	Spindle positioning, spindle elongation during anaphase, microtubule disassembly
00927.1	Kip1p, Cin8p	Klp6 (SPBC649.01C) Cut7 (SPAC25G10.07C)	BimC/Eg5	Spindle assembly and centrosome separation
05180.1	NF	SPBC15D4.01C (also named	Fast evolving/pavarotti	Organization of the mitotic spindle
02626.1	Kip2p	Klp4/Tea2 (SPBC1604.20C)	Kip2	Heterogenous group: Kip2p has mitotic functions (partly overlapping with Kip3p), while Tea2 seems to alter the dynamics of interphase microtubules
05028.1	NF	NF	KID	Chromosome alignment in metaphase
Myosins				
01440.1	Myo2p, Myo4p	Myo5 (SPCC1919.10c) Myo4 (SPBC2D10.14c)	Class V	Organelle transport
02111.1	Myo3p, Myo5p	Myo1 (SPBC146.13c)	Class I	Endo-/exocytosis
00551.1	Myo1p	Myo2 (SPCC645.05c) Myo3 (SPAC4A8.05c)	Class II	Actin organisation, cytokinesis
04350.1	NF	NF	Chitin sythase-myosin fusion protein	Specific for filamentous fungi
Dynein subunits				Nuclear movement, spindle elongation, retrograde vesicle transport
06976.1	Dyn1p	Dhc1	Dynein heavy chain	
09142.1	Pac11p	SPBC646.17	Dynein intermediate chain	
09982.1	NF Dyn2n	NF $D1_{2}$ (SPAC026.07)	Dynein light intermediate chain	
03882 1	NF	Dlc (SPAC1805.08)	Dynein light chain, ECo	
09095.1	NF	NF	Dynein light chain, LC7	
Dynactin subunits				Dynein-cargo interaction, nuclear movement, spindle elongation, retrograde vesicle transport
03483.1	NIP100p	NF	Dynactin p150 ^{Glued}	
00257.1	NF	NF	Dynactin p62	
03563.1	NF	NF	Dynactin Arp11	
08375.1	NF Amula	NF Actin like metain	Dynactin p50/dynamitin	
04247.1	Arp1p	(SPBC1347.12)		
04043.1	NF	NF	Dynactin p27	
Not defined	NF NF	NF NF	Dynactin p25 Dynactin p24	
Lis-1 complex				Dynein regulation, nuclear movement, spindle
04534.1	Pac1p	NF	LIS1	ciongation, retrograde vesicle transport
04312.1	Pac1p	NF	LIS1	
08566.1	NF	NF	NUDE/RO11	

TADIT	_	_	~	•	c .	C 1		
		<u> </u>	1 om	DOPICOD.	ot.	tungol	motor	protoinc
LADLEY	,		A CHILL	JALISUIL	())	Innigat	111021021	DIDICHIS
	~	~ •	~~~		~	I CHIIMCOI		proteino.

^a NF, not found by conventional BLAST searches.

compared to other organisms. Especially interesting is one kinesin (NCU05180.1) that shows only weak sequence conservation in the normally well-conserved motor domain. The *D. melanogaster* orthologue of this kinesin ("pavarotti") was originally identified in a screen for fast-evolving proteins by cross-hybridization with *D. virilis* and *D. yakuba* cDNA libraries (687) and subsequently shown to act in mitosis. It seems that

their fast-evolving nature and not their sequence similarity is a key feature of this subfamily. NCU02626.1 is part of a heterogenous subfamily of kinesins that is defined by a centrally located motor domain; however, sequence conservation in this subfamily is restricted to the motor domain. The *S. cerevisiae* member of this subfamily, Kip2p, functions in spindle assembly and nuclear positioning (530), while the corresponding *S*. *pombe tea2* mutants display altered microtubule dynamics during interphase (112). *U. maydis kin1* mutants have no discernible phenotype (464).

(iii) Myosins. Neurospora has only four myosins, with a single member in each of the three classes found in S. cerevisiae (Table 55). Analysis of the A. nidulans protein corresponding to the Neurospora class I myosin, MyoA, suggests that it functions in endocytosis and secretion (513, 578, 870). Work with both yeasts and several vertebrate systems suggests that the other two Neurospora myosins (NCU00551.1, class II; NCU01440.1, class V) are probably involved in cytokinesis and organelle transport, respectively (111, 718, 856); however, experimental data are lacking for Neurospora or other filamentous fungi. In addition, Neurospora encodes an unconventional myosin domain linked to a class V chitin synthase domain. This unusual myosin has also been identified in other filamentous fungi and appears to function in cell wall synthesis and maintenance of cell wall integrity (263, 599). Mutational analysis has indicated that both the myosin and chitin synthase domains are required for correct cellular function (349).

(iv) Dynein. Cytoplasmic dynein is the most complex of the motor proteins operating in the cytoplasm (341). Purified mammalian cytoplasmic dynein consists of a heavy chain (>4,000 residues), an intermediate chain, a light intermediate chain, and three distinct light chains (347, 416). An additional multisubunit complex, known as dynactin, is required for all known dynein functions and consists of at least distinct 10 subunits (Table 55) (345). In mammals, cytoplasmic dynein is required for numerous intracellular transport processes; however, in S. cerevisiae, dynein function is restricted to ensuring proper nuclear movement and distribution between mother and daughter cells during cell division. Consistent with this restricted role, S. cerevisiae dynein and dynactin subunits are highly diverged relative to those of higher eukaryotes. In contrast, cytoplasmic dynein in Neurospora is required for retrograde transport of membranous organelles, as well as for nuclear movement (702). Examination of the Neurospora genome shows that dynein/dynactin subunits of Neurospora are more similar to those of metazoans than to those of S. cerevisiae. Some of the dynein/dynactin subunits present in filamentous fungi and metazoans (DLIC, the "roadblock" DLC, p150Glued, dynamitin/p50, p62, p27, and p25) are not detectable in either S. cerevisiae or S. pombe (Table 55). Two of the dynactin subunits, p24/p22 of the shoulder/sidearm subcomplex and Arp11 of the Arp1 pointed-end complex (211, 693), have undergone significant change in all organisms, with clear matches seen only between closely related organisms. Haploinsufficiency of LIS1, a dynein regulator, results in a defect in neuronal migration and subsequent brain development (818). Interestingly, Neurospora appears to possess two LIS1 proteins, while other unicellular and filamentous fungi and metazoans appear to have only one. The significance of this duplication is not known.

Cyclin/CDK Machinery

Oscillations in the activity of cyclin-dependent kinases (Cdk) drive the eukaryotic cell cycle (549, 561). These enzymes are complexes of catalytic (Cdk) and regulatory (cyclin) subunits. In most cases it is cyclin abundance that oscillates in the cell

cycle. In *S. cerevisiae*, the cell cycle alternates between two states, depending on whether mitotic B-type cyclin/Cdk activity is high (in the S, G_2 , and M phases) or low (in G_1). The G_1 cyclins are thought to elevate mitotic cyclin/Cdk activity by promoting the degradation of mitotic cyclin/Cdk inhibitors, whereas the anaphase-promoting complex reduces kinase activity by degrading mitotic cyclins, thereby completing the cycle (549, 561). In many eukaryotes the proteins and pathways involved in cell cycle transitions are highly redundant and have specialized functions. In *S. cerevisiae*, for example, most of the current cyclin genes are derived from gene duplications. Because of RIP (709), we anticipated that *Neurospora* should have fewer, presumably less specialized cyclins than the yeasts or higher eukaryotes do.

In both *S. cerevisiae* and *S. pombe*, there is only one major Cdk responsible for cell cycle transitions (*CDC28* and *cdc2*, respectively), while in humans there are five Cdks with cell cycle roles (550). There are additional Cdks that do not play major roles in cell cycle progression; some of these proteins are involved in phosphate metabolism, transcription, or less well defined processes. *Neurospora*, like *S. cerevisiae*, has a Cdk family that includes a Cdc2-like Cdk (*cdc-2*), a Pho85p-like Cdk, and Cdks likely to be involved in transcription (NCU07172.1, NCU06685.1, and NCU03659.1). It also has a Cdk (NCU07880.1) with a PITSLRE motif, which is absent from *S. cerevisiae* but present in *S. pombe*. We conclude that *Neurospora* has just one Cdk (*cdc-2*) that is primarily responsible for cell cycle progression.

Neurospora has three cyclin genes (Table 56) that are likely to be involved in cell cycle control. By comparison, S. cerevisiae and S. pombe have nine and five cyclins, respectively, involved in cell cycle progression, while humans have four classes of cell cycle cyclins (cyclins A, B, D, and E), with more than one member in each class (550). NCU02114.1 (cln-1) is likely to be a G_1 cyclin because the S. cerevisiae (CLN1 to CLN3) and S. *pombe* (*puc1*) G_1 cyclins are more closely related to it than to the other two Neurospora cyclins (Table 56). In fact, CLN1 and CLN3 show no significant similarity to any other Neurospora cyclin (Table 56). Both NCU02758.1 (clb-1) and NCU01242.1 (clb-3) are B-type cyclins. CLB-1 contains the mitotic destruction motif typical of mitotic cyclins (652). It is related to G_2/M cyclins from S. cerevisiae (CLB1/CLB2) and S. pombe (cdc13) (Table 56) and the NIME B-type cyclin from A. nidulans (574). CLB-3 may play an earlier role-perhaps in S phase-because it is more similar to CLB3/CLB4 from S. cerevisiae and cig1 from S. pombe (Table 56). It appears that A. fumigatus also contains these three cyclins-all closely related to the Neurospora proteins (data not shown). We conclude that the cyclin/ Cdk complexes in Neurospora are very streamlined compared to the better-studied model organisms. However, the Neurospora life cycle is far more complex than that of S. cerevisiae and S. pombe. How Neurospora manages to achieve more complex developmental programs with apparently less diversity in protein complexes is a question worth addressing in future cell cycle studies.

Even with a limited set of cyclins, however, most of the key components of the cell cycle machinery are present in *Neurospora*. Thus, *Neurospora* conforms to the view that the machinery and the "wiring" of processes that regulate cell division are conserved among eukaryotes. An apparent lack of Cdk inhib-

Neurospora loc gei	us vs <i>S. cerevisiae</i> nome	S. cerev	<i>isiae</i> locus vs <i>Neurospora</i> genome	Neurospora lo gen	ocus vs S. pombe	S. por	nbe locus vs Neurospora genome
NCU02114.1	CLB3 (1e-27) CLB4 (5e-26) CLB6 (1e-23)	CLN1	NCU02114.1 (1e-16) NCU01242.1 (>e-5) NCU02758 1 (>e-5)	NCU02114.1	<i>cdc13</i> (4e-32) <i>puc1</i> (1e-31) <i>cig1</i> (2e-30)	puc1	NCU02114.1 (1e-30) NCU01242.1 (8e-18) NCU02758 (5e-16)
	CLN3 (2e-22) CLB2 (3e-22) CLB5 (6e 20)	CLN2	NCU02114.1 (1e-14) NCU02758.1 (1e-08) NCU01242.1 (>e.5)	NCU01242 1	cig2 (3e-26) rem1 (9e-18)	cig1	NCU01242.1 (8e-76) NCU02758 (4e-66) NCU02114 1 (7e 25)
	CLB3 (00-20) CLB1 (1e-19) CLN1 (3e-14) CLN2 (2e-12)	CLN3	NCU02114.1 (9e-15) NCU021242.1 (>e-5) NCU02758.1 (>e-5)	NC001242.1	cig1 (4e-72) cig2 (1e-67) cim1 (3e-47)	cig2	NCU02114.1 (7e-23) NCU02758 (2e-79) NCU01242.1 (1e-65) NCU02114.1 (8e-28)
NCU01242.1	CLB3 (7e-78) CLB4 (6e-75) CLB2 (5e-57)	CLB1	NCU02758.1 (2-53) NCU02758.1 (4e-72) NCU01242.1 (2e-53) NCU02114.1 (3e-19)	NCU02758.1	puc1 (4e-17) cdc13 (2e-94)	rem1	NCU01242.1 (2e-48) NCU02758 (1e-41) NCU02114 1 (7e-19)
	CLB2 (3e-57) CLB1 (5e-53) CLB6 (5e-48) CLB5 (3e-47)	CLB2	NCU02758.1 (2e-85) NCU01242.1 (2e-57) NCU02114.1 (3e-22)		cig1 (4e-64) rem1 (2e-41) nuc1 (3e-15)	cdc13	NCU02758 (1e-101) NCU01242.1 (9e-74) NCU02114 1 (2e-30)
	CLN3 (5e-07) CLN2 (>e-5) CLN1 (>e-5)	CLB3	NCU01242.1 (3e-82) NCU01242.1 (3e-82) NCU02758.1 (7e-67)		puer (30-13)		140002114.1 (20-50)
NCU02758.1	CLB2 (4e-81) CLB1 (1e-74) CLB3 (6e 67)	CLB4	NCU02114.1 (2e-26) NCU01242.1 (8e-80) NCU02758.1 (1e-65) NCU02114.1 (2e-25)				
	CLB5 (6e-67) CLB4 (2e-62) CLB6 (3e-58)	CLB5	NCU02114.1 (3e-23) NCU02758.1 (3e-51) NCU01242.1 (3e-48)				
	CLN3 (2e-07) CLN3 (2e-05) CLN1 (>e-5)	CLB6	NCU02114.1 (1e-19) NCU02758.1 (1e-59) NCU01242.1 (1e-48) NCU02114.1 (5e-23)				

TABLE 56. The cyclin family of Neurospora^a

^a Sequences were compared using BLAST, and the corresponding *P* value in each case is indicated in parentheses. The databases used in the analysis are as follows: *S. cerevisiae*, http://www.yeastgenome.org/; *N. crassa*, http://www.genome.wi.mit.edu/annotation/fungi/neurospora/; *S. pombe*, http://www.sanger.ac.uk/Projects/S_pombe/.

itors from BLAST searches is not surprising, since these proteins have very little, if any, sequence similarity. NCU07565.1, however, has weak similarity to budding yeast *FAR1*. Three *Neurospora* proteins with predicted ankyrin repeats (NCU07934.1, NCU02967.1, and NCU01098.1) all display weak similarity to human p16/INK4, an ankyrin repeat protein. Ankyrin repeats are important for Cdk inhibition, but not all ankyrin repeat proteins are Cdk inhibitors (53).

Finally, in addition to cyclin binding, Cdks need to be phosphorylated on a conserved threonine residue to be activated. In *S. cerevisiae*, Cak1p catalyzes this phosphorylation, while mammals use a different Cdk for this activation (550). *Neurospora* probably uses the same mechanism found in *S. cerevisiae* for Cdk activation, since it contains a Cak1p-like predicted protein (NCU04426.1).

Asexual and Sexual Sporulation

Spore formation is a common mechanism among fungi for reproduction, dispersal, and survival under harsh conditions (7, 184). For most pathogenic fungi, spores are the major source of infection of their hosts. Many filamentous fungi reproduce both sexually and asexually. The asexual cycle is mitotic, while the sexual cycle involves mating and meiosis.

Macroconidiation. In the filamentous ascomycetes, asexual reproduction involves the production of macroconidia at the tips of specialized hyphae called conidiophores (7, 751). Macroconidiation begins with the differentiation of aerial hyphae that grow perpendicular to the surface mycelium. Following a period of apical aerial growth, the aerial hyphae switch to a

budding mode of growth that results in the formation of proconidial chains within the conidiophore. As the conidia reach maturity, conidial separation takes place. Free conidia are released and are dispersed primarily by air currents. The macroconidia germinate rapidly, allowing efficient reproduction of the fungus. Macroconidiation can be induced by environmental signals including heat shock, desiccation, and carbon or nitrogen starvation (808). Furthermore, the endogenous circadian clock regulates the timing of conidiation in Neurospora (484). Asexual development in the filamentous ascomycetes is considered to be a relatively simple process and is not essential for viability. Thus, these organisms serve as excellent models for uncovering the genes involved in, and the mechanisms of, asexual spore differentiation. Using both forward and reverse genetic techniques, key components of the macroconidiation pathway have been identified in Neurospora and A. nidulans.

The *A. nidulans* FadA and FluG proteins are considered to be upstream regulators of conidiation, since they affect the expression levels of a key regulator of development, *brlA* (7). FluG produces an extracellular factor that signals conidiophore development, while FadA is a heterotrimeric G-protein α subunit which, when activated, blocks sporulation. Homologues of both FluG and FadA are present in the *Neurospora* genome (NCU04264.1 and GNA-1, respectively), and mutation of *gna-1* leads to multiple developmental phenotypes, including defects in macroconidiation (369). In contrast, there appears to be significantly less conservation, between these two species, of key components that act downstream of FadA and FluG (see below).

In A. nidulans, the FlbC, FlbD, BrlA, AbaA, and WetA proteins are required for the normal production of conidiospores and StuA and MedA are developmental modifiers that are necessary for the normal spatial organization of the conidiophore (7). There are no *Neurospora* homologues of BrlA or AbaA. There is a hypothetical Neurospora protein (NCU06975.1) with weak homology to the C terminus of WetA and a hypothetical protein (NCU03043.1) with similarity to FlbC, but the identity is limited primarily to the zinc finger domain. A homologue of the MedA transcription factor is present in Neurospora (NCU07617.1) and in M. grisea (ACR1). ACR1 is a stage-specific negative regulator of conidiation in Magnaporthe (454). A putative Neurospora homolog of flbD, rca-1, was shown to complement the A. nidulans flbD mutation. Surprisingly, mutation of *rca-1* has no effect on conidiation in Neurospora (726). Similarly, a stuA homologue, asm-1, is required for sexual development but not macroconidiation in Neurospora (25). In Neurospora, the ACON-2, ACON-3, FL, and FLD proteins are required for macroconidiation (752). Of these, the only corresponding gene that has been cloned is fl(40). No homologues of FL are found in other ascomycetes, including A. nidulans and M. grisea. The closest match to FL in the database is to a hypothetical Neurospora protein (NCU09205.1).

The con genes are a set of cloned Neurospora loci that are preferentially expressed during conidiation but are not essential for development (69). In most cases, searches using the con gene sequences yielded few clues to their cellular functions. However, the closest match to CON-7 is another hypothetical Neurospora protein (NCU07846.1). NCUO7846.1 has similarity to a putative transcriptional regulator from M. grisea (AAB69694.1), suggesting that CON-7 and the related Neurospora protein may be involved in transcriptional regulation. EAS (CCG-2) is a fungal hydrophobin that is induced during macroconidiation and coats the mature condiospore, rendering the spores hydrophobic and air dispersible (62, 455). Searches using EAS identified a putative second Neurospora hydrophobin (NCU08457.1). This finding is surprising, since deletion of eas (ccg-2) results in wet, clumpy spores that lack the hydrophobic rodlet layer (62), and suggests either a different role for the NCU08457.1 hydrophobin or a requirement for EAS (CCG-2) for its expression and activity.

In summary, the apparent lack of conservation of key regulators of conidiation in *A. nidulans* and *Neurospora* suggests the possibility that macroconidiation has evolved independently in these two organisms. In contrast, there appears to be significant conservation among the upstream signaling components, perhaps reflecting the similar ways in which these fungi respond to the same environmental signals to initiate macroconidial development.

Meiosis and the sexual cycle. In fungi, meiosis is intimately associated with sporulation (436). The sexual development of filamentous ascomycetes is characterized by the formation of a fertilized fruiting body containing asci, which, in turn, enclose the progeny spores. In contrast, yeasts form an ascus directly from a single diploid cell without the involvement of a fruiting body. Numerous genes have been identified that are involved in meiosis and ascus development in *Neurospora* and closely related filamentous ascomycetes (456, 638; see also "Meiotic recombination" above).

In Neurospora, a single isolate can produce both female and male reproductive structures, but sexual reproduction can take place only between strains of opposite mating type, matA and mata (reviewed in references 436 and 638). The formation of female reproductive structures (protoperithecia) is induced by nitrogen limitation. These structures produce specialized hyphae (trichogynes) that exhibit chemotropic growth toward male cells (usually conidia or hyphae) of the opposite mating type in an apparent pheromone response pathway (81, 87, 414). Contact between the trichogyne and the male cell leads to entry of the male nucleus into the trichogyne and its subsequent transport to the ascogonium cell of the protoperithecium (reviewed in references 169, 184, and 638). The male and female nuclei do not fuse immediately after fertilization but instead undergo a series of mitotic divisions to produce an ascogenous hyphal mass. Later, nuclei of opposite mating types pair and undergo simultaneous mitotic divisions at the tips of ascogenous hyphae to yield distinct cell types, including the binucleate cell in which karyogamy (nuclear fusion) takes place. Each resulting diploid cell immediately enters meiosis, followed by a postmeiotic mitosis, to yield an ascus containing eight ascospores.

S. cerevisiae has been extensively used for the analysis of meiosis and sporulation. To identify genes that are regulated during meiosis and conserved across yeast species, the *S. cerevisiae* and *S. pombe* meiotic transcriptomes were compared (507, 686). These analyses identified a group of approximately 75 similarly regulated meiotic genes, including components of the anaphase-promoting complex and genes involved in recombination, sister chromatid cohesion, and synapsis (686). From these analyses, a list of 74 yeast genes that are essential for meiosis and sporulation, but not for mitotic growth, was compiled and the sequences were compared to the fly, worm, mouse, and human genomes (www.biozentrum.unibas.ch/personal/primig/gamates). A comparison of several of these core meiotic gene products to the *Neurospora* sequence database revealed several interesting features (Table 57).

Of the three known transcription factors specifically involved in meiotic gene transcription in S. cerevisiae (Abf1p, Ume6p, and Ndt80p [147, 268, 333]), only Ndt80p, a meiosisspecific transcription factor that induces genes at the end of prophase, appears to be conserved in Neurospora. Transcription of Ndt80p is itself dependent on Ime1p, which activates the expression of early sporulation genes; an ImeIp homologue is also present in Neurospora. Ume6p, which interacts with Ime1p and recognizes a conserved URS1 site in the promoters of many genes that are activated early in meiosis and are associated with chromosome pairing and recombination, is absent from Neurospora. Consistent with the lack of Ume6p, several of the products of the genes in S. cerevisiae known to be regulated by this factor (854) are not present in Neurospora (with the exception of Spo11p). Ime2p is the founding member of a family of protein kinases that are required for effective progression through meiotic development. Ime2p is essential for the induction of meiosis-specific genes and for the activation of meiotic DNA replication in S. cerevisiae. Orthologs of Ime2p and Rim15p (required for Ime2p expression) are present in Neurospora; however, a regulator of Ime2p expression, Rim4p, is absent.

TABLE 57.	Meiosis	and	sexual	sporulation
-----------	---------	-----	--------	-------------

S. cerevisiae	Evention	Neurospora	BLAST	Best overall BLAST match	Homologue/
enzyme	Function	gene name	e value	to Neurospora	and animals?
Meiosis					
Abf1	ARS1 binding protein/transcriptional	None			
Ama1	Activator of meiotic anaphase- promoting complex	1572.1	2e-42	S. cerevisiae AMA1 AAK61800	Yes
Cdc16	Subunit of anaphase-promoting complex	1377.1	1e-102	S. pombe Cut9 NP593301	Yes
Csm1	Chromosome segregation in meiosis	None			No
Doc1	Component of the anaphase- promoting complex	8731.1	5e-17	A. thaliana expressed protein NP565433	Yes
Dmc1	Meiotic recombination	No match	4 4 40	AAB39323	
Hmf1 Hop1	DNA helicase meiotic crossing over	9793.1 None	1e-149	S. cerevisiae HMF1 NP588310	Yes
Hop1 Hop2	Synaptonemal complex component	None			INO
Isd2	IME2-dependent signaling	None			
Ime2	Serine/threonine kinase; positive regulator of meiosis	1498.1	7e-69	S. pombe Ser/Thr protein kinase NP593607	Yes
Ime4	Activates IME1	None			No
Mam1	Monoorientation of sister kinetochores	7984.1	2e-8	<i>S. pombe</i> monopolin complex component CAD88639	No
Mei4	Chromosome pairing	None			No
Mei5	Synapsis and meiotic recombination	None	2- 25 2- 24	II anniana anatain hinasa CUIV2 isafamma	No
Mih2	Mismatch rongin	2814.1, 9123.1	2e-35, 2e-34	H. saptens protein kinase CHK2 isoform a NP009125	Yes
Mnd1	Meiotic recombination	None			105
Mpc54	Meiotic spindle pole body component	0658.1, 9063.1	4e-08, 9e-07	E. histolytica myosin heavy chain T18296	Yes
Mre11	Meiotic DNA DSB formation	8730.1	1e-123	S. cerevisiae Smc4 NP013187	Yes
Msh4	Meiotic recombination	2230.1, Msh2	1e-40	S. cerevisiae homologue of MutS AAA34802	Yes
Msh5	Reciprocal recombination between homologs	9384.1	1e-58	L. maculans mismatch repair protein	Yes
Mum2	Premeiotic DNA synthesis	None			
Ndt80	Meiosis-specific gene	9915.1	4e-8	S cerevisiae NDT80 P38830	No
Pch2	Pachytene checkpoint	None		5. 2010/15/201001200000	No
Rec8	Recombination and sister chromatid cohesion	None			Yes
Rec102	Meiotic DNA DSB formation	None			No
Rec104	Meiotic DNA DSB formation	None			No
Rec10/ Rec114	Meiotic DNA DSB formation	None			No
Red1	Synaptonemal complex formation	None			No
Rim4	Regulator of IME2 expression	None			No
Rim11	IME1/transcriptional regulator	4185.1	1e-106	C. gloeosporioides protein kinase GSK AAN32716	Yes
Rim15	Required for IME2 expression	7378.1	1e-103	S. pombe Cek1p NP588310	Yes
Sae2	Meiotic DNA DSB processing	None			No
Sae3	Meiotic recombination	None	70.46	<i>R</i> always group a phospholipose P D20457	No
Spo11 Spo11	Endoribonuclease meiotic DSB formation	1120.1/REC12	2e-10	C. cinereus Spo11 AAF26720	Yes
Spo12	Regulates meiosis	None			
Spo19	Meiosis-specific GPI protein	None			No
Spo69	Sister chromatid cohesin component	None	2 2 72		
Sps1	Transcriptional regulator	//2.1 None	3 3-12	D. discoideum severin kinase AAC24522	Yes
Zin1	Synaptonemal complex formation	658.1	8e-24	E. histolytica myosin heavy chain T18296	Yes
Zip2	Synaptonemal complex formation	None	0021	2. Autorijaca mjesin neavj enam 1102/0	No
Zip3	Recombination nodules and synapses	None			No
Sporulation					
Ady2 Ady3	YaaH family of putative transporters Mediates assembly of the Don1p con- taining structure at the leading edge of the prospore membrane via inter- action with components of the	6043.1 None	2e-53	P. anserina CAD60593	No No
	spindle pole body				
Ady4	Sporulation	None			
Dit1 Dtr1	Spore wall maturation protein Dityrosine transporter spore wall assembly	1411.1	9e-39	S. cerevisiae A (acid, azole) Q (quinidine)	No
Isc10	Required for spore formation	None		105151a1100 141 009399	No

Continued on following page

S. cerevisiae enzyme	Function	<i>Neurospora</i> NCU no., gene name	BLAST e value	Best overall BLAST match to Neurospora	Homologue/ orthologue in plants and animals?
Pfs1	Prospore formation	None			No
Sma1	Spore membrane assembly	None			No
Smk1	MAPK involved in cell wall formation	9842.1	1e-75	C. lagenarium MAPK AAL50116	Yes
Spo14	Phospholipase D	3955.1	4e-51	S. pombe putative phospholipase D1 NP592986	Yes
Spo16	Early meiotic protein required for efficient spore formation	None			No
Spo20	v-SNARE, spore wall maturation	9243.1	5e-8	S. cerevisiae SNAP 25 homologue NP013730	No
Spo22	Meiosis-specific phospholipase A2 homolog	None		Ũ	
Spo71	Spore wall maturation	None			
Spo73	Sporulation	None			
Spo74	Sporulation	None			
Spo75	Sporulation	8776.1	1e-33	S. cerevisiae NP013993	Yes
Spo77	Sporulation	None			
Spr1	Glucan 1,3-β-glucosidase	3914.1	3e-53	C. immitis B-glucosidase 6 AAL09830	Yes
Spr3	Spore wall assembly	3795.1	4e-65	A. nidulans septin AAK21000	Yes
Sps18	Transcription factor	7734.1	9e-20	S. cerevisiae Gcs1p NP010055	Yes
Sps100	Spore wall maturation	None		x	No
Ssp1	Spore wall maturation	None			No
Ssp2	Spore wall maturation	None			No
Swm1	Spore wall maturation	None			No

TABLE 57-Continued

Given that the process of meiosis is similar in distantly related organisms, it is surprising to find that very few proteins specific to meiotic chromosome behavior in S. cerevisiae appear to be conserved in Neurospora (Table 57). When the proteins are shared between Neurospora and S. cerevisiae, they are in most cases also present in higher eukaryotes. Thus, while several of the regulators of meiosis appear conserved, many of the other proteins are not. Conspicuously absent from Neurospora are proteins with similarity to those required in S. cerevisiae for premeiotic DNA synthesis, chromosome segregation and pairing, sister chromatid cohesion, and pachytene checkpoint control. However, a cursory comparison of Neurospora proteins to proteins in other higher eukaryotes known to be involved in meiosis identified potential Neurospora homologues of gene products required for pachytene checkpoint control (nim-1, related to the mouse NimA kinase; CHK2, a C. elegans checkpoint control protein required for meiosis; and NCU02814.1), chromatid adhesion (human CDCA1 and NCU06568.1), and chromosome segregation and pairing (mouse SMC11 and NCU01323.1). These data point to Neurospora as an evolutionary intermediate between yeast and higher organisms with respect to the proteins involved in meiosis and suggest that there are only a limited number of core meiotic genes conserved among eukaryotic organisms.

Less than half of the conserved *S. cerevisiae* sporulationspecific gene products examined are present in the *Neurospora* genome (Table 57). Similar to meiosis-specific proteins, the signaling components required for sporulation appear to be more strongly conserved than the structural proteins. In any case, proteins that do show similarity are likely to be involved in spore or ascus formation in *Neurospora*.

FUNGAL PATHOGENESIS AND HUMAN DISEASE

Relationship to Animal and Plant Pathogens

Ideally, antifungal compounds should target gene products essential for the growth and development of the fungus in the host without affecting the function of the host cells. In addition, drugs that target a large variety of fungi are especially useful; however, many identified fungal virulence factors are pathogen specific. Development of a broad-acting drug involves the identification of gene products which have essential functions in many fungi but which are absent in or otherwise greatly differ from those found in humans. Nonetheless, this does not rule out the development and application of compounds that exhibit antifungal efficacy, even though they target cellular components common to the pathogen and the host (for example, benzimidazoles that inhibit β -tubulin polymerization). Systems which fungi possess and which mammals lack include cell walls and a variety of membranous components. Even though plants possess carbohydrate polymer-based cell walls, there are sufficient structural differences that provide a basis for specific antifungal agents that can be used for plant protection. Existing drugs take advantage of these differences; for example, azoles and polyenes are compounds that inhibit the function of the fungal cell membrane. Inhibitors of $\beta(1,3)$ -glucan synthesis, the candins, are being used to inhibit fungal cell wall function (237), and the investigation of cell wall synthesis continues to uncover potential antifungal targets (705).

Animal pathogens. Even though *Neurospora* is not a pathogen, it shares numerous properties with related fungal pathogens, and its genome can be used to identify targets that generally differ between humans and fungi. In addition, the genome sequence allows a comparison between nonpathogens and pathogens, which will aid in the identification of putative virulence factors used to adapt to human hosts. One interesting finding from analysis of the genome sequence is that *Neurospora* has several genes similar to those used to synthesize the polysaccharide capsule, a known virulence factor found in the pathogen *Cryptococcus neoformans*, the causative agent of fatal meningoencophalitis in AIDS patients (Table 58) (850). The capsule of *C. neoformans* surrounds the cell wall and mediates the immune response with the host. Multiple effects of the capsule include the ability to inhibit inflammatory cyto-

TABLE 58. Human pathogenesis-associated genes in Neurospora

Neurospora gene	Homologue	Organism	Pathogenesis function
NCU06430.1	CAP10	C. neoformans	Capsule
NCU02336.1		0	1
NCU05123.1			
NCU02119.1			
NCU05916.1	CAP59	C. neoformans	Capsule
NCU04473.1	CAP60	C. neoformans	Capsule
nik-1/os-1	COS1	C. albicans	Unknown
NCU07221.1	fos-1	A. fumigatus	Unknown

kine production and complement factor and leukocyte migration, all of which contribute to evasion of the host immune system. In addition, the capsule prevents efficient phagocytosis by macrophages and thus leads to persistence of the pathogen in the host. Neurospora possesses proteins (Table 58) similar to those encoded by three genes (CAP10, CAP59, and CAP60) that are implicated in capsule formation in C. neoformans and are absent from S. cerevisiae (98). The finding that Neurospora lacks a fourth cap gene (CAP64), a glycosyltransferase required for O acetylation (CAS1) of the main capsule polysaccharide, and a UDP-glucuronate decarboxylase (CAS2), may explain the absence of this structure in *Neurospora* and also suggests noncapsule functions for CAP10/CAP59/CAP60 in filamentous fungi. Laccase is also a known virulence factor for C. neoformans, and Neurospora has several laccase genes (see below).

Emerging antifungal targets are processes regulated by twocomponent signal transduction systems (see "Environmental sensing" above) (Table 58). HKs found in *C. albicans* (Cos1/ CaNik1) and *A. fumigatus* (Fos-1), which have been shown to affect virulence, have homologues in *Neurospora*: NIK-1 and NCU07221.1, respectively (153, 704). Compounds with activity against these types of systems have been described but are not currently in use (492). The analysis of the *Neurospora* genome has shown that 63% of the predicted proteins lack homologues in *S. cerevisiae*, a species that is commonly used in antifungal screening processes. Unshared components that are possible antifungal targets include a wealth of two-component signal transduction systems, multicomponent cytoskeletal motor complexes, etc. It is likely that additional potential targets for antifungal development are present within the approximately 40% of the putative *Neurospora* ORFs without homologues in other organisms.

Plant pathogens. Members of the fungal kingdom exhibit a broad spectrum of lifestyles, ranging from saprophytic to obligate parasitism. Furthermore, the same fungal species can sometimes exhibit different lifestyles depending on the host range and/or environmental conditions. In some instances, pathogencity and virulence genes were defined as such because when discovered, their only observable role was in determining the pathogenic attributes of the organism studied. In other instances, analysis of many gene products involved in functions that were not immediately or intuitively associated with pathogenicity later resulted in the appreciation that they are required for the pathogenic phase.

Even though *Neurospora* is not known to be intimately associated with living plants, the genome sequence has revealed the presence of genes whose putative products are highly similar to those shown to be strictly associated with pathogenicity in other organisms (Table 59). Genera containing these proteins include *Botrytis* (292), *Colletotrichum* (360), *Magnaporthe* (195), *Nectria* (311), and others (174). All of the listed putative proteins, with the exception of NCU5730.1, which resembles the *N. haematococca* PEP2 gene product (311), have apparent homologues in *M. grisea* and *F. graminearum*. Furthermore, similar proteins (with the exception of NCU07432.1, a tetraspanin-like protein) are also found in the human pathogen *A. fumigatus*, supporting the possibility that the respective gene products are likely to function in ways that are not limited to plant-pathogen interactions.

Destain	NCU	BLAST match							
Protein	no.	Best overall	S. cerevisiae	S. pombe	Animal ^a	Plant ^l			
Nitrogen deprivation	03370.1	B. cinerea; 6e-38	None	None	None	None			
Probable cutinase percursor ^c	09663.1	<i>M. tuberculosis</i> probable cutinase; 9e-6	None	None	None	None			
Pathogenicity-related protein	08038.1	M. grisea MAS3/ASG1; 3e-41	None	No hits	None	None			
N. crassa Histidine kinase (NIK1)	02815.1	N. crassa NIK1 and 8 other fungi; 0.0	3e-16	5e-59	2e-7; H. sapiens	2e-68			
Membrane protein that mediates differentiation in response to inductive substrate cues	02903.1	M. grisea PTH11; 3e-11	None	None	None	None			
N. crassa PLS1	07432.1	Colletotrichum lindemuthianum; 3e-73	None	None	None	None			
Pathogenicity-related protein	06170.1	M. grisea MAS3/ASG1; e-17	None	None	8e-9; D. melanogaster	None			
Pathogenicity protein	05521.1	M. grisea PATH531; 4e-70	5e-17	6e-13	5e-19; M. musculus	e-13			
Pathogenicity protein	02604.1	M. grisea PTH10; 2e-55	2e-20	2e-35	7e-16; D. melanogaster	2e-11			
Putative transcription factor	07846.1	Botrytis cinerea; 8e-28	None	None	4e-8; D. melanogaster	2e-7			
PEP2 Pathogenicity cluster	05730.1	Nectria haematococca; 7e-40	None	None	None	None			
Putative transcription factor	01422.1	Botrytis cinerea; 3e-47	e-9	3e-29	6e-21; M. musculus	8e-14			

TABLE 59. Neurospora plant pathogenicity-related proteins

^a Caenorhabditis elegans, Drosophila melanogaster, Mus musculus, or Homo sapiens.

^b Arabidopsis thaliana.

^c Not required for virulence in F. solani, but still a potential cutinase in Neurospora.

Enguno	Disease or	NCU	BLAST match				
Elizyille	disorder	no.	Best overall	S. cerevisiae	S. pombe	Human	
With homologues in yeast							
10 chloride channel protein	Dent's disease (renal)	06624.1	S. pombe/e-124	e-135	e-162	e-124	
DNA ligase I polydeoxyribo- nucleotide synthase [ATP]	Immunodeficiency	09706.10	S. pombe, R. norvegicus DNL1/7e-61	4e-46	7e-61	3e-60	
		06481.1	S. cerevisiae Cdc9/0.0	0.0	e-116	e-146	
Calcium/calmodulin-depen- dent protein kinase type I (CAM kinase 1)	Cancer	09212.1	S. pombe Srk1/e-133	8e-92	e-133	5e-57	
· · · · · ·		09123.1	E. nidulans KCC1/e-166	e-96	2e-64	7e-69	
Probable ATP-dependent permease (multidrug resis- tance-associated protein 1)	Cancer related	09012.1	S. cerevisiae YCFI/0.0	0.0	0.0	0.0	
Probable ATP-dependent permease (multidrug resis- tance-associated protein 2)	Cancer related	08358.1	S. cerevisiae YBT1/0.0	0.0	e-146	e-180	
Copper-transporting ATPase (copper pump 2)	Wilson's disease	07531.1	S. cerevisiae Pca1/0.0	0.0	7e-70	8e-66	
		08341.1	A. thaliana RAN1/0.0	e-175	0.0	e-170	
Carnitine acetyltransferase	CPT2 deficiency/metabolic	08002.1	S. cerevisiae Cat2/e-123	e-123	None	6e-86	
TSC2	Tuberous sclerosis type 2	4105.1	S. pombe SPAC630.13c/e-100	None	e-100	NP_066399.1; 3e-36	
COT1 kinase	Myotonic dystrophy	07296.1	S. cerevisiae Cbk1/e-151	e-151	e-144	2e-67 ^a	
Without homologues in yeast							
Tripeptidyl-peptidase I pre- cursor, TPP-I (CLN2)	Ceroid-lipofuscinosis	08656.1	A. oryzae aorsin/7e-33	None	None	4e-18	
		08418.1	<i>A. oryzae</i> tripeptidylpeptidase A/e-145	None	None	5e-59	
		04903.1	C. brassiana tripeptidylpepti- dase precursor/1e-89	None	None	5e-38	
Calpain-like thiol protease	Limb girdle MD ⁺ 2A- CAPN3	1151.1	N. crassa NCU00251.1/e-120	None	None	5e-38	
		0251.1	N. crassa NCU01151.1/e-110	None	None	9e-30	
		3355.1	N. crassa NCU01151.1/8e-98	None	None	8e-37	

TABLE 60. Neurospora proteins	exhibiting	similarities to	human	proteins	associated	with	disease
-------------------------------	------------	-----------------	-------	----------	------------	------	---------

^a Also similar to human NDR kinase (e-108).

In addition to the listed gene products, many components known to be involved in both pathogen- and non-pathogenrelated functions of fungi are clearly present in Neurospora. The potentials for metabolism or efflux of toxic plant metabolites (e.g., P450 cytochrome oxidases and ABC transporters, respectively), biotic sensing and stress signal transduction, and polyketide biosynthesis are just a few examples of cellular machinery capable of performing such functions that are present in Neurospora. It is clear that the differences in the gene content of Neurospora and Magnaporthe will very soon be a focus of investigation. These fungi are considered close relatives but are estimated to have evolved from a common ancestor 50 to 150 Mya ago. Given this evolutionary distance, it is not surprising that they appear to share only about 60% of their genes. The question of whether their common ancestor was a plant pathogen, a saprophyte, or a nonpathogenic symbiont may remain unanswered until additional fungal genome sequences are available. The presence of apparent homologues of genes for secondary metabolism and plant pathogen virulence factors in Neurospora suggests that the lineage leading to Neurospora is just as likely to have lost its ancestral ability to parasitize plants as the lineage leading to Magnaporthe is to have gained parasitism (269).

Human Disease Genes

There are over 200 predicted Neurospora proteins exhibiting significant similarity to human gene products that, when altered, have been demonstrated to cause disease (Table 60). The range of proteins associated with human diseases that is represented in the Neurospora genome is vast and includes those implicated in diseases of the immune system, metabolic disorders, neurological impairments, and cancer (for a list of examples, see Table 60). Although many of these proteins have counterparts in S. cerevisiae and S. pombe, some of the features of Neurospora (multinucleate, multicellular, and exhibiting distinct and diverse developmental phenotypes) provide an attractive platform for investigating the functions of these genes with the intention of increasing our understanding of the relevant human disease. Furthermore, the observation that in some instances the Neurospora proteins have high similarity to proteins in one (but not both) of the yeasts (e.g., TSC2 and carnitine acetyltransferase) may have evolutionary significance. In many cases, Neurospora has more than one predicted protein that is associated with the same disorder (e.g., DNA ligase, CAM kinase 1, and copper-transporting ATPase), raising the question of whether both (or any) of the structurally related proteins are relevant to the respective human diseases.

Two human disease genes associated with ceroid lipofusci-

nosis and limb-girdle muscular dystrophies (LGMD) are represented in the *Neurospora* genome but are absent from both S. cerevisiae and S. pombe. Ceroid lipofuscinosis is an inherited degenerative disease characterized by neuronal cytoplasmic inclusions that stain positively for ceroid and lipofuscin (168). Affected individuals develop retinal degeneration, seizures, myoclonus, ataxia, rigidity, and progressive dementia. Individual genes mutated in six forms of ceroid lipofuscinosis have been identified. The products of these genes fall into two distinct categories comprising either soluble lysosomal enzymes (CLN1 and CLN2) or predicted transmembrane proteins of unknown structure and function (CLN3, CLN5, CLN6, and CLN8). Neurospora has three predicted polypeptides that show significant structural similarity to the CLN2 class of proteins. The CLN2 gene encodes tripeptidyl peptidase 1, which cleaves tripeptides from the N-terminus of small proteins before their degradation by other lysosomal proteases.

The LGMDs are a heterogenous group of genetically determined progressive disorders of skeletal muscle with a primary or predominant involvement of the pelvic or shoulder girdle musculature. At least 15 genes have been identified which, when mutated, can cause LGMD, among them calpain-3, a Ca^{2+} -activated cysteine protease, responsible for LGMD2A (650). The fact that two such proteins (NCU03355.1 and NCU01151.1) are present in *Neurospora* but not in yeasts is in line with the general observation that significantly more Ca^{2+} signaling proteins are present in *Neurospora* than in *S. cerevisiae* (Table 47) A. Zelter, M. Bencina, B. J. Bowman, O. Yarden, and N. D. Read, unpublished data).

Many of the other gene products known to be associated with LGMD (e.g., dysferlin, telethonin, myotilin, FKRP, and sarcoglycan beta [894]) do not have structural homologues in *Neurospora* or in *S. cerevisiae* and *S. pombe*. The absence of these other LGMD proteins in fungi could be because they are involved in sarcomeric functions, and fungi lack sarcomeres. Should these proteins be involved in additional (nonsarcomeric) cellular activities, it is conceivable that their *Neurospora* functional homologues would not be identified by BLAST-based structural similarity searches.

The observation that six of the unique *Neurospora* gene products associated with human disease are proteases is intriguing. It could be argued that *Neurospora* would be expected to have an abundance of proteases due to its ability to metabolize a very diverse set of nutrients compared to yeasts. However, the six proteases are not predicted to be secreted (Table 26) and are therefore not likely to be involved in extracellular digestion processes. This suggests the existence of fundamental cellular processes involving these proteases that are shared between *Neurospora* and humans but are absent from yeasts; it also suggests that *Neurospora* is an excellent system for the study of these processes.

The observation that *Neurospora* has an abundance of transporters (see "Metabolic processes and transport" above) (269) is also relevant to human disease. For example, there are at least two highly similar copper transporters (NCU07531.1 and NCU08341.1) associated with Wilson's disease, an inherited disorder that causes the body to retain copper and can result in severe brain damage, liver failure, and death. The structural and functional dissection of these two transporters may provide important information concerning copper transport and

the metabolic consequences of impaired copper transport in eukaryotes. The finding of two related proteins in *Neurospora* suggests the potential for studying subtleties in the copper uptake process in this organism. Another exciting medically related potential of studying the multitude of transporters in *Neurospora* is the elucidation of drug uptake/resistance processes transporters (e.g., NCU09012.1 and NCU08358.1), which are associated with the success or failure of drug treatments in humans. These, of course, are also of prime interest from the viewpoint of successful application of antifungals (see "Relationship to animal and plant pathogens" above).

Some of the *Neurospora* proteins that have significant homology to human disease gene products are also very similar to additional human proteins. For example, the *Neurospora* COT1 kinase (877) is highly similar to the human myotonic dystrophy kinase but may be even more homologous to the human NDR kinase, which is involved in cell proliferation and tumor development (779). Thus, on the one hand, the similarity of *Neurospora* proteins to specific human counterparts poses an exciting avenue for functional analysis in relation to human diseases. However, in most cases, determining the extent of functional homology of the *Neurospora* (or any given model organism) gene product to the human disease-associated protein awaits analysis.

PERSPECTIVES AND FUTURE DIRECTIONS

This analysis of the Neurospora genome is provided as a starting point for further understanding of filamentous fungi and other multicellular eukaryotes and to facilitate progress not only in basic research but also in study of animal and plant pathogenesis, biotechnology, and biodegradation. With just over 10% of the gene complement of 10,000 ORFs analyzed, the investigation has already revealed many unexpected and exciting avenues for future studies. Many predicted Neurospora proteins have no homologues in the yeasts S. cerevisiae and S. pombe but are similar to proteins in animals, plants, and other filamentous fungi. Furthermore, Neurospora contains numerous gene products that are found in the two yeasts but for which the Neurospora protein is a better match for the corresponding animal or plant protein. These features suggest that Neurospora is an excellent model system for studies of numerous aspects of biology.

The related arenas of chromosome structure and gene regulation appear ripe for future productive analysis. For example, Neurospora possesses a virtually complete repertoire of vertebrate histone-modifying enzymes, suggesting that this organism may be an excellent system in which to describe the histone code and its role in epigenetic regulation of gene expression and development. The compilation of transcription factors offers clues that will benefit researchers working on a multitude of biological processes that contain a transcriptional regulatory component. Remarkably, only 14 of the 186 annotated transcription factor genes had previously been cloned and characterized. The Zn(II)₂Cys₆ fungal binuclear cluster family appears to have several family members that have partners, suggesting their involvement in a combinatorial regulatory process. The C2H2 zinc finger transcription factor family was divided into two groups, one that was most similar to factors from unicellular yeasts and one that was most similar to

proteins from filamentous fungi and animals. This dichotomy could represent a point of divergence in the evolution of promoters. In the future, it will be interesting to determine which genes are controlled by these two groups of factors and to investigate their relative importance to gene regulation in *Neurospora*.

The formation and maintenance of silent chromatin in Neurospora appears to be a variation of the HP1/Swi-6-mediated pathway of S. pombe. A connection between DNA methylation and the HP1/Swi-6 pathway was first experimentally determined for Neurospora and has now also been established in plants and animals. In contrast to Neurospora, the formation of silent chromatin in S. cerevisiae relies on complexes of Sir proteins, which appear absent from the Neurospora genome. While some of the processes are shared between Neurospora, fission yeast, plants, and animals, some differences have also been revealed from the genome sequence. It will be interesting to determine exactly how Neurospora recognizes and maintains methylated DNA, given the fact that it lacks recognizable homologues of plant and animal proteins involved in methylated DNA binding. In addition, the cytosine DNA methyltransferases of Neurospora are distinct from those found in other eukaryotes.

Besides DNA methylation, an additional three distinct genome defense mechanisms have been identified in Neurospora. RIP is a process that extensively mutates and methylates duplicated DNA sequences in the haploid genome during the sexual phase of development. RIP is thought to protect the genome from invasion by transposons. Two other mechanisms which rely on RNA-mediated gene silencing have also been discovered: quelling and meiotic silencing. In fact, genome analysis indicates that there are orthologues of the known components of eukaryotic RNA-dependent silencing in Neurospora (Argonaute-like translation initiation factors, Dicerlike RNases, etc.). In addition, further analysis of the Neurospora proteins indicates that they, like the other fungal proteins, fall into two distinct clades, suggesting that they function in similar but distinct pathways. These predictions can now be tested.

The recombination machinery of *Neurospora* is more like that of other filamentous fungi and mammals than that of *S. cerevisiae*; which has many genes required for recombination that have no identified homologue in the more complex organisms. Even *S. cerevisiae* Spo11p, responsible for initiating recombination by a DSB, has limited homology to the equivalent proteins of higher eukaryotes and requires several accessory proteins that are not needed in other organisms. These data suggest that recombination may be achieved in a variety of ways in different species and emphasize the need for detailed analysis in other tractable species such as *Neurospora*. Additionally, *Neurospora* has genes known to exert a level of recombination control (133, 134) not seen in *S. cerevisiae*, and the question whether such regulation occurs in other eukaryotes might be answered by their cloning.

The genome contains a unique diversity of proteins, both animal-like and plant-like, involved in various areas of metabolism that will make *Neurospora* an attractive eukaryotic system for study. However, functional studies are needed to fully exploit and augment the information from the genome sequence. For example, the identity of the sensors for carbon, nitrogen and sulfur, as well as the subsequent signal transduction pathways that regulate the assimilation of these critical metabolites, cannot be predicted from genome analysis and remain unknown. Similarly, an accurate description of the mitochondrial proteome is needed, since analysis of the genome sequence did not yield a complete inventory of the proteins in this organelle.

Neurospora possesses a wide array of environmental sensory capabilities and promises to be an outstanding model for investigations of signaling. The larger number of HKs and GPCRs in Neurospora compared to S. cerevisiae and S. pombe predicts the existence of novel signaling pathways and/or networking interactions not found in yeasts. The completely unexpected identification of cryptochrome and phytochrome homologues has sparked renewed interest in the photobiology of Neurospora, leading investigators to look more closely at the role of light in the development and real-world biology of this organism and of filamentous fungi in general. Homologues of many of the key proteins involved in the release of Ca^{2+} from internal Ca²⁺ stores could not be recognized in Neurospora, indicating that there are significant differences between the Ca²⁺-signaling machinery in filamentous fungi and that in animals and plants. The different intracellular Ca²⁺ release mechanisms in filamentous fungi may provide novel targets for antifungal agents.

The polarized hyphal cell growth of *Neurospora* is reflected in its complement of proteins involved in generating the hyphal structure that in many cases is quite distinct from proteins that carry out similar functions in yeast cells. Analysis of cell wall polymer synthesis genes demonstrated that *Neurospora* lacks the (1,6) β -glucan synthesis pathway found in *S. cerevisiae*. In addition, components of the chitin synthesis machinery are more complex in *Neurospora*. Together, these results confirm the long-held idea that cell wall assembly in yeasts is not a universal model for fungal cell wall assembly.

Although S. cerevisiae has polarized modes of growth-production of psuedohyphae-the mechanisms used to generate these structures and the true hyphae in filamentous fungi are very different. The genome analysis bears this out, since key transcription factors necessary for the switch from budding to filamentous growth in yeast are not present in Neurospora. In addition, proteins involved in bud site selection are absent from Neurospora, suggesting novel processes for the determination of emerging hyphal branches and their spacing. The presence of the small GTPase Rac in the Neurospora genome, which is absent from S. cerevisiae, may have implications for actin-dependent polarized hyphal cell growth and division. In addition, Neurospora has a larger complement of cytoskeletal components than does S. cerevisiae. An unusual myosin protein containing a chitin synthase domain has been identified and has been shown to function in cell wall synthesis and integrity in other filamentous fungi.

An intriguing question arises from analysis of the cyclin/ CDK machinery. *Neurospora* has one Cdk (CDC-2) that is primarily responsible for cell cycle progression. However, it has only three cyclins, in contrast to the nine in *S. cerevisiae* and three in *S. pombe*. It may be that RIP has limited the set of *Neurospora* cyclins. However, given the more complex life cycle of *Neurospora*, this finding also suggests the possibility of novel mechanisms for regulation of cell cycle progression. In addition, functional analysis of the cyclin and Cdk genes may reveal the mechanisms leading to the observed asynchrony between the cell cycle and cell division in *Neurospora*.

The differentation of asexual spores, or conidia, has been well characterized in *Neurospora* and *A. nidulans*. The genes involved in these two processes appear to be quite different, suggesting that they have evolved independently. The production of sexual spores, ascospores, requires meiosis. Comparison of genes known to be involved in meiosis in *S. cerevisiae* revealed that only one transcription factor is conserved between the two species (Ndt80p) and that a key transcriptional regulator of meiotic genes in *S. cerevisiae* (Ume1p) is absent from *Neurospora*.

In most cases, the roles of *Neurospora* genes exhibiting structural similarity to human disease genes have yet to be determined. Are the biochemical and cellular functions of these genes similar to those observed in humans? If so, the amenability of *Neurospora* to genetic and physiological manipulations may prove extremely useful in the functional dissection of these proteins (especially in cases when these are not produced by the yeast model systems) with the long-term objective of intervening in the function of these genes or gene products. Are the consequences of mutations in such genes as detrimental in filamentous fungi as they are in higher eukaryotes? The answer to this question not only may provide additional insight into the biology of fungi but also may help to establish new links between developmental complexity and genome evolution.

Finally, as full fungal genome sequences become available, there will be increased interest in and need for comparative analysis at all levels (from locus organization to nucleotide sequence [see also reference 876]). In the future, data analysis rather than data acquisition will be the limiting factor to progress. Appropriate software must be developed to minimize the need for repeated manual (or even program-based) analyses, since many of the requirements can be defined a priori. There are numerous choices to be made regarding how genome data are curated, maintained, and funded, with consideration given to existing models for effectively organizing the data. Addressing these issues in a timely and organized manner represents an important milestone in the quest to maximize and speed the application of fungal genome information to medicine, agriculture, industry, and the environment.

ACKNOWLEDGMENTS

We are indebted to Matthew Springer and David Perkins for the use of previously published figures and legends, and we thank David Jacobson and David Perkins for providing a high-resolution version of Fig. 1. We thank Markus Aebi and Albert Courey for their comments and suggestions concerning the Dolichol pathway and the transcription factor analyses, respectively.

This analysis was supported by National Institutes of Health grant GM48626 and National Science Foundation grant MCB-0296055 (to K. A. Borkovich), National Institutes of Health grant S06 GM53933 (to L. A. Alex), Israel Science Foundation (to O. Yarden), Royal Society grant 066392/Z/01/Z and the Darwin Trust (to N. D. Read), Deutsche Forschungsgemeinschaft grant SPP1111 (to S. Seiler), National Institutes of Health grants GM58529 and NS39546 (to D. Bell-Pedersen), National Science Foundation grant MCB-0235871 (to M. Plamann), National Research Initiative of the USDA Competitive Research Grants Office 2001-35100-10618 (to N. Plesofsky), Deutsche Forschungsgemeinschaft grant Schu698/3 (to G. Mannhaupt and U. Schulte), Natural Sciences and Engineering Research Council 3041

and the Canadian Institutes of Health Research MOP-13170 (to F. E. Nargang), National Institutes of Health grant R37 GM34985 (to J. C. Dunlap), National Science Foundation grant MCB-0084509 (to J. J. Loros), National Institutes of Health grant MH44651 and the Norris Cotton Cancer Center Core Grant at Dartmouth Medical School (to J. C. Dunlap and J. J. Loros), Australian Research Council grant DP0345994 (to D. E. Catcheside), National Institutes of Health grant GM56770 (to R. Aramayo), National Institutes of Health grant GM062377 (to M. Polymenis), and National Institutes of Health grant GM35690 (to E. U. Selker).

REFERENCES

- Aaronson, L. R., and C. E. Martin. 1983. Temperature-induced modifications of glycosphingolipids in plasma membranes of *Neurospora crassa*. Biochim. Biophys. Acta 735:252–258.
- Abbas-Terki, T., O. Donze, P. A. Briand, and D. Picard. 2001. Hsp104 interacts with Hsp90 cochaperones in respiring yeast. Mol. Cell. Biol. 21: 7569–7575.
- Abbott, R. J., and G. A. Marzluf. 1984. Major extracellular protease of Neurospora crassa. J. Bacteriol. 159:505–510.
- Adam, M., F. Robert, M. Larochelle, and L. Gaudreau. 2001. H2A.Z is required for global chromatin integrity and for recruitment of RNA polymerase II under specific conditions. Mol. Cell. Biol. 21:6270–6279.
- 5. Adams, M. D., S. E. Celniker, R. A. Holt, C. A. Evans, J. D. Gocayne, P. G. Amanatides, S. E. Scherer, P. W. Li, R. A. Hoskins, R. F. Galle, R. A. George, S. E. Lewis, S. Richards, M. Ashburner, S. N. Henderson, G. G. Sutton, J. R. Wortman, M. D. Yandell, Q. Zhang, L. X. Chen, R. C. Brandon, Y. H. Rogers, R. G. Blazej, M. Champe, B. D. Pfeiffer, K. H. Wan, C. Doyle, E. G. Baxter, G. Helt, C. R. Nelson, G. L. Gabor, J. F. Abril, A. Agbayani, H. J. An, C. Andrews-Pfannkoch, D. Baldwin, R. M. Ballew, A. Basu, J. Baxendale, L. Bayraktaroglu, E. M. Beasley, K. Y. Beeson, P. V. Benos, B. P. Berman, D. Bhandari, S. Bolshakov, D. Borkova, M. R. Botchan, J. Bouck, P. Brokstein, P. Brottier, K. C. Burtis, D. A. Busam, H. Butler, E. Cadieu, A. Center, I. Chandra, J. M. Cherry, S. Cawley, C. Dahlke, L. B. Davenport, P. Davies, B. de Pablos, A. Delcher, Z. Deng, A. D. Mays, I. Dew, S. M. Dietz, K. Dodson, L. E. Doup, M. Downes, S. Dugan-Rocha, B. C. Dunkov, P. Dunn, K. J. Durbin, C. C. Evangelista, C. Ferraz, S. Ferriera, W. Fleischmann, C. Fosler, A. E. Gabrielian, N. S. Garg, W. M. Gelbart, K. Glasser, A. Glodek, F. Gong, J. H. Gorrell, Z. Gu, P. Guan, M. Harris, N. L. Harris, D. Harvey, T. J. Heiman, J. R. Hernandez, J. Houck, D. Hostin, K. A. Houston, T. J. Howland, M. H. Wei, C. Ibegwam, et al. 2000. The genome sequence of Drosophila melanogaster. Science 287:2185-2195.
- Adams, R. R., A. A. Tavares, A. Salzberg, H. J. Bellen, and D. M. Glover. 1998. pavarotti encodes a kinesin-like protein required to organize the central spindle and contractile ring for cytokinesis. Genes Dev. 12:1483– 1494.
- Adams, T. H., J. K. Wieser, and J. H. Yu. 1998. Asexual sporulation in Aspergillus nidulans. Microbiol. Mol. Biol. Rev. 62:35–54.
- Agrios, G. N. 1997. Plant pathology, 4th ed. Academic Press, London, United Kingdom.
- Agsteribbe, E., M. Hartog, and H. de Vries. 1989. Duplication of the tRNA(MMet) and tRNA(Cys) genes and of fragments of a gene encoding a subunit of the NADH dehydrogenase complex in *Neurospora crassa* mitochondrial DNA. Curr. Genet. 15:57–62.
- Ahearn, D. G., S. P. Meyers, and R. A. Nichols. 1968. Extracellular proteinases of yeasts and yeastlike fungi. Appl. Microbiol. 16:1370–1374.
- Aign, V., U. Schulte, and J. D. Hoheisel. 2001. Hybridization-based mapping of Neurospora crassa linkage groups II and V. Genetics 157:1015– 1020.
- Aist, J. R. 2002. Mitosis and motor proteins in the filamentous ascomycete, Nectria haematococca, and some related fungi. Int. Rev. Cytol. 212:239– 263.
- Alex, L. A., K. A. Borkovich, and M. I. Simon. 1996. Hyphal development in *Neurospora crassa:* involvement of a two-component histidine kinase. Proc. Natl. Acad. Sci. USA 93:3416–3421.
- Alex, L. A., C. Korch, C. P. Selitrennikoff, and M. I. Simon. 1998. COS1, a two-component histidine kinase that is involved in hyphal development in the opportunistic pathogen *Candida albicans*. Proc. Natl. Acad. Sci. USA 95:7069–7073.
- Aligue, R., H. Akhavan-Niak, and P. Russell. 1994. A role for Hsp90 in cell cycle control: Weel tyrosine kinase activity requires interaction with Hsp90. EMBO J. 13:6099–6106.
- Althaus, F. R., L. Hofferer, H. E. Kleczkowska, M. Malanga, H. Naegeli, P. L. Panzeter, and C. A. Realini. 1994. Histone shuttling by poly ADPribosylation. Mol. Cell. Biochem. 138:53–59.
- Altschul, S. F., T. L. Madden, A. A. Schaffer, J. Zhang, Z. Zhang, W. Miller, and J. Lipman. 1997. Gapped BLAST and PSI-BLAST, a new generation of protein database search programs. Nucleic Acids Res. 25:3389–3492.
- 18. Altschuler, D. L., A. Muro, A. Schijman, F. B. Almonacid, and H. N. Torres.

- protein family. FEBS Lett. 273:103–106.
 19. Amedeo, P., Y. Habu, K. Afsar, O. M. Scheid, and J. Paszkowski. 2000. Disruption of the plant gene MOM releases transcriptional silencing of methylated genes. Nature 405:203–206.
- Anand, M., K. Chakraburtty, M. J. Marton, A. G. Hinnebusch, and T. G. Kinzy. 2003. Functional interactions between yeast translation eukaryotic elongation factor (eEF) 1A and eEF3. J. Biol. Chem. 278:6985–6991.
- Ando, S., H. Yang, N. Nozaki, T. Okazaki, and K. Yoda. 2002. CENP-A, -B, and -C chromatin complex that contains the I-type alpha-satellite array constitutes the prekinetochore in HeLa cells. Mol. Cell. Biol. 22:2229–2241.
- Andoh, T., T. Yoko, Y. Matsui, and A. Toh. 1995. Molecular cloning of the plc1⁺ gene of Schizosaccharomyces pombe, which encodes a putative phos-phoinositide-specific phospholipase C. Yeast 11:179–185.
- Ang, D., K. Liberek, D. Skowyra, M. Zylicz, and C. Georgopoulos. 1991. Biological role and regulation of the universally conserved heat shock proteins. J. Biol. Chem. 266:24233–24236.
- Aramayo, R., and R. L. Metzenberg. 1996. Meiotic transvection in fungi. Cell 86:103–113.
- Aramayo, R., Y. Peleg, R. Addison, and R. Metzenberg. 1996. Asm-1⁺, a Neurospora crassa gene related to transcriptional regulators of fungal development. Genetics 144:991–1003.
- Argueso, J. L., D. Smith, J. Yi, M. Waase, S. Sarin, and E. Alani. 2002. Analysis of conditional mutations in the *Saccharomyces cerevisiae MLH1* gene in mismatch repair and in meiotic crossing over. Genetics 160:909– 921.
- Armstrong, J. A., O. Papoulas, G. Daubresse, A. S. Sperling, J. T. Lis, M. P. Scott, and J. W. Tamkun. 2002. The *Drosophila* BRM complex facilitates global transcription by RNA polymerase II. EMBO J. 21:5245–5254.
- Arndt, K. T., C. Styles, and G. R. Fink. 1987. Multiple global regulators control HIS4 transcription in yeast. Science 237:874–880.
- Aronson, B. D., K. A. Johnson, and J. C. Dunlap. 1994. Circadian clock locus frequency: protein encoded by a single open reading frame defines period length and temperature compensation. Proc. Natl. Acad. Sci. USA 91:7683–7687.
- Aronson, B. D., K. A. Johnson, J. J. Loros, and J. C. Dunlap. 1994. Negative feedback defining a circadian clock: autoregulation of the clock gene frequency. Science 263:1578–1584.
- Asai, D. J., and M. P. Koonce. 2001. The dynein heavy chain: structure, mechanics and evolution. Trends Cell Biol. 11:196–202.
- Atchley, W. R., and F. W. M. 1997. A natural classification of the basic helix-loop-helix class of transcription factors. Proc. Natl. Acad. Sci. USA 94:5172–5176.
- Aubry, L., and R. Firtel. 1999. Integration of signaling networks that regulate *Dictyostelium* differentiation. Annu. Rev. Cell Dev. Biol. 15:469–517.
- Avramova, Z. V. 2002. Heterochromatin in animals and plants. Similarities and differences. Plant Physiol. 129:40–49.
 Awald, P. D., D. Frost, R. R. Drake, and C. P. Selitrennikoff. 1994.
- 35. Award, P. D., D. Frost, K. K. Drake, and C. P. Sentremnikon. 1994. (1,3)beta-Glucan synthase activity of *Neurospora crassa:* identification of a substrate-binding protein. Biochim Biophys Acta **1201**:312–320.
- 36. Azumi, Y., D. Liu, D. Zhao, W. Li, G. Wang, Y. Hu, and H. Ma. 2002. Homolog interaction during meiotic prophase 1 in *Arabidopsis* requires the SOLO DANCERS gene encoding a novel cyclin-like protein. EMBO J. 21:3081–3095.
- Baasiri, R. A., X. Lu, P. S. Rowley, G. E. Turner, and K. A. Borkovich. 1997. Overlapping functions for two G protein α subunits in *Neurospora crassa*. Genetics 147:137–145.
- Bachman, B. J., and W. N. Strickland. 1965. *Neurospora* bibliography and index. Yale University Press, New Haven, Conn.
- Bachman, K. E., M. R. Rountree, and S. B. Baylin. 2001. Dnmt3a and Dnmt3b are transcriptional repressors that exhibit unique localization properties to heterochromatin. J. Biol. Chem. 276:32282–32287.
- Bailey, L. A., and D. J. Ebbole. 1998. The *fluffy* gene of *Neurospora crassa* encodes a Gal4p-type C6 zinc cluster protein required for conidial development. Genetics 148:1813–1820.
- Baker, N., and F. Lynen. 1971. Factors involved in fatty acyl CoA desaturation by fungal microsomes. The relative roles of acyl CoA and phospholipids as substrates. Eur. J. Biochem. 19:200–210.
- Ballario, P., P. Vittorioso, A. Magrelli, C. Talora, A. Cabibbo, and G. Macino. 1996. White collar-1, a central regulator of blue light responses in *Neurospora*, is a zinc finger protein. EMBO J. 15:1650–1657.
- 43. Bao, S., Y. Qyang, P. Yang, H. Kim, H. Du, G. Bartholomeusz, J. Henkel, R. Pimental, F. Verde, and S. Marcus. 2001. The highly conserved protein methyltransferase, Skb1, is a mediator of hyperosmotic stress response in the fission yeast *Schizosaccharomyces pombe*. J. Biol. Chem. 276:14549– 14552.
- 44. Barra, J. L., L. Rhounim, J. L. Rossignol, and G. Faugeron. 2000. Histone H1 is dispensable for methylation-associated gene silencing in ascobolus immersus and essential for long life span. Mol. Cell. Biol. 20:61–69.
- Barral, Y., V. Mermall, M. S. Mooseker, and M. Snyder. 2000. Compartmentalization of the cell cortex by septins is required for maintenance of cell polarity in yeast. Mol. Cell 5:841–851.

- Barthelmess, I. B. 1982. Mutants affecting amino acid crosspathway control in *Neurospora crassa*. Genet. Res. 39:169–185.
- Bartnicki-Garcia, S. 1968. Cell wall chemistry, morphogenesis and taxonomy of filamentous fungi. Annu. Rev. Microbiol. 22:97–108.
- Bartnicki-Garcia, S., D. D. Bartnicki, G. Gierz, R. Lopez-Franco, and C. E. Bracker. 1995. Evidence that Spitzenkorper behavior determines the shape of a fungal hypha: a test of the hyphoid model. Exp. Mycol. 19:153–159.
- Bartnicki-Garcia, S., F. Hergert, and G. Gierz. 1989. Computer simulation of fungal morphogenesis and the mathimatical basis for hyphal (tip) growth. Protoplasma 153:46–57.
- Batzoglou, S., D. B. Jaffe, K. Stanley, J. Butler, S. Gnerre, E. Mauceli, B. Berger, J. P. Mesirov, and E. S. Lander. 2002. ARACHNE: a wholegenome shotgun assembler. Genome Res. 12:177–189.
- Bauer, U. M., S. Daujat, S. J. Nielsen, K. Nightingale, and T. Kouzarides. 2002. Methylation at arginine 17 of histone H3 is linked to gene activation. EMBO Rep. 3:39–44.
- Baum, J. A., R. Geever, and N. H. Giles. 1987. Expression of qa-1F activator protein: identification of upstream binding sites in the qa gene cluster and localization of the DNA-binding domain. Mol. Cell. Biol. 7:1256–1266.
- 53. Baumgartner, R., C. Fernandez-Catalan, A. Winoto, R. Huber, R. A. Engh, and T. A. Holak. 1998. Structure of human cyclin-dependent kinase inhibitor p191NK4d: comparison to known ankyrin-repeat-containing structures and implications for the dysfunction of tumor suppressor p161NK4a. Structure 6:1279–1290.
- Baylin, S. B., and J. G. Herman. 2000. DNA hypermethylation in tumorigenesis: epigenetics joins genetics. Trends Genet. 16:168–174.
- Beadle, G. W., and E. L. Tatum. 1941. Genetic control of chemical reactions in *Neurospora*. Proc. Natl. Acad. Sci. USA 27:499–506.
- 56. Bean, L. E., W. H. Dvorachek, Jr., E. L. Braun, A. Errett, G. S. Saenz, M. D. Giles, M. Werner-Washburne, M. A. Nelson, and D. O. Natvig. 2001. Analysis of the *pdx-1* (*snz-1/sno-1*) region of the *Neurospora crassa* genome: correlation of pyridoxine-requiring phenotypes with mutations in two structural genes. Genetics 157:1067–1075.
- Beauvais, A., J. M. Bruneau, P. C. Mol, M. J. Buitrago, R. Legrand, and J. P. Latge. 2001. Glucan synthase complex of *Aspergillus fumigatus*. J. Bacteriol. 183:2273–2279.
- Becker, P. B., and W. Horz. 2002. ATP-dependent nucleosome remodeling. Annu. Rev. Biochem. 71:247–273.
- Beisel, C., A. Imhof, J. Greene, E. Kremmer, and F. Sauer. 2002. Histone methylation by the *Drosophila* epigenetic transcriptional regulator Ash1. Nature 419:857–862.
- 60. Beisson, F., A. J. Koo, S. Ruuska, J. Schwender, M. Pollard, J. J. Thelen, T. Paddock, J. J. Salas, L. Savage, A. Milcamps, V. B. Mhaske, Y. Cho, and J. B. Ohlrogge. 2003. *Arabidopsis* genes involved in acyl lipid metabolism. A 2003 census of the candidates, a study of the distribution of expressed sequence tags in organs, and a web-based database. Plant Physiol. 132:681–697.
- Bell-Pedersen, D., S. K. Crosthwaite, P. L. Lakin-Thomas, M. Merrow, and M. Okland. 2001. The *Neurospora* circadian clock: simple or complex? Philos. Trans. R. Soc. London B Ser. 356:1697–1709.
- Bell-Pedersen, D., J. C. Dunlap, and J. J. Loros. 1992. The *Neurospora* circadian clock-controlled gene, *ccg-2*, is allelic to *eas* and encodes a fungal hydrophobin required for formation of the conidial rodlet layer. Genes Dev. 6:2382–2394.
- Bell-Pedersen, D., M. L. Shinohara, J. J. Loros, and J. C. Dunlap. 1996. Circadian clock-controlled genes isolated from *Neurospora crassa* are late night- to early morning-specific. Proc. Natl. Acad. Sci. USA 93:13096– 13101.
- Benito, B., B. Garciadeblas, and A. Rodriguez-Navarro. 2000. Molecular cloning of the calcium and sodium ATPases in *Neurospora crassa*. Mol. Microbiol. 35:1079–1088.
- Bennetzen, J. L., and B. D. Hall. 1982. Codon selection in yeast. J. Biol. Chem. 257:3026–3031.
- Benton, B. K., A. Tinkelenberg, I. Gonzalez, and F. R. Cross. 1997. Cla4p, a Saccharomyces cerevisiae Cdc42p-activated kinase involved in cytokinesis, is activated at mitosis. Mol. Cell. Biol. 17:5067–5076.
- Berger, S. L. 2002. Histone modifications in transcriptional regulation. Curr. Opin. Genet. Dev. 12: 142–148.
- Bergerat, A., B. de Massy, D. Gadelle, P. C. Varoutas, A. Nicolas, and P. Forterre. 1997. An atypical topoisomerase II from Archaea with implications for meiotic recombination. Nature 386:414–417.
- Berlin, V., and C. Yanofsky. 1985. Isolation and characterization of genes differentially expressed during conidiation of *Neurospora crassa*. Mol. Cell. Biol. 5:849–855.
- Berridge, M. J., M. D. Bootman, and H. L. Roderick. 2003. Calcium signalling: dynamics, homeostasis and remodelling. Nat. Rev. Mol. Cell. Biol. 4:517–529.
- Berridge, M. J., P. Lipp, and M. D. Bootman. 2000. The versatility and universality of calcium signalling. Nat. Rev. Mol. Cell. Biol. 1:11–21.
- Bertrand, H., C. A. Argan, and N. A. Szakacs. 1983. Genetic control of the biogenesis of cyanide insensitive respiration in *Neurospora crassa*, p. 495–

507. In R. J. Schweyen, K. Wolf, and F. Kaudewitz (ed.), Mitochondria. Walter de Gruyter Co., Berlin, Germany.

- Bestor, T. H. 2000. The DNA methyltransferases of mammals. Hum. Mol. Genet. 9:2395–2402.
- 74. Beth Din, A., C. A. Specht, P. W. Robbins, and O. Yarden. 1996. chs-4, a class IV chitin synthase gene from *Neurospora crassa*. Mol. Gen. Genet. 250:214–222.
- Beth Din, A., and O. Yarden. 1994. The Neurospora crassa chs-2 gene encodes a non-essential chitin synthase. Microbiology 140:2189–2197.
- Beth Din, A., and O. Yarden. 2000. The *Neurospora crassa chs-3* gene encodes an essential class I chitin synthase. Mycologia 92:65–73.
- Bhargava, J., J. Engebrecht, and G. S. Roeder. 1992. The rec102 mutant of yeast is defective in meiotic recombination and chromosome synapsis. Genetics 130:59–69.
- Bieszke, J. A., E. L. Braun, L. E. Bean, S. Kang, D. O. Natvig, and K. A. Borkovich. 1999. The *nop-1* gene of *Neurospora crassa* encodes a seven transmembrane helix retinal-binding protein homologous to archaeal rhodopsins. Proc. Natl. Acad. Sci. USA 96:8034–8039.
- Bilwes, A. M., L. A. Alex, B. R. Crane, and M. I. Simon. 1999. Structure of CheA, a signal-transducing histidine kinase. Cell 96:131–141.
- Bilwes, A. M., C. M. Quezada, L. R. Croal, B. R. Crane, and M. I. Simon. 2001. Nucleotide binding by the histidine kinase CheA. Nat. Struct. Biol. 8:353–360.
- Bistis, G. N. 1981. Chemotropic interactions between trichogynes and conidia of opposite mating-type in *Neurospora crassa*. Mycologia 73:959– 975.
- Bistis, G. N., D. D. Perkins, and N. D. Read. 2003. Cell types of *Neurospora crassa*. Fungal Genet. Newsl. 50:17–19
- Bittner-Eddy, P., A. F. Monroy, and R. Brambl. 1994. Expression of mitochondrial genes in the germinating conidia of *Neurospora crassa*. J. Mol. Biol. 235:881–897.
- Bloom, G. S. 2001. The UNC-104/KIF1 family of kinesins. Curr. Opin. Cell Biol. 13:36–40.
- Blower, M. D., and G. H. Karpen. 2001. The role of *Drosophila* CID in kinetochore formation, cell-cycle progression and heterochromatin interactions. Nat. Cell Biol. 3:730–739.
- Blower, M. D., B. A. Sullivan, and G. H. Karpen. 2002. Conserved organization of centromeric chromatin in flies and humans. Dev. Cell 2:319–330.
- Bobrowicz, P., R. Pawlak, A. Corea, D. Bell-Pedersen, and D. J. Ebbole. 2002. The *Neurospora crassa* pheromone precursor genes are regulated by the mating type locus and the circadian clock. Mol. Microbiol. 45:795–804.
- Bochtler, M., L. Ditzel, M. Groll, C. Hartmann, and R. Huber. 1999. The proteasome. Annu. Rev. Biophys. Biomol. Struct. 28:295–317.
- Boehm, M., and J. S. Bonifacino. 2001. Adaptins: the final recount. Mol. Biol. Cell 12:2907–2920.
- Boehm, M., and J. S. Bonifacino. 2002. Genetic analyses of adaptin function from yeast to mammals. Gene 286:175–186.
- Bohm, L., G. Briand, P. Sautiere, and C. Crane-Robinson. 1982. Proteolytic digestion studies of chromatin core-histone structure. Identification of limit peptides from histone H2B. Eur. J. Biochem. 123:299–303.
- Bokoch, G. M. 2003. Biology of the p21-activated kinases. Annu. Rev. Biochem. 72:743–781.
- Bolliger, L., O. Deloche, B. S. Glick, C. Georgopoulos, P. Jeno, N. Kronidou, M. Horst, N. Morishima, and G. Schatz. 1994. A mitochondrial homolog of bacterial GrpE interacts with mitochondrial hsp70 and is essential for viability. EMBO J. 13:1998–2006.
- Boone, C., S. S. Sommer, A. Hensel, and H. Bussey. 1990. Yeast KRE genes provide evidence for a pathway of cell wall beta-glucan assembly. J. Cell Biol. 110:1833–1843.
- Bootman, M. D., T. J. Collins, C. M. Peppiatt, L. S. Prothero, L. Mac-Kenzie, P. De Smet, M. Travers, S. C. Tovey, J. T. Seo, M. J. Berridge, F. Ciccolini, and P. Lipp. 2001. Calcium signalling—an overview. Semin. Cell Dev. Biol. 12:3–10.
- 96. Borneman, A. R., M. J. Hynes, and A. Andrianopoulos. 2001. A STE12 homolog from the asexual, dimorphic fungus *Penicillium marneffei* complements the defect in sexual development of an *Aspergillus nidulans steA* mutant. Genetics 157:1003–1014.
- Borts, R. H., S. R. Chambers, and M. F. Abdullah. 2000. The many faces of mismatch repair in meiosis. Mutat. Res. 451:129–150.
- Bose, I., A. J. Reese, J. J. Ory, G. Janbon, and T. L. Doering. 2003. A yeast under cover: the capsule of *Cryptococcus neoformans*. Eukaryot. Cell 2:655– 663.
- Bouhouche, N., M. Syvanen, and C. I. Kada. 2000. The origin of prokaryotic C2H2 zinc finger regulators. Trends Microbiol. 8:77–81.
- Bourc'his, D., and T. H. Bestor. 2002. Helicase homologues maintain cytosine methylation in plants and mammals. Bioessays 24:297–299.
- Boyce, K. J., M. J. Hynes, and A. Andrianopoulos. 2001. The CDC42 homolog of the dimorphic fungus *Penicillium mameffei* is required for correct cell polarization during growth but not development. J. Bacteriol. 183:3447–3457.
- 102. Boyce, K. J., M. J. Hynes, and A. Andrianopoulos. 2003. Control of mor-

phogenesis and actin localization by the *Penicillium marneffei* RAC homolog. J. Cell Sci. 116:1249–1260.

- Branscombe, T. L., A. Frankel, J. H. Lee, J. R. Cook, Z. Yang, S. Pestka, and S. Clarke. 2001. PRMT5 (Janus kinase-binding protein 1) catalyzes the formation of symmetric dimethylarginine residues in proteins. J. Biol. Chem. 276:32971–32976.
- Braunstein, M., R. E. Sobel, C. D. Allis, B. M. Turner, and J. R. Broach. 1996. Efficient transcriptional silencing in *Saccharomyces cerevisiae* requires a heterochromatin histone acetylation pattern. Mol. Cell. Biol. 16:4349– 4356.
- Bridger, W. A., W. T. Wolodko, W. Henning, C. Upton, R. Majumdar, and S. P. Williams. 1987. The subunits of succinyl-coenzyme A synthetase function and assembly. Biochem. Soc. Symp. 54:103–111.
- 106. Briggs, S. D., M. Bryk, B. D. Strahl, W. L. Cheung, J. K. Davie, S. Y. Dent, F. Winston, and C. D. Allis. 2001. Histone H3 lysine 4 methylation is mediated by Set1 and required for cell growth and rDNA silencing in *Saccharomyces cerevisiae*. Genes Dev. 15:3286–3295.
- 107. Broun, P., J. Shanklin, E. Whittle, and C. Somerville. 1998. Catalytic plasticity of fatty acid modification enzymes underlying chemical diversity of plant lipids. Science 282:1315–1317.
- Brown, D. W., T. H. Adams, and N. P. Keller. 1996. Aspergillus has distinct fatty acid synthases for primary and secondary metabolism. Proc. Natl. Acad. Sci. USA 93:14873–14877.
- 109. Brown, E. M., G. Gamba, D. Riccardi, M. Lombardi, R. Butters, O. Kifor, A. Sun, M. A. Hediger, J. Lytton, and S. C. Hebert. 1993. Cloning and characterization of an extracellular Ca²⁺-sensing receptor from bovine parathyroid. Nature 366:575–580.
- 110. Brown, J. L., Z. Kossaczka, B. Jiang, and H. Bussey. 1993. A mutational analysis of killer toxin resistance in *Saccharomyces cerevisiae* identifies new genes involved in cell wall (1→6)-beta-glucan synthesis. Genetics 133:837– 849.
- 111. Brown, S. S. 1997. Myosins in yeast. Curr. Opin. Cell Biol. 9:44-48.
- 112. Browning, H., J. Hayles, J. Mata, L. Aveline, P. Nurse, and J. R. McIntosh. 2000. Tea2p is a kinesin-like protein required to generate polarized growth in fission yeast. J. Cell Biol. 151:15–28.
- 113. Bruno, K. S., R. Aramayo, P. F. Minke, R. L. Metzenberg, and M. Plamann. 1996. Loss of growth polarity and mislocalization of septa in a *Neurospora* mutant altered in the regulatory subunit of cAMP-dependent protein kinase. EMBO J. 15:5772–5782.
- Bull, J. H., and J. C. Wootton. 1984. Heavily methylated amplified DNA in transformants of *Neurospora crassa*. Nature 310:701–704.
- Bullerwell, C. E., G. Burger, and B. F. Lang. 2000. A novel motif for identifying *rps3* homologs in fungal mitochondrial genomes. Trends Biochem. Sci. 25:363–365.
- Burda, P., and M. Aebi. 1999. The dolichol pathway of N-linked glycosylation. Biochem. Biophys. Acta 1426:239–257.
- Burger, G., and S. Werner. 1986. Mitochondrial gene URFN of Neurospora crassa codes for a long polypeptide with highly repetitive structure. J. Mol. Biol. 191:589–599.
- 118. Burks, E. A., P. P. Bezerra, H. Le, D. R. Gallie, and K. S. Browning. 2001. Plant initiation factor 3 subunit composition resembles mammalian initiation factor 3 and has a novel subunit. J. Biol. Chem. 276:2122–2131.
- Cabib, E., D. H. Roh, M. Schmidt, L. B. Crotti, and A. Varma. 2001. The yeast cell wall and septum as paradigms of cell growth and morphogenesis. J. Biol. Chem. 276:19679–19682.
- Cairns, B. R., H. Erdjument-Bromage, P. Tempst, F. Winston, and R. D. Kornberg. 1998. Two actin-related proteins are shared functional components of the chromatin-remodeling complexes RSC and SWI/SNF. Mol. Cell 2:639–651.
- 121. Cambareri, E. B., R. Aisner, and J. Carbon. 1998. Structure of the chromosome VII centromere region in *Neurospora crassa:* degenerate transposons and simple repeats. Mol. Cell. Biol. 18:5465–5477.
- 122. Cambareri, E. B., J. Helber, and J. A. Kinsey. 1994. Tad-1, an active LINE-like element of *Neurospora crassa*. Mol. Gen. Genet. 242:658–665.
- Cambareri, E. B., B. C. Jensen, E. Schabtach, and E. U. Selker. 1989. Repeat-induced G-C to A-T mutations in *Neurospora*. Science 244:1571– 1575.
- Cambareri, E. B., M. J. Singer, and E. U. Selker. 1991. Recurrence of repeat-induced point mutation (RIP) in *Neurospora crassa*. Genetics 127: 699–710.
- 125. Capelli, N., D. van Tuinen, R. Ortega Perez, J. F. Arrighi, and G. Turian. 1993. Molecular cloning of a cDNA encoding calmodulin from *Neurospora crassa*. FEBS Lett. 321:63–68.
- 126. Caplan, A. J., J. Tsai, P. J. Casey, and M. G. Douglas. 1992. Farnesylation of YDJ1p is required for function at elevated growth temperatures in *Saccharomyces cerevisiae*. J. Biol. Chem. 267:18890–18895.
- Carbone, I., and L. M. Kohn. 1999. A method for designing primer sets for speciation studies in filamentous ascomycetes. Mycologia 91:553–556.
- Carman, G. M., and S. A. Henry. 1999. Phospholipid biosynthesis in the yeast *Saccharomyces cerevisiae* and interrelationship with other metabolic processes. Prog. Lipid Res. 38:361–399.
- 129. Carotti, C., L. Ferrario, C. Roncero, M. H. Valdivieso, A. Duran, and L.

Popolo. 2002. Maintenance of cell integrity in the *gas1* mutant of *Saccharomyces cerevisiae* requires the Chs3p-targeting and activation pathway and involves an unusual Chs3p localization. Yeast **19**:1113–1124.

- 130. Carr, A. M., S. M. Dorrington, J. Hindley, G. A. Phear, S. J. Aves, and P. Nurse. 1994. Analysis of a histone H2A variant from fission yeast: evidence for a role in chromosome stability. Mol. Gen. Genet. 245:628–635.
- Catalanotto, C., G. Azzalin, G. Macino, and C. Cogoni. 2000. Gene silencing in worms and fungi. Nature 404:245.
- 132. Catalanotto, C., G. Azzalin, G. Macino, and C. Cogoni. 2002. Involvement of small RNAs and role of the *qde* genes in the gene silencing pathway in *Neurospora*. Genes Dev. 16:790–795.
- Catcheside, D. E. A. 1986. A restriction and modification model for the initiation and control of recombination in *Neurospora*. Genet. Res. Camb. 47:157–165.
- Catcheside, D. G. 1977. The genetics of recombination. Edward Arnold, London, United Kingdom.
- 135. Catty, P., A. de Kerchove d'Exaerde, and A. Goffeau. 1997. The complete inventory of the yeast *Saccharomyces cerevisiae* P-type transport ATPases. FEBS Lett. 409:325–332.
- Cech, T. R. 1991. RNA editing: world's smallest introns? Cell 64:667–669.
 Centola, M., and J. Carbon. 1994. Cloning and characterization of centro-
- meric DNA from *Neurospora crassa*. Mol. Cell. Biol. 14:1510–1519.
 138. Chang, L., and M. Karin. 2001. Mammalian MAP kinase signalling cascades. Nature 410:37–40.
- Chang, L., S. S. Loranger, C. Mizzen, S. G. Ernst, C. D. Allis, and A. T. Annunziato. 1997. Histones in transit: cytosolic histone complexes and diacetylation of H4 during nucleosome assembly in human cells. Biochemistry 36:469–80.
- Chavrier, P., and B. Goud. 1999. The role of ARF and Rab GTPases in membrane transport. Curr. Opin. Cell Biol. 11:466–475.
- 141. Cheeseman, I. M., C. Brew, M. Wolyniak, A. Desai, S. Anderson, N. Muster, J. R. Yates, T. C. Huffaker, D. G. Drubin, and G. Barnes. 2001. Implication of a novel multiprotein Dam1p complex in outer kinetochore function. J. Cell Biol. 155:1137–1145.
- 142. Cheeseman, I. M., D. G. Drubin, and G. Barnes. 2002. Simple centromere, complex kinetochore: linking spindle microtubules and centromeric DNA in budding yeast. J. Cell Biol. 157:199–203.
- 143. Chen, Y., R. E. Baker, K. C. Keith, K. Harris, S. Stoler, and M. Fitzgerald-Hayes. 2000. The N terminus of the centromere H3-like protein Cse4p performs an essential function distinct from that of the histone fold domain. Mol. Cell. Biol. 20:7037–7048.
- 144. Cheng, P., Q. He, Y. Yang, L. Wang, and Y. Liu. 2003. Functional conservation of light, oxygen, or voltage domains in light sensing. Proc. Natl. Acad. Sci. USA 100:5938–5943.
- 145. Cheung, P., C. D. Allis, and P. Sassone-Corsi. 2000. Signaling to chromatin through histone modifications. Cell 103:263–271.
- 146. Chiang, T.-Y., and G. A. Marzluf. 1994. DNA Recognition by the NIT2 nitrogen regulatory protein: importance of the number, spacing, and orientation of GATA core elements and their flanking sequences upon NIT2 binding. Biochemistry 33:576–582.
- 147. Chu, S., and I. Herskowitz. 1998. Gametogenesis in yeast is regulated by a transcriptional cascade dependent on Ndt80. Mol. Cell 1:685–696.
- 148. Chua, P. R., and G. S. Roeder. 1998. Zip2, a meiosis-specific protein required for the initiation of chromosome synapsis. Cell 93:349–359.
- 149. Chung, K. S., M. Won, S. B. Lee, Y. J. Jang, K. L. Hoe, D. U. Kim, J. W. Lee, K. W. Kim, and H. S. Yoo. 2001. Isolation of a novel gene from *Schizosac-charomyces pombe: stm1*⁺ encoding a seven-transmembrane loop protein that may couple with the heterotrimeric Galpha 2 protein, Gpa2. J. Biol. Chem. 276:40190–40201.
- Cid, V. J., M. J. Shulewitz, K. L. McDonald, and J. Thorner. 2001. Dynamic localization of the Swe1 regulator Hs17 during the *Saccharomyces cerevisiae* cell cycle. Mol. Biol. Cell 12:1645–1669.
- 151. Citterio, E., V. Van Den Boom, G. Schnitzler, R. Kanaar, E. Bonte, R. E. Kingston, J. H. Hoeijmakers, and W. Vermeulen. 2000. ATP-dependent chromatin remodeling by the Cockayne syndrome B DNA repair-transcription-coupling factor. Mol. Cell. Biol. 20:7643–7653.
- Claros, M. G., and P. Vincens. 1996. Computational method to predict mitochondrially imported proteins and their targeting sequences. Eur J. Biochem. 241:779–786.
- Clemons, K. V., T. K. Miller, C. P. Selitrennikoff, and D. A. Stevens. 2002. fos-1, a putative histidine kinase as a virulence factor for systemic aspergillosis. Med. Mycol. 40:259–262.
- Cleveland, D. W., Y. Mao, and K. F. Sullivan. 2003. Centromeres and kinetochores: from epigenetics to mitotic checkpoint signaling. Cell 112: 407–421.
- 155. Coffman, J. A., R. Rai, T. Cunningham, V. Svetlov, and T. G. Cooper. 1996. Gat1p, a GATA family protein whose production is sensitive to nitrogen catabolite repression, participates in transcriptional activation of nitrogencatabolite genes in *Saccharomyces cerevisiae*. Mol. Cell. Biol. 16:847–858.
- Coghlan, A., and K. H. Wolfe. 2000. Relationship of codon bias to mRNA concentration and protein length in *Saccharomyces cerevisiae*. Yeast 16: 1131–1145.

- Cogoni, C. 2001. Homology-dependent gene silencing mechanisms in fungi. Annu. Rev. Microbiol. 55:381–406.
- 158. Cogoni, C. 2002. Unifying homology effects. Nat. Genet. 30:245-246.
- 159. Cogoni, C., J. T. Irelan, M. Schumacher, T. J. Schmidhauser, E. U. Selker, and G. Macino. 1996. Transgene silencing of the *al-1* gene in vegetative cells of *Neurospora* is mediated by a cytoplasmic effector and does not depend on DNA-DNA interactions or DNA methylation. EMBO J. 15:3153– 3163.
- Cogoni, C., and G. Macino. 1999. Gene silencing in *Neurospora crassa* requires a protein homologous to RNA-dependent RNA polymerase. Nature 399:166–169.
- 161. Cogoni, C., and G. Macino. 1997. Isolation of quelling-defective (qde) mutants impaired in posttranscriptional transgene-induced gene silencing in *Neurospora crassa*. Proc. Natl. Acad. Sci. USA 94:10233–10238.
- Cogoni, C., and G. Macino. 2000. Post-transcriptional gene silencing across kingdoms. Curr. Opin. Genet. Dev. 10:638–643.
- 163. Cogoni, C., and G. Macino. 1999. Posttranscriptional gene silencing in *Neurospora* by a RecQ DNA helicase. Science 286:2342–2344.
- 164. Cohen, P. T. 1997. Novel protein serine/threonine phosphatases: variety is the spice of life. Trends Biochem. Sci. 22:245–251.
- Cokol, M., R. Nair, and B. Rost. 2000. Finding nuclear localization signals. EMBO Rep. 1:411–415.
- 166. Colaiacovo, M. P., F. Paques, and J. E. Haber. 1999. Removal of one nonhomologous DNA end during gene conversion by a RAD1- and MSH2independent pathway. Genetics 151:1409–1423.
- 167. Conrad, M. N., A. M. Dominguez, and M. E. Dresser. 1997. Ndj1p, a meiotic telomere protein required for normal chromosome synapsis and segregation in yeast. Science 276:1252–1255.
- Cooper, J. D. 2003. Progress towards understanding the neurobiology of Batten disease or neuronal ceroid lipofuscinosis. Curr. Opin. Neurol. 16: 121–128.
- Coppin, E., R. Debuchy, S. Arnaise, and M. Picard. 1997. Mating types and sexual development in filamentous ascomycetes. Microbiol Mol. Biol. Rev. 61:411–428.
- Cornelius, G., G. Gebauer, and D. Techel. 1989. Inositol trisphosphate induces calcium release from *Neurospora crassa* vacuoles. Biochem. Biophys. Res. Commun. 162:852–856.
- 171. Craig, K. L., and M. Tyers. 1999. The F-box: a new motif for ubiquitin dependent proteolysis in cell cycle regulation and signal transduction. Prog. Biophys. Mol. Biol. 72:299–328.
- 172. Crosio, C., G. M. Fimia, R. Loury, M. Kimura, Y. Okano, H. Zhou, S. Sen, C. D. Allis, and P. Sassone-Corsi. 2002. Mitotic phosphorylation of histone H3: spatio-temporal regulation by mammalian Aurora kinases. Mol. Cell. Biol. 22:874–885.
- Crosthwaite, S. K., J. C. Dunlap, and J. J. Loros. 1997. Neurospora wc-1 and wc-2: transcription, photoresponses, and the origins of circadian rhythmicity. Science 276:763–769.
- 174. Crowhurst, R. N., S. J. Binnie, J. K. Bowen, B. T. Hawthorne, K. M. Plummer, J. Rees-George, E. H. Rikkerink, and M. D. Templeton. 1977. Effect of disruption of a cutinase gene (*cutA*) on virulence and tissue specificity of *Fusarium solani* f. sp. cucurbitae race 2 toward *Cucurbita maxima* and *C. moschata*. Mol. Plant-Microbe Interact. 10:355–368.
- 175. Cunningham, T. S., and T. G. Cooper. 1993. The Saccharomyces cerevisiae Da180 repressor binds to multiple copies of GATAA-containing sequences (URS GATA). J. Bacteriol. 175:5851–5861.
- 176. Cvrckova, F., C. De Virgilio, E. Manser, J. R. Pringle, and K. Nasmyth. 1995. Ste20-like protein kinases are required for normal localization of cell growth and for cytokinesis in budding yeast. Genes Dev. 9:1817–1830.
- Dan, I., N. M. Watanabe, and A. Kusumi. 2001. The Ste20 group kinases as regulators of MAP kinase cascades. Trends Cell Biol. 11:220–230.
- Dasgupta, A., R. P. Darst, K. J. Martin, C. A. Afshari, and D. T. Auble. 2002. Mot1 activates and represses transcription by direct, ATPase-dependent mechanisms. Proc. Natl. Acad. Sci. USA 99:2666–2671.
- 179. Daubresse, G., R. Deuring, L. Moore, O. Papoulas, I. Zakrajsek, W. R. Waldrip, M. P. Scott, J. A. Kennison, and J. W. Tamkun. 1999. The Drosophila kismet gene is related to chromatin-remodeling factors and is required for both segmentation and segment identity. Development 126: 1175–1187.
- Daujat, S., U. M. Bauer, V. Shah, B. Turner, S. Berger, and T. Kouzarides. 2002. Crosstalk between CARM1 methylation and CBP acetylation on histone H3. Curr. Biol. 12:2090–2097.
- Daum, G., N. D. Lees, M. Bard, and R. Dickson. 1998. Biochemistry, cell biology and molecular biology of lipids of *Saccharomyces cerevisiae*. Yeast 14:1471–1510.
- 182. Davies, J. P., F. H. Yildiz, and A. R. Grossman. 1999. Sac3, an Snf1-like serine/threonine kinase that positively and negatively regulates the responses of *Chlamydomonas* to sulfur limitation. Plant Cell 11:1179–1190.
- 183. Davis, C. R., R. R. Kempainen, M. S. Srodes, and C. R. McClung. 1994. Correlation of the physical and genetic maps of the centromeric region of the right arm of linkage group III of *Neurospora crassa*. Genetics 136:1297– 1306.

- Davis, R. H. 2000. *Neurospora*: contributions of a model organism. Oxford University Press, New York, N.Y.
- 185. Davis, R. H., and D. D. Perkins. 2002. Timeline: *Neurospora*: a model of model microbes. Nat. Rev. Genet. 3:397–403.
- de los Santos, T., and N. M. Hollingsworth. 1999. Red1p, a MEK1-dependent phosphoprotein that physically interacts with Hop1p during meiosis in yeast. J. Biol. Chem. 274:1783–1790.
- Deneka, M., M. Neeft, and P. van der Sluijs. 2003. Regulation of membrane transport by rab GTPases. Crit. Rev. Biochem. Mol. Biol. 38:121–142.
- 188. d'Enfert, C., B. M. Bonini, P. D. Zapella, T. Fontaine, A. M. da Silva, and H. F. Terenzi. 1999. Neutral trehalases catalyse intracellular trehalose breakdown in the filamentous fungi *Aspergillus nidulans* and *Neurospora crassa*. Mol. Microbiol. 32:471–483.
- Dennis, K., T. Fan, T. Geiman, Q. Yan, and K. Muegge. 2001. Lsh, a member of the SNF2 family, is required for genome-wide methylation. Genes Dev. 15:2940–2944.
- 190. de Ruijter, A. J., A. H. van Gennip, H. N. Caron, S. Kemp, and A. B. van Kuilenburg. 2003. Histone deacetylases (HDACs): characterization of the classical HDAC family. Biochem. J. 370:737–749.
- 191. Deshaies, R. J., B. D. Koch, M. Werner-Washburne, E. A. Craig, and R. Schekman. 1988. A subfamily of stress proteins facilitates translocation of secretory and mitochondrial precursor polypeptides. Nature 332:800–805.
- 192. De Souza, C. P., A. H. Osmani, L. P. Wu, J. L. Spotts, and S. A. Osmani. 2000. Mitotic histone H3 phosphorylation by the NIMA kinase in *Aspergillus nidulans*. Cell **102**:293–302.
- 193. Deuring, R., L. Fanti, J. A. Armstrong, M. Sarte, O. Papoulas, M. Prestel, G. Daubresse, M. Verardo, S. L. Moseley, M. Berloco, T. Tsukiyama, C. Wu, S. Pimpinelli, and J. W. Tamkun. 2000. The ISWI chromatin-remodeling protein is required for gene expression and the maintenance of higher order chromatin structure in vivo. Mol. Cell 5:355–365.
- Devlin, P. F., and S. A. Kay. 2001. Circadian photoperception. Annu. Rev. Physiol. 63:677–694.
- 195. DeZwaan, T. M., A. M. Carroll, B. Valent, and J. A. Sweigard. 1999. Magnaporthe grisea pth11p is a novel plasma membrane protein that mediates appressorium differentiation in response to inductive substrate cues. Plant Cell 11:2013–2030.
- Dickman, M. B., and O. Yarden. 1999. Serine/threonine protein kinases and phosphatases in filamentious fungi. Fungal Genet. Biol. 26:99–117.
- 197. Dixon, D. P., A. Lapthorn, and R. Edwards. 2002. Plant glutathione transferases. Genome Biol. 3:3004.1–3004.10.
- Dohlman, H. G., J. Thorner, M. G. Caron, and R. J. Lefkowitz. 1991. Model systems for the study of seven-transmembrane-segment receptors. Annu. Rev. Biochem. 60:653–688.
- Dohlman, H. G., and J. W. Thorner. 2001. Regulation of G protein-initiated signal transduction in yeast: paradigms and principles. Annu. Rev. Biochem. 70:703–754.
- Dolan, J. W., C. Kirkman, and S. Fields. 1989. The yeast STE12 protein binds to the DNA sequence mediating pheromone induction. Proc. Natl. Acad. Sci. USA 86:5703–5707.
- Dolinski, K. J., M. E. Cardenas, and J. Heitman. 1998. CNS1 encodes an essential p60/Sti1 homolog in *Saccharomyces cerevisiae* that suppresses cyclophilin 40 mutations and interacts with Hsp90. Mol. Cell. Biol. 18:7344– 7352.
- Dombrádi, V. 1997. Comparative analysis of Ser/Thr protein phosphatases. Trends Comp. Biochem. 3:23–48.
- 203. Drees, B. L., B. Sundin, E. Brazeau, J. P. Caviston, G. C. Chen, W. Guo, K. G. Kozminski, M. W. Lau, J. J. Moskow, A. Tong, L. R. Schenkman, A. McKenzie III, P. Brennwald, M. Longtine, E. Bi, C. Chan, P. Novick, C. Boone, J. R. Pringle, T. N. Davis, S. Fields, and D. G. Drubin. 2001. A protein interaction map for cell polarity development. J. Cell Biol. 154:549–571.
- Drewes, G., A. Ebneth, and E. M. Mandelkow. 1998. MAPs, MARKs and microtubule dynamics. Trends Biochem. Sci. 23:307–311.
 Duina, A. A., H. C. Chang, J. A. Marsh, S. Lindquist, and R. F. Gaber. 1996.
- Duina, A. A., H. C. Chang, J. A. Marsh, S. Lindquist, and R. F. Gaber. 1996. A cyclophilin function in Hsp90-dependent signal transduction. Science 274:1713–1715.
- 206. Dunlap, J. C. 1999. Molecular bases for circadian clocks. Cell 96:271-290.
- Duret, L., and L. D. Hurst. 2001. The elevated GC content at exonic third sites is not evidence against neutralist models of isochore evolution. Mol. Biol. Evol. 18:757–762.
- Duret, L., and D. Mouchiroud. 1999. Expression pattern and, surprisingly, gene length shape codon usage in *Caenorhabditis*, *Drosophila*, and *Arabidopsis*. Proc. Natl. Acad. Sci. USA 96:4482–4487.
- Durrenberger, F., K. Wong, and J. W. Kronstad. 1998. Identification of a cAMP-dependent protein kinase catalytic subunit required for virulence and morphogenesis in *Ustilago maydis*. Proc Natl Acad Sci USA 95:5684– 5689.
- Eberharter, A., S. Ferrari, G. Langst, T. Straub, A. Imhof, P. Varga-Weisz, M. Wilm, and P. B. Becker. 2001. Acf1, the largest subunit of CHRAC, regulates ISWI-induced nucleosome remodelling. EMBO J. 20:3781–3788.
- Eckley, D. M., S. R. Gill, K. A. Melkonian, J. B. Bingham, H. V. Goodson, J. E. Heuser, and T. A. Schroer. 1999. Analysis of dynactin subcomplexes

reveals a novel actin-related protein associated with the arp1 minifilament pointed end. J. Cell Biol. **147:**307–320.

- Ehrnsperger, M., S. Graber, M. Gaestel, and J. Buchner. 1997. Binding of non-native protein to Hsp25 during heat shock creates a reservoir of folding intermediates for reactivation. EMBO J. 16:221–229.
- Eichhorn, E., J. R. van der Ploeg, M. A. Kertesz, and T. Leisinger. 1997. Characterization of alpha-ketoglutarate-dependent taurine dioxygenase from *Escherichia coli*. J. Biol. Chem. 272:23031–23036.
- Eichhorn, E., J. R. van der Ploeg, and T. Leisinger. 1999. Characterization of a two-component alkanesulfonate monooxygenase from *Escherichia coli*. J. Biol. Chem. 274:26639–26646.
- Eichinger, L., M. Bahler, M. Dietz, C. Eckerskorn, and M. Schleicher. 1998. Characterization and cloning of a *Dictyostelium* Ste20-like protein kinase that phosphorylates the actin-binding protein severin. J. Biol. Chem. 273: 12952–12959.
- 216. Eilers, T., G. Schwarz, H. Brinkmann, C. Witt, T. Richter, J. Nieder, B. Koch, R. Hille, R. Hansch, and R. R. Mendel. 2001. Identification and biochemical characterization of *Arabidopsis thaliana* sulfite oxidase. A new player in plant sulfur metabolism. J. Biol. Chem. 276:46989–46994.
- Eisen, J. A., K. S. Sweder, and P. C. Hanawalt. 1995. Evolution of the SNF2 family of proteins: subfamilies with distinct sequences and functions. Nucleic Acids Res. 23:2715–2723.
- 218. Elion, E. A. 2001. The Ste5p scaffold. J. Cell Sci. 114:3967-3978.
- Elovson, J. 1975. Purification and properties of the fatty acid synthetase complex from *Neurospora crassa*, and the nature of the *fas* mutation. J. Bacteriol. 124:524–533.
- 220. Emanuelsson, O., H. Nielsen, S. Brunak, and G. von Heijne. 2000. Predicting subcellular localization of proteins based on their N-terminal amino acid sequence. J. Mol. Biol. 300:1005–1016.
- 221. Endo, A., K. Kakiki, and T. Misato. 1970. Feedback inhibition of L-glutamine D-fructose 6-phosphate amidotransferase by uridine diphosphate N-acetylglucosamine in Neurospora crassa. J. Bacteriol. 103:588–594.
- 222. Errede, B. E., and G. Ammerer. 1989. STE12 a protein involved in cell type specific transcription and signal transduction in yeast, is part of protein-DNA complexes. Genes Dev. 3:1349–1361.
- 223. Eshed, Y., S. F. Baum, and J. L. Bowman. 1999. Distinct mechanisms promote polarity establishment in carpels of *Arabidopsis*. Cell 99:199–209.
- Etienne-Manneville, S., and A. Hall. 2002. Rho GTPases in cell biology. Nature 420:629–635.
- 225. Evans, C. S., and J. N. Hedger. 2001. Degradation of plant cell wall polymers, p. 1–26. *In* G. M. Gadd (ed.), Fungi in bioremediation. Cambridge University Press, Cambridge, United kingdom.
- Evans, T., and G. Felsenfeld. 2003. Differential expansion of zinc-finger transcription factor loci in homologous human and mouse gene clusters. Cell 58:877–885.
- 227. Fabbrizio, E., S. El Messaoudi, J. Polanowska, C. Paul, J. R. Cook, J. H. Lee, V. Negre, M. Rousset, S. Pestka, A. Le Cam, and C. Sardet. 2002. Negative regulation of transcription by the type II arginine methyltransferase PRMT5. EMBO Rep. 3:641–645.
- 228. Fassler, J. D., A. Landsman, A. Acharya, J. R. Moll, M. Bonovich, and C. Vinson. 2002. B-ZIP proteins encoded by the *Drosophila* genome: evaluation of potential dimerization partners. Genome Res. 12:1190–1200.
- 229. Feng, B., H. Haas, and G. A. Marzluf. 2000. ASD4, a new GATA factor of *Neurospora crassa*, displays sequence-specific DNA binding and functions in ascus and ascospore development. Biochemistry **39**:11065–11073.
- Feng, B., and V. Marzluf. 1998. Interaction between major nitrogen regulatory protein NIT2 and pathway-specific regulatory factor NIT4 is required for their synergistic activation of gene expression in *Neurospora crassa*. Mol. Cell. Biol. 18:3983–3990.
- 231. Feng, B., X. Xiadong, and G. A. Marzluf. 1993. Recognition of specific nucleotide bases and cooperative DNA binding by the trans-acting nitrogen regulatory protein NIT2 of *Neurospora crassa*. Nucleic Acids Res. 21:3989– 3996.
- 232. Feng, Q., H. Wang, H. H. Ng, H. Erdjument-Bromage, P. Tempst, K. Struhl, and Y. Zhang. 2002. Methylation of H3-lysine 79 is mediated by a new family of HMTases without a SET domain. Curr. Biol. 12:1052–1058.
- 233. Ferreira, A. V., S. Saupe, and N. L. Glass. 1996. Transcriptional analysis of the mtA idiomorph of *Neurospora crassa* identifies two genes in addition to mtA-1. Mol. Gen. Genet. 250:767–774.
- Fields, S., and I. Herskowitz. 1985. The yeast STE12 product is required for expression of two sets of cell type specific genes. Cell 42:923–930.
- 235. Finger, F. P., T. E. Hughes, and P. Novick. 1998. Sec3p is a spatial landmark for polarized secretion in budding yeast. Cell 92:559–571.
- Finger, F. P., and P. Novick. 1998. Spatial regulation of exocytosis: lessons from yeast. J. Cell Biol. 142:609–612.
- Firon, A., and C. d'Enfert. 2002. Identifying essential genes in fungal pathogens of humans. Trends Microbiol 10:456–462.
- Fischer, M., N. Schnell, J. Chattaway, P. Davies, G. Dixon, and D. Sanders. 1997. The Saccharomyces cerevisiae CCH1 gene is involved in calcium influx and mating. FEBS Lett. 419:259–262.
- 239. Fischer-Parton, S., R. M. Parton, P. C. Hickey, J. Dijksterhuis, H. A. Atkinson, and N. D. Read. 2000. Confocal microscopy of FM4–64 as a tool

for analysing endocytosis and vesicle trafficking in living fungal hyphae. 20 J. Microsc. **198:**246–259.

- 240. Flaus, A., and T. Owen-Hughes. 2001. Mechanisms for ATP-dependent chromatin remodelling. Curr. Opin. Genet. Dev. 11:148–154.
- Flores, C. L., C. Rodriguez, T. Petit, and C. Gancedo. 2000. Carbohydrate and energy-yielding metabolism in non-conventional yeasts. FEMS Microbiol. Rev. 24:507–529.
- 242. Folco, H. D., M. Freitag, A. Ramon, E. D. Temporini, M. E. Alvarez, I. Garcia, C. Scazzocchio, E. U. Selker, and A. L. Rosa. 2003. Histone H1 is required for proper regulation of pyruvate decarboxylase gene expression in *Neurospora crassa*. Eukaryot. Cell 2:341–350.
- 243. Fontaine, T., C. Simenel, G. Dubreucq, O. Adam, M. Delepierre, J. Lemoine, C. E. Vorgias, M. Diaquin, and J. P. Latge. 2000. Molecular organization of the alkali-insoluble fraction of *Aspergillus fumigatus* cell wall. J. Biol. Chem. 275:27594–27607.
- Foos, T. M., and J. Y. Wu. 2002. The role of taurine in the central nervous system and the modulation of intracellular calcium homeostasis. Neurochem. Res. 27:21–26.
- Formosa, T. 2003. Changing the DNA landscape: putting a SPN on chromatin. Curr. Top. Microbiol. Immunol. 274:171–201.
- Foss, H. M., C. J. Roberts, K. M. Claeys, and E. U. Selker. 1993. Abnormal chromosome behavior in *Neurospora mutants* defective in DNA methylation. Science 262:1737–1741.
- 247. Foss, H. M., C. J. Roberts, K. M. Claeys, and E. U. Selker. 1995. Abnormal chromosome behavior in *Neurospora* mutants defective in DNA methylation Science 267:316. (Erratum.)
- 248. Fossa, A., A. Beyer, E. Pfitzner, B. Wenzel, and W. H. Kunau. 1995. Molecular cloning, sequencing and sequence analysis of the *fox-2* gene of *Neurospora crassa* encoding the multifunctional beta-oxidation protein. Mol. Gen. Genet. 247:95–104.
- Frankel, A., and S. Clarke. 2000. PRMT3 is a distinct member of the protein arginine N-methyltransferase family. Conferral of substrate specificity by a zinc-finger domain. J. Biol. Chem. 275:32974–32982.
- Fraser, M. J., T. Y. Chow, H. Cohen, and H. Koa. 1986. An immunochemical study of *Neurospora* nucleases. Biochem. Cell Biol. 64:106–116.
- 251. Free, S. J., P. W. Rice, and R. L. Metzenberg. 1979. Arrangement of the genes coding for ribosomal ribonucleic acids in *Neurospora crassa*. J. Bacteriol. 137:1219–1226.
- Freidkin, I., and D. J. Katcoff. 2001. Specific distribution of the Saccharomyces cerevisiae linker histone homolog HHO1p in the chromatin. Nucleic Acids Res. 29:4043–4051.
- 253. Freitag, M., R. L. Williams, G. O. Kothe, and E. U. Selker. 2002. A cytosine methyltransferase homologue is essential for repeat-induced point mutation in *Neurospora crassa*. Proc. Natl. Acad. Sci. USA 99:8802–8807.
- 254. Friedberg, E. C., G. C. Walker, and W. Siede. 1995. DNA repair and mutagenesis. Academic Press, Inc., San Diego, Calif.
- 255. Friesen, H., R. Lunz, S. Doyle, and J. Segall. 1994. Mutation of the SPS1encoded protein kinase of *Saccharomyces cerevisiae* leads to defects in transcription and morphology during spore formation. Genes Dev. 8:2162– 2175.
- Froehlich, A. C., Y. Liu, J. J. Loros, and J. C. Dunlap. 2002. White Collar-1, a circadian blue light photoreceptor, binding to the frequency promoter. Science 297:815–819.
- 257. Fu, Y. H., B. Feng, S. Evans, and G. A. Marzluf. 1995. Sequence-specific DNA binding by NIT4, the pathway-specific regulatory protein that mediates nitrate induction in *Neurospora*. Mol. Microbiol. 15:935–942.
- Fu, Y. H., J. Y. Knessi, and G. A. Marzluf. 1989. Isolation of *nit-4*, the minor nitrogen regulatory gene which mediates nitrate induction in *Neurospora crassa*. J. Bacteriol. 171:4067–4070.
- 259. Fu, Y. H., and G. A. Marzluf. 1987. Molecular cloning and analysis of the regulation of *nit-3*, the structural gene for nitate reductase in *Neurospora crassa*. Proc. Natl. Acad. Sci. USA 84:8243–8247.
- 260. Fu, Y.-H., J. V. Paietta, D. G. Mannix, and G. A. Marzluf. 1989. cys-3, the positive-acting sulfur regulatory gene of *Neurospora crassa*, encodes a protein with a putative leucine zipper DNA-binding element. Mol. Cell. Biol. 9:1120–1127.
- 261. Fu, Y. H., J. L. Young, and G. A. Marzluf. 1988. Molecular cloning and characterization of a negative-acting regulatory gene of *Neurospora crassa*. Mol. Gen. Genet. 214:74–79.
- 262. Fu, Y.-SH., and G. A. Marzluf. 1990. nit-2, the major nitrogen regulatory gene of *Neurospora crassa*, encodes a protein with a putative zinc finger DNA-binding domain. Mol. Cell. Biol. 10:1056–1065.
- 263. Fujiwara, M., H. Horiuchi, A. Ohta, and M. Takagi. 1997. A novel fungal gene encoding chitin synthase with a myosin motor-like domain. Biochem. Biophys. Res. Commun. 236:75–78.
- 264. Fuks, F., P. J. Hurd, R. Deplus, and T. Kouzarides. 2003. The DNA methyltransferases associate with HP1 and the SUV39H1 histone methyl-transferase. Nucleic Acids Res. 31:2305–2312.
- 265. Funabiki, H., and A. W. Murray. 2000. The *Xenopus* chromokinesin Xkid is essential for metaphase chromosome alignment and must be degraded to allow anaphase chromosome movement. Cell **102**:411–424.

- 266. Futcher, B., G. I. Latter, P. Monardo, C. S. McLaughlin, and J. I. Garrels. 1999. A sampling of the yeast proteome. Mol. Cell. Biol. 19:7357–7368.
- 267. Gadd, G. M. 1994. Signal transduction in fungi, p. 183–210. *In* N. A. R. Gow and G. M. Gadd (ed.), The growing fungus. Chapman & Hall, London, United Kingdom.
- Callus-Durner, V., J. Xie, C. Chintamaneni, and A. K. Vershon. 1996. Participation of the yeast activator Abf1 in meiosis-specific expression of the HOP1 gene. Mol. Cell. Biol. 16:2777–2786.
- 269. Galagan, J. E., S. E. Calvo, K. A. Borkovich, E. U. Selker, N. D. Read, D. Jaffe, W. FitzHugh, L. J. Ma, S. Smirnov, S. Purcell, B. Rehman, T. Elkins, R. Engels, S. Wang, C. B. Nielsen, J. Butler, M. Endrizzi, D. Qui, P. Ianakiev, D. Bell-Pedersen, M. A. Nelson, M. Werner-Washburne, C. P. Selitrennikoff, J. A. Kinsey, E. L. Braun, A. Zelter, U. Schulte, G. O. Kothe, G. Jedd, W. Mewes, C. Staben, E. Marcotte, D. Greenberg, A. Roy, K. Foley, J. Naylor, N. Stange-Thomann, R. Barrett, S. Gnerre, M. Kamal, M. Kamvysselis, E. Mauceli, C. Bielke, S. Rudd, D. Frishman, S. Krystofova, C. Rasmussen, R. L. Metzenberg, D. D. Perkins, S. Kroken, C. Cogoni, G. Macino, D. Catcheside, W. Li, R. J. Pratt, S. A. Osmani, C. P. DeSouza, L. Glass, M. J. Orbach, J. A. Berglund, R. Voelker, O. Yarden, M. Plamann, S. Seiler, J. Dunlap, A. Radford, R. Aramayo, D. O. Natvig, L. A. Alex, G. Mannhaupt, D. J. Ebbole, M. Freitag, I. Paulsen, M. S. Sachs, E. S. Lander, C. Nusbaum, and B. Birren. 2003. The genome sequence of the filamentous fungus *Neurospora crassa*. Nature 422:859–868.
- Galbraith, A. M., S. A. Bullard, K. Jiao, J. J. Nau, and R. E. Malone. 1997. Recombination and the progression of meiosis in *Saccharomyces cerevisiae*. Genetics 146:481–489.
- 271. Gangloff, Y. G., C. Romier, S. Thuault, S. Werten, and I. Davidson. 2001. The histone fold is a key structural motif of transcription factor TFIID. Trends Biochem. Sci. 26:250–257.
- Gao-Rubinelli, F., and G. A. Marzluf. 2004. Identification and characterization of a nitrate transporter gene in *Neurospora crassa*. Biochem. Genet. 42:21–34.
- 273. Gardiner, J. M., S. A. Bullard, C. Chrome, and R. E. Malone. 1997. Molecular and genetic analysis of REC103, an early meiotic recombination gene in yeast. Genetics 146:1265–1274.
- 274. Gardner, R. D., A. Poddar, C. Yellman, P. A. Tavormina, M. C. Monteagudo, and D. J. Burke. 2001. The spindle checkpoint of the yeast Saccharomyces cerevisiae requires kinetochore function and maps to the CBF3 domain. Genetics 157:1493–1502.
- Gasser, S. M., and M. M. Cockell. 2001. The molecular biology of the SIR proteins. Gene 279:1–16.
- Geli, M. I., and H. Riezman. 1998. Endocytic internalization in yeast and animal cells: similar and different. J. Cell Sci. 111:1031–1037.
- 277. Gendrel, A. V., Z. Lippman, C. Yordan, V. Colot, and R. A. Martienssen. 2002. Dependence of heterochromatic histone H3 methylation patterns on the *Arabidopsis* gene *DDM1*. Science 297:1871–1873
- Gething, M. J., and J. Sambrook. 1992. Protein folding in the cell. Nature 355:33–45.
- Ghannoum, M. A., and L. B. Rice. 1999. Antifungal agents: mode of action, mechanisms of resistance, and correlation of these mechanisms with bacterial resistance. Clin. Microbiol. Rev. 12:501–517.
- Giaever, G., A. M. Chu, L. Ni, C. Connelly, L. Riles, S. Veronneau, S. Dow, A. Lucau-Danila, K. Anderson, B. Andre, A. P. Arkin, A. Astromoff, M. El-Bakkoury, R. Bangham, R. Benito, S. Brachat, S. Campanaro, M. Curr tiss, K. Davis, A. Deutschbauer, K. D. Entian, P. Flaherty, F. Foury, D. J. Garfinkel, M. Gerstein, D. Gotte, U., Guldener, J. H. Hegemann, S. Hempel, Z. Herman, D. F. Jaramillo, D. E. Kelly, S. L. Kelly, P. Kotter, D. LaBonte, D. C. Lamb, N. Lan, H. Liang, H. Liao, L. Liu, C. Luo, M. Lussier, R. Mao, P. Menard, S. L. Ooi, J. L. Revuelta, C. J. Roberts, M. Rose, P. Ross-Macdonald, B. Scherens, G. Schimmack, B. Shafer, D. D. Shoemaker, S. Sookhai-Mahadeo, R. K. Storms, J. N. Strathern, G. Valle, M. Voet, G. Volckaert, C. Y. Wang, T. R. Ward, J. Wilhelmy, E. A. Winzeler, Y. Yang, G. Yen, E. Youngman, K. Yu, H. Bussey, J. D. Boeke, M. Snyder, P. Philippsen, R. W. Davis, and M. Johnston. 2002. Functional profiling of the Saccharomyces cerevisiae genome. Nature 418:387–391.
- 281. Gibbons, R. J., T. L. McDowell, S. Raman, D. M. O'Rourke, D. Garrick, H. Ayyub, and D. R. Higgs. 2000. Mutations in ATRX, encoding a SWI/SNF-like protein, cause diverse changes in the pattern of DNA methylation. Nat. Genet. 24:368–371.
- Gilbertson, L. A., and F. W. Stahl. 1996. A test of the double-strand break repair model for meiotic recombination in *Saccharomyces cerevisiae*. Genetics 144:27–41.
- Girbardt, M. 1969. Die Ultrastruktur der Apikalregion von Pilzhyphen. Protoplasma 67:413–441.
- 284. Glazer, L., and D. H. Brown. 1957. The synthesis of chitin in cell free extracts of *Neurospora crassa*. J. Biol. Chem. 228:729–742.
- Glickman, M. H., D. M. Rubin, V. A. Fried, and D. Finley. 1998. The regulatory particle of the *Saccharomyces cerevisiae* proteasome. Mol. Cell. Biol. 18:3149–3162.
- Goffeau, A., B. G. Barrell, H. Bussey, R. W. Davis, B. Dujon, H. Feldmann, F. Galibert, J. D. Hoheisel, C. Jacq, M. Johnston, E. J. Louis, H. W. Mewes,

Y. Murakami, P. Philippsen, H. Tettelin, and S. G. Oliver. 1996. Life with 6000 genes. Science 274:546, 563–567.

- 287. Goldman, G. H., S. L. McGuire, and S. D. Harris. 2002. The DNA damage response in filamentous fungi. Fungal Genet. Biol. 35:183–195.
- Goodrich-Tanrikulu, M., D. J. Jacobson, A. E. Stafford, J. T. Lin, and T. A. McKeon. 1999. Characterization of *Neurospora crassa* mutants isolated following repeat-induced point mutation of the beta subunit of fatty acid synthase. Curr. Genet. 36:147–152.
- Goodrich-Tanrikulu, M., A. E. Stafford, J. T. Lin, M. I. Makapugay, G. Fuller, and T. A. McKeon. 1994. Fatty acid biosynthesis in novel ufa mutants of *Neurospora crassa*. Microbiology 140:2683–2690.
- Gorbalenya, A. E., and E. V. Koonin. 1993. Helicases: aminoacid sequence comparisons and structure-function relationships. Curr. Opin. Struct. Biol. 3:419–429.
- 291. Gorl, M., M. Merrow, B. Huttner, J. Johnson, T. Roenneberg, and M. Brunner. 2001. A PEST-like element in FREQUENCY determines the length of the circadian period in *Neurospora crassa*. EMBO J. 20:7074–7084.
- 292. Gourgues, M., P. H. Clergeot, C. Veneault, J. Cots, S. Sibuet, A. Brunet-Simon, C. Levis, T. Langin, and M. H. Lebrun. 2002. A new class of tetraspanins in fungi. Biochem. Biophys. Res. Commun. 297:1197–1204.
- 293. Gow, L. A., and C. P. Selitrennikoff. 1984. Chitin synthetase of *Neurospora crassa:* inhibition by nikkomycin, polyoxin B, and UDP. Curr. Microbiol. 11:211–216.
- 294. Gow, N. A. R. 1994. Tip growth and polarity, p. 277–299. In N. A. R. Gow and G. M. Gadd (ed.), The growing fungus. Chapman & Hall, London, United Kingdom.
- 295. Grant, S. R. 1999. Dissecting the mechanisms of posttranscriptional gene silencing: divide and conquer. Cell 96:303–306.
- Gratzner, H., and D. N. Sheehan. 1969. Neurospora mutant exhibiting hyperproduction of amylase and invertase. J. Bacteriol. 97:544–549.
- 297. Graul, R. C., and W. Sadee. 1997. Evolutionary relationships among proteins probed by an iterative neighborhood cluster analysis (INCA). Alignment of bacteriorhodopsins with the yeast sequence YRO2. Pharm. Res. 14:1533–1541.
- Grayburn, W. S., and E. U. Selker. 1989. A natural case of RIP: degeneration of DNA sequence in an ancestral tandem duplication. Mol. Cell. Biol. 9:4416–4421.
- 299. Grewal, S. I., and S. C. Elgin. 2002. Heterochromatin: new possibilities for the inheritance of structure. Curr. Opin. Genet. Dev. 12:178–187.
- Griffen, D. H. 1994. Fungal physiology, 2nd ed. Wiley-Liss, Inc., New York, N.Y.
- 301. Griffiths, A. J. F., R. A. Collins, and F. E. Nargang. 1995. The mitochondrial genetics of *Neurospora*, p. 93–105. *In* U. Kück (ed.), The Mycota, vol. II. Genetics and biotechnology. Springer-Verlag, KG, Berlin, Germany.
- 302. Grishok, A., A. E. Pasquinelli, D. Conte, N. Li, S. Parrish, I. Ha, D. L. Baillie, A. Fire, G. Ruvkun, and C. C. Mello. 2001. Genes and mechanisms related to RNA interference regulate expression of the small temporal RNAs that control *C. elegans* developmental timing. Cell 106:23–34.
- 303. Groll, M., L. Ditzel, J. Lowe, D. Stock, M. Bochtler, H. D. Bartunik, and R. Huber. 1997. Structure of 20S proteasome from yeast at 2.4 A resolution. Nature 386:463–471.
- Gupta, G. D., and I. Brent Heath. 2002. Predicting the distribution, conservation, and functions of SNAREs and related proteins in fungi. Fungal Genet. Biol. 36:1–21.
- Gygi, S. P., Y. Rochon, B. R. Franza, and R. Aebersold. 1999. Correlation between protein and mRNA abundance in yeast. Mol. Cell. Biol. 19:1720– 1730.
- 306. Hagemann, A. T., and E. U. Selker. 1996. Control and function of DNA methylation in *Neurospora crassa*, p. 335–344. *In* V. E. A. Russo, R. A. Martienssen, and A. D. Riggs (ed.), Epigenetic mechanisms of gene regulation. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y.
- 307. Hall, I. M., G. D. Shankaranarayana, K. I. Noma, N. Ayoub, A. Cohen, and S. I. Grewal. 2002. Establishment and maintenance of a heterochromatin domain. Science 297:2232–2237.
- Hall, R. A., R. T. Premont, and R. J. Lefkowitz. 1999. Heptahelical receptor signaling: beyond the G protein paradigm. J. Cell Biol. 145:927–932.
- 309. Hammond, S. M., A. A. Caudy, and G. J. Hannon. 2001. Post-transcriptional gene silencing by double-stranded RNA. Nat. Rev. Genet. 2:110–119.
- Han, B.-K., R. Aramayo, and M. Polymenis. 2003. The G1 cyclin Cln3p controls vacuolar biogenesis in *Saccharomyces cerevisiae*. Genetics 165:467– 476.
- Han, Y., X. Liu, U. Benny, H. C. Kistler, and H. D. VanEtten. 2001. Genes determining pathogenicity to pea are clustered on a supernumerary chromosome in the fungal plant pathogen *Nectria haematococca*. Plant J. 25: 305–314.
- 312. Handa, N., Y. Noguchi, Y. Sakuraba, P. Ballario, G. Macino, N. Fujimoto, C. Ishii, and H. Inoue. 2000. Characterization of the *Neurospora crassa mus-25* mutant: the gene encodes a protein which is homologous to the *Saccharomyces cerevisiae* Rad54 protein. Mol. Gen. Genet. 264:154–163.
- 313. Hanson, M. A., and G. A. Marzluf. 1975. Control of the synthesis of a single

enzyme by multiple regulatory circuits in *Neurospora crassa*. Proc. Natl. Acad. Sci. USA 72:1240-1244.

- Harashima, T., and J. Heitman. 2002. The Galpha protein Gpa2 controls yeast differentiation by interacting with kelch repeat proteins that mimic Gbeta subunits. Mol. Cell 10:163–173.
- Harris, S. D. 2001. Septum formation in *Aspergillus nidulans*. Curr. Opin. Microbiol. 4:736–739.
- Hasegawa, T., H. Xiao, F. Hamajima, and K. Isobe. 2000. Interaction between DNA-damage protein GADD34 and a new member of the Hsp40 family of heat shock proteins that is induced by a DNA-damaging reagent. Biochem. J. 352:795–800.
- 317. Hass, H., K. Angermayr, and G. Stöffler. 1997. Molecular analysis of a *Penicillium chrysogenum* GATA factor encoding gene (*srcP*) exhibiting significant homology to the *Ustilago maydis urbs1* gene. Gene 184:33–37.
- 318. Hass, H., I. Zadra, G. Stöffler, and K. Angermayr. 1999. The Aspergillus nidulans GATA factor SREA is involved in regulation of siderophore biosynthesis and control of iron uptake. J. Biol. Chem. 274:4613–4619.
- Hatakeyama, S., Y. Ito, A. Shimane, C. Ishii, and H. Inoue. 1998. Cloning and characterization of the yeast RAD1 homolog gene (*mus-38*) from *Neurospora crassa*: evidence for involvement in nucleotide excision repair. Curr. Genet. 33:276–283.
- Hawkesford, M. J. 2003. Transporter gene families in plants: the sulphate transporter gene family—redundancy or specialization? Physiol. Plant. 117: 155–163.
- Hawksworth, D. L. 2001. The magnitude of fungal diversity: the 1.5 million species estimate revisited. Mycol. Res. 105:1422–1432.
- 322. Hays, S. M., J. Swanson, and E. U. Selker. 2002. Identification and characterization of the genes encoding the core histones and histone variants of *Neurospora crassa*. Genetics 160:961–973.
- 323. He, Q., P. Cheng, Y. Yang, L. Wang, K. H. Gardner, and Y. Liu. 2002. White collar-1, a DNA binding transcription factor and a light sensor. Science 297:840–843.
- Heath, B., A. Bonham, A. Akram, and G. Gupta. 2003. The interrelationships of actin and hyphal tip growth in the ascomycete *Geotrichum candidum*. Fungal Genet. Biol. 38:85–97.
- Hector, R. F. 1993. Compounds active against cell walls of medically important fungi. Clin. Microbiol. Rev. 6:1–21.
- Hegde, V., and H. Klein. 2000. Requirement for the SRS2 DNA helicase gene in non-homologous end joining in yeast. Nucleic Acids Res. 28:2779– 2783.
- 327. Heinemeyer, W., N. Trondle, G. Albrecht, and D. H. Wolf. 1994. PRE5 and PRE6, the last missing genes encoding 20S proteasome subunits from yeast? Indication for a set of 14 different subunits in the eukaryotic proteasome core. Biochemistry 33:12229–12237.
- Heintzen, C., J. J. Loros, and J. C. Dunlap. 2001. The PAS protein VIVID defines a clock-associated feedback loop that represses light input, modulates gating, and regulates clock resetting. Cell 104:453–464.
- Hellauer, K., E. Sirard, and B. Turcotte. 2001. Decreased expression of specific genes in yeast cells lacking histone H1. J. Biol. Chem. 276:13587– 13592.
- 330. Henikoff, S., K. Ahmad, and H. S. Malik. 2001. The centromere paradox: stable inheritance with rapidly evolving DNA. Science 293:1098–1102.
- Henikoff, S., K. Ahmad, J. S. Platero, and B. van Steensel. 2000. Heterochromatic deposition of centromeric histone H3-like proteins. Proc. Natl. Acad. Sci. USA 97:716–721.
- Henrissat, B., and A. Bairoch. 1996. Updating the sequence-based classification of glycosyl hydrolases. Biochem. J. 316:695–696.
- 333. Hepworth, S. R., H. Friesen, and J. Segall. 1998. NDT80 and the meiotic recombination checkpoint regulate expression of middle sporulation-specific genes in *Saccharomyces cerevisiae*. Mol. Cell. Biol. 18:5750–5761.
- 334. Herrmann, G., T. Lindahl, and P. Schar. 1998. Saccharomyces cerevisiae LIF1: a function involved in DNA double-strand break repair related to mammalian XRCC4. EMBO J. 17:4188–4198.
- 335. Hettema, E. H., C. C. Ruigrok, M. G. Koerkamp, M. van den Berg, H. F. Tabak, B. Distel, and I. Braakman. 1998. The cytosolic DnaJ-like protein djp1p is involved specifically in peroxisomal protein import. J. Cell Biol. 142:421–434.
- 336. Hickey, P. C., D. J. Jacobson, N. D. Read, and N. L. Glass. 2002. Live-cell imaging of vegetative hyphal fusion in *Neurospora crassa*. Fungal Genet. Biol. 37:109–119.
- 337. Higuchi, S., J. Tamura, P. R. Giri, J. W. Polli, and R. L. Kincaid. 1991. Calmodulin-dependent protein phosphatase from *Neurospora crassa*. Molecular cloning and expression of recombinant catalytic subunit. J. Biol. Chem. 266:18104–18112.
- Hildebrandt, E. R., and M. A. Hoyt. 2000. Mitotic motors in Saccharomyces cerevisiae. Biochim. Biophys. Acta 1496:99–116.
- 339. Hinnebusch, A. G., G. Lucchini, and G. R. Fink. 1985. A synthetic HIS4 regulatory element confers general amino acid control on cytochrome c gene (CYC1) of yeast. Proc. Natl. Acad. Sci. USA 82:498–502.
- 340. Hinnebusch, A. G., and K. Natarajan. 2002. Gcn4p, a master regulator of gene expression, is controlled at multiple levels by diverse signals of starvation and stress. Eukaryot. Cell 1:22–32.

- Hirokawa, N., Y. Noda, and Y. Okada. 1998. Kinesin and dynein superfamily proteins in organelle transport and cell division. Curr. Opin. Cell Biol. 10:60–73.
- 342. Hodge, T., and M. J. Cope. 2000. A myosin family tree. J. Cell Sci. 113: 3353–3354.
- 343. Hoffmann, B., W. Zuo, A. Liu, and N. R. Morris. 2001. The LIS1-related protein NUDF of *Aspergillus nidulans* and its interaction partner NUDE bind directly to specific subunits of dynein and dynactin and to alpha- and gamma-tubulin. J. Biol. Chem. 276;38877–38884.
- Hohfeld, J., and S. Jentsch. 1997. GrpE-like regulation of the hsc70 chaperone by the anti-apoptotic protein BAG-1. EMBO J. 16:6209–6216.
- 345. Holleran, E. A., S. Karki, and E. L. Holzbaur. 1998. The role of the dynactin complex in intracellular motility. Int. Rev. Cytol. 182:69–109.
- Hollingsworth, N. M., L. Goetsch, and B. Byers. 1990. The HOP1 gene encodes a meiosis-specific component of yeast chromosomes. Cell 61:73– 84.
- Holzbaur, E. L. F., and R. B. Vallee. 1994. Dyneins—molecular structure and cellular function. Annu. Rev. Cell Biol. 10:339–372.
- Hopwood, D. A., and D. H. Sherman. 1990. Molecular genetics of polyketides and its comparison to fatty acid biosynthesis. Annu. Rev. Genet. 24:37–66.
- 349. Horiuchi, H., M. Fujiwara, S. Yamashita, A. Ohta, and M. Takagi. 1999. Proliferation of intrahyphal hyphae caused by disruption of *csmA*, which encodes a class V chitin synthase with a myosin motor-like domain in *Aspergillus nidulans*. J. Bacteriol. 181:3721–3729.
- 350. Horton, L. E., P. James, E. A. Craig, and J. O. Hensold. 2001. The yeast hsp70 homologue Ssa is required for translation and interacts with Sis1 and Pab1 on translating ribosomes. J. Biol. Chem. 276:14426–14433.
- 351. Hsu, J. Y., Z. W. Šun, X. Li, M. Reuben, K. Tatchell, D. K. Bishop, J. M. Grushcow, C. J. Brame, J. A. Caldwell, D. F. Hunt, R. Lin, M. M. Smith, and C. D. Allis. 2000. Mitotic phosphorylation of histone H3 is governed by Ip11/aurora kinase and Glc7/PP1 phosphatase in budding yeast and nema-todes. Cell 102:279–291.
- 352. Hu, Y., and N. F. Mivechi. 2003. HSF-1 interacts with Ral-binding protein 1 in a stress-responsive, multiprotein complex with HSP90 in vivo. J. Biol. Chem. 278:17299–17306.
- 353. Huang, Y. 2002. Transcriptional silencing in Saccharomyces cerevisiae and Schizosaccharomyces pombe. Nucleic Acids Res. 30:1465–1482.
- Huelsenbeck, J. P., and F. Ronquist. 2001. MRBAYES: Bayesian inference of phylogenetic trees. Bioinformatics 17:754–755.
- 355. Huelsenbeck, J. P., F. Ronquist, R. Nielsen, and J. P. Bollback. 2001. Bayesian inference of phylogeny and its impact on evolutionary biology. Science 294:2310–2314.
- 356. Hundley, H., H. Eisenman, W. Walter, T. Evans, Y. Hotokezaka, M. Wiedmann, and E. Craig. 2002. The in vivo function of the ribosome-associated Hsp70, Ssz1, does not require its putative peptide-binding domain. Proc. Natl. Acad. Sci. USA 99:4203–4208.
- 357. Hurtado, C. A., J. M. Beckerich, C. Gaillardin, and R. A. Rachubinski. 2000. A rac homolog is required for induction of hyphal growth in the dimorphic yeast *Yarrowia lipolytica*. J. Bacteriol. **182**:2376–2386.
- 358. Hutchinson, E. G., W. Tichelaar, G. Hofhaus, H. Weiss, and K. R. Leonard. 1989. Identification and electron microscopic analysis of a chaperonin oligomer from *Neurospora crassa* mitochondria. EMBO J. 8:1485–1490.
- Hutvagner, G., and P. D. Zamore. 2002. RNAi: nature abhors a doublestrand. Curr. Opin. Genet. Dev. 12:225–232.
- 360. Hwang, C. S., M. A. Flaishman, and P. E. Kolattukudy. 1995. Cloning of a gene expressed during appressorium formation by *Colletotrichum gloeosporioides* and a marked decrease in virulence by disruption of this gene. Plant Cell 7:183–193.
- Iizuka, M., and M. M. Smith. 2003. Functional consequences of histone modifications. Curr. Opin. Genet. Dev. 13:154–160.
- 362. Ikemura, T. 1981. Correlation between the abundance of *Escherichia coli* transfer RNAs and the occurrence of the respective codons in its protein genes: a proposal for a synonymous codon choice that is optimal for the *E. coli* translational system. J. Mol. Biol. 151:389–409.
- 363. Ingolia, T. D., and E. A. Craig. 1982. Four small *Drosophila* heat shock proteins are related to each other and to mammalian alpha-crystallin. Proc. Natl. Acad. Sci. USA 79:2360–2364.
- Inoue, H. 1999. DNA repair and specific-locus mutagenesis in *Neurospora crassa*. Mutat. Res. 437:121–133.
- Irelan, J. T., and E. U. Selker. 1997. Cytosine methylation associated with repeat-induced point mutation causes epigenetic gene silencing in *Neuro*spora crassa. Genetics 146:509–523.
- 366. Isnard, A. D., D. Thomas, and Y. Surdin-Kerjan. 1996. The study of methionine uptake in *Saccharomyces cerevisiae* reveals a new family of amino acid permeases. J. Mol. Biol. 262:473–484.
- 367. Ito, S., Y. Matsui, A. Toh-e, T. Harashima, and H. Inoue. 1997. Isolation and characterization of the *krev-1* gene, a novel member of ras superfamily in *Neurospora crassa*: involvement in sexual cycle progression. Mol. Gen. Genet. 255:429–437.
- 368. Ito, T., M. Bulger, R. Kobayashi, and J. T. Kadonaga. 1996. Drosophila NAP-1 is a core histone chaperone that functions in ATP-facilitated as-

sembly of regularly spaced nucleosomal arrays. Mol. Cell. Biol. 16:3112-3124.

- 369. Ivey, F. D., P. N. Hodge, G. E. Turner, and K. A. Borkovich. 1996. The Gα_i homologue *gna-1* controls multiple differentiation pathways in *Neurospora crassa*. Mol. Biol. Cell 7:1283–1297.
- 370. Ivey, F. D., A. M. Kays, and K. A. Borkovich. 2002. Shared and independent roles for a Gα_i protein and adenylyl cyclase in regulating development and stress responses in *Neurospora crassa*. Eukaryot. Cell 1:634–642.
- 371. Ivey, F. D., Q. Yang, and K. A. Borkovich. 1999. Positive regulation of adenylyl cyclase activity by a Gαi homologue in *Neurospora crassa*. Fungal Genet. Biol. 26:48–61.
- 372. Jabri, E., D. Taft, D. Quigley, M. Hrmova, P. Phelps, M. Alders, and C. P. Selitrennikoff. 1989. (1–3)-β-glucan synthesis of *Neurospora*. Curr. Microbiol. 19:153–161.
- 373. Jackl, G., and W. Sebald. 1975. Identification of two products of mitochondrial protein synthesis associated with mitochondrial adenosine triphosphatase from *Neurospora crassa*. Eur. J. Biochem. 54:97–106.
- 374. Jackson, J. D., and M. A. Gorovsky. 2000. Histone H2A.Z has a conserved function that is distinct from that of the major H2A sequence variants. Nucleic Acids Res. 28:3811–3816.
- Jackson, J. P., A. M. Lindroth, X. Cao, and S. E. Jacobsen. 2002. Control of CpNpG DNA methylation by the KRYPTONITE histone H3 methyltransferase. Nature 416:556–560.
- Jacobsen, S. E., M. P. Running, and E. M. Meyerowitz. 1999. Disruption of an RNA helicase/RNAse III gene in *Arabidopsis* causes unregulated cell division in floral meristems. Development 126:5231–5243.
- 377. Jae Yoo, E., Y. Kyu Jang, M. Ae Lee, P. Bjerling, J. Bum Kim, K. Ekwall, R. Hyun Seong, and S. Dai Park. 2002. Hrp3, a chromodomain helicase/ ATPase DNA binding protein, is required for heterochromatin silencing in fission yeast. Biochem. Biophys. Res. Commun. 295:970–974.
- 378. Jaffe, D. B., J. Butler, S. Gnerre, E. Mauceli, K. Lindblad-Toh, J. P. Mesirov, M. C. Zody, and E. S. Lander. 2003. Whole-genome sequence assembly for mammalian genomes: Arachne 2. Genome Res. 13:91–96.
- 379. Janke, C., J. Ortiz, J. Lechner, A. Shevchenko, M. M. Magiera, C. Schramm, and E. Schiebel. 2001. The budding yeast proteins Spc24p and Spc25p interact with Ndc80p and Nuf2p at the kinetochore and are important for kinetochore clustering and checkpoint control. EMBO J. 20:777–791.
- Janke, C., J. Ortiz, T. U. Tanaka, J. Lechner, and E. Schiebel. 2002. Four new subunits of the Dam1-Duo1 complex reveal novel functions in sister kinetochore biorientation. EMBO J. 21:181–193.
- 381. Jason, L. J., S. C. Moore, J. D. Lewis, G. Lindsey, and J. Ausio. 2002. Histone ubiquitination: a tagging tail unfolds? Bioessays 24:166–174.
- Jeddeloh, J. A., T. L. Stokes, and E. J. Richards. 1999. Maintenance of genomic methylation requires a SWI2/SNF2-like protein. Nat. Genet. 22: 94–97.
- 383. Jeenes, D. J., D. A. Mackenzie, I. N. Roberts, and D. B. Archer. 1991. Heterologous protein production by filamentous fungi. Biotechnol. Genet. Eng. Rev. 9:327–367.
- Jenuwein, T., and C. D. Allis. 2001. Translating the histone code. Science 293:1074–1080.
- Johnson, C. H. 2001. Endogenous timekeepers in photosynthetic organisms. Annu. Rev. Physiol. 63:695–728.
- Johnson, L., X. Cao, and S. Jacobsen. 2002. Interplay between two epigenetic marks. DNA methylation and histone H3 lysine 9 methylation. Curr. Biol. 12:1360–1367.
- 387. Jones, P. A., and P. W. Laird. 1999. Cancer epigenetics comes of age. Nat. Genet. 21:163–167.
- Josefsson, L. G., and L. Rask. 1997. Cloning of a putative G-proteincoupled receptor from *Arabidopsis thaliana*. Eur. J. Biochem. 249:415–420.
- 389. Jumpponen, A., J. M. Trappe, and E. Cazares. 2002. Occurrence of ectomycorrhizal fungi on the forefront of retreating Lyman Glacier (Washington, USA) in relation to time since deglaciation. Mycorrhiza 12:43–49.
- Kabani, M., J. M. Beckerich, and J. L. Brodsky. 2002. Nucleotide exchange factor for the yeast Hsp70 molecular chaperone Ssa1p. Mol. Cell. Biol. 22: 4677–4689.
- 391. Kabani, M., J. M. Beckerich, and C. Gaillardin. 2000. Sls1p stimulates Sec63p-mediated activation of Kar2p in a conformation-dependent manner in the yeast endoplasmic reticulum. Mol. Cell. Biol. 20:6923–6934.
- 392. Kadam, S., and B. M. Emerson. 2003. Transcriptional specificity of human SWI/SNF BRG1 and BRM chromatin remodeling complexes. Mol. Cell 11:377–389.
- 393. Kafer, E., and M. Fraser. 1988. Isolation and genetic analysis of nuclease halo (nuh) mutants in *Neurospora*. Mol. Gen. Genet. 169:117–127.
- 394. Kafer, E., and G. R. Witchell. 1984. Effects of *Neurospora nuclease* halo (nuh) mutants on secretion of two phosphate-repressible alkaline deoxyribonucleases. Biochem. Genet. 22:403–417.
- 395. Kaliraman, V., J. R. Mullen, W. M. Fricke, S. A. Bastin-Shanower, and S. J. Brill. 2001. Functional overlap between Sgs1-Top3 and the Mms4-Mus81 endonuclease. Genes Dev. 15:2730–2740.
- 396. Kanaan, M. N., and G. A. Marzluf. 1991. Mutational analysis of the DNA-

binding domain of the CYS3 regulatory protein of *Neurospora crassa*. Mol. Cell. Biol. **11**:4356–4362.

- 397. Kana-uchi, A., C. T. Yamashiro, S. Tanabe, and T. Murayama. 1997. A ras homologue of *Neurospora crassa* regulates morphology. Mol. Gen. Genet. 254:427–432.
- 398. Kang, M. S., and E. Cabib. 1986. Regulation of fungal cell wall growth: a guanine nucleotide-binding, proteinaceous component required for activity of (1→3)-beta-D-glucan synthase. Proc. Natl. Acad. Sci. USA 83:5808–5812.
- 399. Kang, S., and R. L. Metzenberg. 1990. Molecular analysis of *nuc-1⁺*, a gene controlling phosphorus acquisition in *Neurospora crassa*. Mol. Cell. Biol. 10:5839–5848.
- 400. Kapoor, M., C. A. Curle, and C. Runham. 1995. The *hsp70* gene family of *Neurospora crassa*: cloning, sequence analysis, expression, and genetic mapping of the major stress-inducible member. J. Bacteriol. 177:212–221.
- 401. Karcher, R. L., S. W. Deacon, and V. I. Gelfand. 2002. Motor-cargo interactions: the key to transport specificity. Trends Cell Biol. 12:21–27.
- 402. Kasahara, S., P. Wang, and D. L. Nuss. 2000. Identification of *bdm-1*, a gene involved in G protein beta-subunit function and alpha-subunit accumulation. Proc. Natl. Acad. Sci. USA 97:412–417.
- 403. Kasinsky, H. E., J. D. Lewis, J. B. Dacks, and J. Ausio. 2001. Origin of H1 linker histones. FASEB J. 15:34–42.
- 404. Katz, M. E., A. Masoumi, S. R. Burrows, C. G. Shirtliff, and B. F. Cheetham. 2000. The *Aspergillus nidulans xprF* gene encodes a hexokinaselike protein involved in the regulation of extracellular proteases. Genetics 156:1559–1571.
- 405. Kays, A. M., and K. A. Borkovich. Severe impairment of growth and differentiation in a *Neurospora crassa* mutant lacking all heterotrimeric $G\alpha$ proteins. Genetics, in press.
- 406. Kays, A. M., P. S. Rowley, R. A. Baasiri, and K. A. Borkovich. 2000. Regulation of conidiation and adenylyl cyclase levels by the Galpha protein GNA-3 in *Neurospora crassa*. Mol. Cell. Biol. 20:7693–7705.
- 407. Keeney, S. 2001. Mechanism and control of meiotic recombination initiation. Curr. Top. Dev. Biol. 52:1–53.
- Keeney, S., C. N. Giroux, and N. Kleckner. 1997. Meiosis-specific DNA double-strand breaks are catalyzed by Spo11, a member of a widely conserved protein family. Cell 88:375–384.
- 409. Kelkar, H. S., J. Griffith, M. E. Case, S. F. Covert, R. D. Hall, C. H. Keith, J. S. Oliver, M. J. Orbach, M. S. Sachs, J. R. Wagner, M. J. Weise, J. K. Wunderlich, and J. Arnold. 2001. The Neurospora crassa genome: cosmid libraries sorted by chromosome. Genetics 157:979–990.
- 410. Kennell, J. C., R. A. Collins, A. J. F. Griffiths, and F. E. Nargang. Mitochondrial genetics of *Neurospora*, p. 95–112. *In* U. Kück (ed.), The Mycota, vol. II. Genetics and biotechnology, 2nd ed., in press. Springer-Verlag, KG, Berlin, Germany.
- Kertesz, M. A. 2000. Riding the sulfur cycle-metabolism of sulfonates and sulfate esters in gram-negative bacteria. FEMS Microbiol. Rev. 24:135–175.
- 412. Ketter, J. S., and G. A. Marzluf. 1988. Molecular cloning and analysis of the regulation of cys-14⁺, a structural gene of the sulfur regulatory circuit of *Neurospora crassa*. Mol. Cell. Biol. 8:1504–1508.
- 413. Kim, A. J., and S. A. Endow. 2000. A kinesin family tree. J. Cell Sci. 113:3681–3682.
- 414. Kim, H., R. L. Metzenberg, and M. A. Nelson. 2002. Multiple functions of *mfa-1*, a putative pheromone precursor gene of *Neurospora crassa*. Eukaryot. Cell 1:987–999.
- 415. Kimura, Y., M. Takaoka, S. Tanaka, H. Sassa, K. Tanaka, B. Polevoda, F. Sherman, and H. Hirano. 2000. N(alpha)-acetylation and proteolytic activity of the yeast 20's proteasome. J. Biol. Chem. 275:4635–4639.
- 416. King, S. M., E. Barbarese, J. F. Dillman, S. E. Benashski, K. T. Do, R. S. Patelking, and K. K. Pfister. 1998. Cytoplasmic dynein contains a family of differentially expressed light chains. Biochemistry 37:15033–15041.
- 417. Kinsey, J. A. 1989. Restricted distribution of the Tad transposon in strains of *Neurospora*. Curr. Genet. 15:271–275.
- 418. Kinsey, J. A., P. W. Garrett-Engele, E. B. Cambareri, and E. U. Selker. 1994. The *Neurospora* transposon Tad is sensitive to repeat-induced point mutation (RIP). Genetics 138:657–664.
- Kinsey, J. A., and J. Helber. 1989. Isolation of a transposable element from Neurospora crassa. Proc. Natl. Acad. Sci. USA 86:1929–1933.
- Kirchhausen, T. 2000. Three ways to make a vesicle. Nat. Rev. Mol. Cell Biol 1:187–198.
- 421. Kirchner, J., S. Seiler, S. Fuchs, and M. Schliwa. 1999. Functional anatomy of the kinesin molecule in vivo. EMBO J. 18:4404–4413.
- 422. Kirchner, J., G. Woehlke, and M. Schliwa. 1999. Universal and unique features of kinesin motors: insights from a comparison of fungal and animal conventional kinesins. Biol. Chem. 380:915–921.
- 423. Klein, P. S., T. J. Sun, C. L. Saxe III, A. R. Kimmel, R. L. Johnson, and P. N. Devreotes. 1988. A chemoattractant receptor controls development in *Dictyostelium discoideum*. Science 241:1467–1472.
- 424. Klis, F. M., P. Mol, K. Hellingwerf, and S. Brul. 2002. Dynamics of cell wall structure in *Saccharomyces cerevisiae*. FEMS Microbiol. Rev. 26:239–256.
- 425. Klug, R. M., and C. Benning. 2001. Two enzymes of diacylglyceryl-O-4'-(N,N,N,-trimethyl)homoserine biosynthesis are encoded by btaA and btaB

in the purple bacterium *Rhodobacter sphaeroides*. Proc. Natl. Acad. Sci. USA **98:**5910–5915.

- 426. Kodama, Y., F. Omura, K. Takahashi, K. Shirahige, and T. Ashikari. 2002. Genome-wide expression analysis of genes affected by amino acid sensor Ssy1p in Saccharomyces cerevisiae. Curr. Genet. 41:63–72.
- 427. Kore-eda, S., T. Murayama, and I. Uno. 1991. Isolation and characterization of the adenylate cyclase structural gene of *Neurospora crassa*. Jpn. J. Genet. 66:317–334.
- Kornberg, H. L. 1966. Anapleurotic sequences and their role in metabolism. Essays Biochem. 2:1–31.
- 429. Kosugi, A., Y. Koizumi, F. Yanagida, and S. Udaka. 2001. MUP1, high affinity methionine permease, is involved in cysteine uptake by *Saccharomyces cerevisiae*. Biosci. Biotechnol. Biochem. 65:728–731.
- Kothe, G. O., and S. J. Free. 1998. Calcineurin subunit B is required for normal vegetative growth in *Neurospora crassa*. Fungal Genet. Biol. 23:248– 258.
- 431. Kothe, G. O., and S. J. Free. 1998. The isolation and characterization of nrc-1 and nrc-2, two genes encoding protein kinases that control growth and development in *Neurospora crassa*. Genetics 149:117–130.
- Kouzarides, T. 2002. Histone methylation in transcriptional control. Curr. Opin. Genet. Dev. 12:198–209.
- 433. Kouzminova, E. A., and E. U. Selker. 2001. Dim-2 encodes a DNA-methyltransferase responsible for all known cytosine methylation in *Neurospora*. EMBO J. 20:4309–4323.
- 434. Kreis, T., and R. Vale. 1999. Guidebook to the cytoskeletal and motor proteins, 2nd ed. Oxford University Press, Oxford, United Kingdom.
- 435. Krems, B., C. Charizanis, and K. D. Entian. 1996. The response regulatorlike protein Pos9/Skn7 of Saccharomyces cerevisiae is involved in oxidative stress resistance. Curr. Genet. 29:327–334.
- Kronstad, J. W., and C. Staben. 1997. Mating type in filamentous fungi. Annu. Rev. Genet. 31:245–276.
- 437. Kruft, V., H. Eubel, L. Jansch, W. Werhahn, and H. P. Braun. 2001. Proteomic approach to identify novel mitochondrial proteins in *Arabidopsis*. Plant Physiol. 127:1694–1710.
- 438. Kuffner, R., A. Rohr, A. Schmiede, C. Krull, and U. Schulte. 1998. Involvement of two novel chaperones in the assembly of mitochondrial NADH: ubiquinone oxidoreductase (complex 1). J. Mol. Biol. 283:409–417.
- 439. Kumar, A., S. Agarwal, J. A. Heyman, S. Matson, M. Heidtman, S. Piccirillo, L. Umansky, A. Drawid, R. Jansen, Y. Liu, K. H. Cheung, P. Miller, M. Gerstein, G. S. Roeder, and M. Snyder. 2002. Subcellular localization of the yeast proteome. Genes Dev. 16:707–719.
- 440. Kumar, A., and J. V. Paietta. 1995. The sulfur controller-2 negative regulatory gene of *Neurospora crassa* encodes a protein with beta-transducin repeats. Proc. Natl. Acad. Sci. USA 92:3343–3347.
- 441. Kundu, M., J. Basu, M. Guchhait, and P. Chakrabarti. 1987. Isolation and characterization of an extracellular lipase from the conidia of *Neurospora crassa*. J. Gen. Microbiol. 133:149–153.
- 442. Kupfer, D. M., C. A. Reece, S. W. Clifton, B. A. Roe, and R. A. Prade. 1997. Multicellular ascomycetous fungal genomes contain more than 8000 genes. Fungal Genet. Biol. 21:364–372.
- 443. Kwon, Y. H., and M. H. Stipanuk. 2001. Cysteine regulates expression of cysteine dioxygenase and gamma-glutamylcysteine synthetase in cultured rat hepatocytes. Am. J. Physiol. Endocrinol. Metab. 280:E804–E815.
- 444. Kyriakis, J. M. 1999. Signaling by the germinal center kinase family of protein kinases. J. Biol. Chem. 274:5259–5262.
- 445. Lachner, M., R. J. O'Sullivan, and T. Jenuwein. 2003. An epigenetic road map for histone lysine methylation. J. Cell Sci. 116:2117–2124.
- Lagunas, R., and J. M. Gancedo. 1973. Reduced pyridine-nucleotides balance in glucose-growing Saccharomyces cerevisiae. Eur. J. Biochem. 37:90– 94.
- 447. Lakin-Thomas, P. L. 1993. Effects of inositol starvation on the levels of inositol phosphates and inositol lipids in *Neurospora crassa*. Biochem. J. 292:805–811.
- 448. Lambowitz, A. M., and M. Belfort. 1993. Introns as mobile genetic elements. Annu. Rev. Biochem. 62:587–622.
- 449. Lambowitz, A. M., J. R. Sabourin, H. Bertrand, R. Nickels, and L. Mc-Intosh. 1989. Immunological identification of the alternative oxidase of *Neurospora crassa* mitochondria. Mol. Cell Biol. 9:1362–1364.
- Lambowitz, A. M., and C. W. Slayman. 1971. Cyanide-resistant respiration in *Neurospora crassa*. J. Bacteriol. 108:1087–1096.
- Lambowitz, A. M., E. W. Smith, and C. W. Slayman. 1972. Electron transport in *Neurospora* mitochondria. Studies on wild type and poky. J. Biol. Chem. 247:4850–4858.
- Langst, G., and P. B. Becker. 2001. Nucleosome mobilization and positioning by ISWI-containing chromatin-remodeling factors. J. Cell Sci. 114: 2561–2696.
- Latge, J. P. 1999. Aspergillus fumigatus and aspergillosis. Clin. Microbiol. Rev. 12:310–350.
- 454. Lau, G., and J. E. Hamer. 1996. Regulatory genes controlling MPG1 expression and pathogenicity in the rice blast fungus *Magnaporthe grisea*. Plant Cell 8:771–781.
- 455. Lauter, F. R., V. E. Russo, and C. Yanofsky. 1992. Developmental and light

regulation of *eas*, the structural gene for the rodlet protein of *Neurospora*. Genes Dev. **6**:2373–2381.

- 456. Le Chevanton, L., and D. Zickler. 1991. Sordaria macrospora: the transition to the age of gene manipulation, p. 291–303. In J. W. Bennet and L. L. Lasure (ed.), More gene manipulations in fungi. Academic Press, Inc., San Diego, Calif.
- 457. Lechner, J., and J. Ortiz. 1996. The Saccharomyces cerevisiae kinetochore. FEBS Lett. 389:70–74.
- 458. Lee, D. W., R. J. Pratt, M. McLaughlin, and R. Aramayo. 2003. An Argonaute-like protein is required for meiotic silencing. Genetics 164:821–828.
- 459. Lee, D. W., K.-Y. Seong, R. Pratt, K. Baker, and R. Aramayo. Properties of unpaired DNA required for efficient silencing in *Neurospora crassa*. Genetics, in press.
- 460. Lee, H., Y. H. Fu, and G. A. Marzluf. 1990. Nucleotide sequence and DNA recognition elements of *alc*, the structural gene which encodes allantoicase, a purine catabolic enzyme of *Neurospora crassa*. Biochemistry 29:8779–8787.
- 461. Lee, K., J. J. Loros, and J. C. Dunlap. 2000. Interconnected feedback loops in the *Neurospora* circadian system. Science 289:107–110.
- 462. Lee, K. S., K. Irie, Y. Gotoh, Y. Watanabe, H. Araki, E. Nishida, K. Matsumoto, and D. E. Levin. 1993. A yeast mitogen-activated protein kinase homolog (Mpk1p) mediates signalling by protein kinase C. Mol. Cell. Biol. 13:3067–3075.
- 463. Lee-Yoon, D., D. Easton, M. Murawski, R. Burd, and J. R. Subjeck. 1995. Identification of a major subfamily of large hsp70-like proteins through the cloning of the mammalian 110-kDa heat shock protein. J. Biol. Chem. 270:15725–15733.
- 464. Lehmler, C., G. Steinberg, K. M. Snetselaar, M. Schliwa, R. Kahmann, and M. Bolker. 1997. Identification of a motor protein required for filamentous growth in *Ustilago maydis*. EMBO J. 16:3464–3473.
- 465. Lehnertz, B., Y. Ueda, A. A. Derijck, U. Braunschweig, L. Perez-Burgos, S. Kubicek, T. Chen, E. Li, T. Jenuwein, and A. H. Peters. 2003. Suv39hmediated histone H3 lysine 9 methylation directs DNA methylation to major satellite repeats at pericentric heterochromatin. Curr. Biol. 13:1192– 1200.
- 466. Leloir, L., and C. Cardini. 1953. The biosynthesis of glucosamine. Biochim. Biophys. Acta 12:15–22.
- 467. Lemmon, M. A., K. M. Ferguson, and J. Schlessinger. 1996. PH domains: diverse sequences with a common fold recruit signaling molecules to the cell surface. Cell 85:621–624.
- 468. Leonhardt, S. A., K. Fearson, P. N. Danese, and T. L. Mason. 1993. HSP78 encodes a yeast mitochondrial heat shock protein in the Clp family of ATP-dependent proteases. Mol. Cell. Biol. 13:6304–6313.
- 469. Leu, J. Y., P. R. Chua, and G. S. Roeder. 1998. The meiosis-specific Hop2 protein of *S. cerevisiae* ensures synapsis between homologous chromosomes. Cell 94:375–386.
- Levine, M., and R. Tijan. 2003. Transcription regulation and animal diversity. Nature 424:147–151.
- 471. Li, M., A. Makkinje, and Z. Damuni. 1996. The myeloid leukemia-associated protein SET is a potent inhibitor of protein phosphatase 2A. J. Biol. Chem. 271:11059–11062.
- 472. Li, Q., R. G. Ritzel, L. L. McLean, L. McIntosh, T. Ko, H. Bertrand, and F. E. Nargang. 1996. Cloning and analysis of the alternative oxidase gene of *Neurospora crassa*. Genetics 142:129–140.
- 473. Li, S., A. Ault, C. L. Malone, D. Raitt, S. Dean, L. H. Johnston, R. J. Deschenes, and J. S. Fassler. 1998. The yeast histidine protein kinase, Sln1p, mediates phosphotransfer to two response regulators, Ssk1p and Skn7p. EMBO J. 17:6952–6962.
- 474. Lin, S. S., J. K. Manchester, and J. I. Gordon. 2003. Sip2, an N-myristoylated beta subunit of Snf1 kinase, regulates aging in *Saccharomyces cerevisiae* by affecting cellular histone kinase activity, recombination at rDNA loci, and silencing. J. Biol. Chem. 278:13390–13397.
- Lindberg, R. A., W. G. Rhodes, L. D. Eirich, and H. Drucker. 1982. Extracellular acid proteases from *Neurospora crassa*. J. Bacteriol. 150:1103–1108.
 Lindegren, C. C. 1936. A six-point map of the sex chromosome of *Neuro-*
- Jandegrein, C. G. 1950. A sharpoint half of the sex enrolling some of Neurospora crassa. J. Genet. 32:243–256.
 Linden, H., P. Ballario, G. Arpaia, and G. Macino. 1999. Seeing the light:
- news in *Neurospora* blue light signal transduction. Adv. Genet. **41**:35–54. **478. Linden, H., and G. Macino.** 1997. White collar 2, a partner in blue-light
- 478. Linder, H., and G. Machio. 1997. White cona 2, a partier in oute-ngnt signal transduction, controlling expression of light-regulated genes in *Neurospora crassa*. EMBO J. 16:98–109.
- 479. Lipke, P. N., and R. Ovalle. 1998. Cell wall architecture in yeast: new structure and new challenges. J. Bacteriol. 180:3735–3740.
- 480. Liu, X. D., K. A. Morano, and D. J. Thiele. 1999. The yeast Hsp110 family member, Sse1, is an Hsp90 cochaperone. J. Biol. Chem. 274:26654–26660.
- Lloyd, A. T., and P. M. Sharp. 1993. Synonymous codon usage in *Kluyvero-myces lactis*. Yeast 9:1219–1228.
- 482. Lo, W. S., L. Duggan, N. C. Tolga, Emre, R. Belotserkovskya, W. S. Lane, R. Shiekhattar, and S. L. Berger. 2001. Snf1—a histone kinase that works in concert with the histone acetyltransferase Gen5 to regulate transcription. Science 293:1142–1146.
- 483. Lo, W. S., R. C. Trievel, J. R. Rojas, L. Duggan, J. Y. Hsu, C. D. Allis, R.

Marmorstein, and S. L. Berger. 2000. Phosphorylation of serine 10 in histone H3 is functionally linked in vitro and in vivo to Gen5-mediated acetylation at lysine 14. Mol. Cell 5:917–926.

- Loros, J. J., and J. C. Dunlap. 2001. Genetic and molecular analysis of circadian rhythms in *Neurospora*. Annu. Rev. Physiol. 63:757–794.
- 485. Losel, D. M. 1988. Fungal lipids, p. 699–806. In C. Ratledge and S. G. Wilkinson (ed.), Microbial lipids, vol. 1. Academic Press, Ltd., London, United Kingdom.
- 486. Lotz, G. P., H. Lin, A. Harst, and W. M. Obermann. 2003. Ahal binds to the middle domain of Hsp90, contributes to client protein activation, and stimulates the ATPase activity of the molecular chaperone. J. Biol. Chem. 278:17228–17235.
- 487. Lowe, T. M., and S. R. Eddy. 1997. tRNAscan-SE: a program for improved detection of transfer RNA genes in genomic sequence. Nucleic Acids Res. 25:955–964.
- 488. Lubben, T. H., A. A. Gatenby, G. K. Donaldson, G. H. Lorimer, and P. V. Viitanen. 1990. Identification of a groES-like chaperonin in mitochondria that facilitates protein folding. Proc. Natl. Acad. Sci. USA 87:7683–7687.
- Lui, H., C. A. Styles, and G. R. Fink. 1993. Elements of the yeast pheromone response pathway required for filamentous growth of diploids. Science 262:1741–1744.
- Lupas, A., M. Van Dyke, and J. Stock. 1991. Predicting coiled coils from protein sequences. Science 252:1162–1164.
- 491. Ma, P., S. Wera, P. Van Dijck, and J. M. Thevelein. 1999. The PDE1encoded low-affinity phosphodiesterase in the yeast *Saccharomyces cerevisiae* has a specific function in controlling agonist-induced cAMP signaling. Mol. Biol. Cell 10:91–104.
- Macielag, M. J., and R. Goldschmidt. 2000. Inhibitors of bacterial twocomponent signalling systems. Expert Opin. Investig. Drugs 9:2351–2369.
- 493. Madden, K., and M. Snyder. 1998. Cell polarity and morphogenesis in budding yeast. Annu. Rev. Microbiol. 52:687–744.
- 494. Madhani, H. D., C. A. Styles, and G. R. Fink. 1997. MAP kinases with distinct inhibitory functions impart signaling specificity during yeast differentiation. Cell 91:673–684.
- 495. Maheshwari, R. 1999. Microconidia of *Neurospora crassa*. Fungal Genet. Biol. 26:1–18.
- 496. Malagnac, F., L. Bartee, and J. Bender. 2002. An *Arabidopsis* SET domain protein required for maintenance but not establishment of DNA methylation. EMBO J. 21:6842–6852.
- 497. Malik, H. S., and S. Henikoff. 2002. Conflict begets complexity: the evolution of centromeres. Curr. Opin. Genet. Dev. 12:711–718.
- 498. Mannhaupt, G., C. Montrone, D. Haase, H. W. Mewes, V. Aign, J. D. Hoheisel, B. Fartmann, G. Nyakatura, F. Kempken, J. Maier, and U. Schulte. 2003. What's in the genome of a filamentous fungus? Analysis of the *Neurospora* genome sequence. Nucleic Acids Res. 31:1944–1954.
- 499. Marcotte, E. M., I. Xenarios, A. M. van Der Bliek, and D. Eisenberg. 2000. Localizing proteins in the cell from their phylogenetic profiles. Proc. Natl. Acad. Sci. USA 97:12115–12120.
- 500. Margolin, B. S., P. W. Garrett-Engele, J. N. Stevens, D. Yen-Fritz, C. Garrett-Engele, R. L. Metzenberg, and E. U. Selker. 1998. A methylated Neurospora 5S rRNA pseudogene contains a transposable element inactivated by RIP. Genetics 149:1787–1797.
- 501. Margolles-Clark, E. E., S. Abreu, and B. J. Bowman. 1999. Characterization of a vacuolar Ca²⁺/H⁺ exchanger (CAX) of *Neurospora crassa*. Fungal Genet. Newsl. 46(Suppl.):137.
- 502. Marszalek, J. R., J. A. Weiner, S. J. Farlow, J. Chun, and L. S. Goldstein. 1999. Novel dendritic kinesin sorting identified by different process targeting of two related kinesins: KIF21A and KIF21B. J. Cell. Biol. 145:469–479.
- Martinek, S., S. Inonog, A. S. Manoukian, and M. W. Young. 2001. A role for the segment polarity gene shaggy/GSK-3 in the *Drosophila* circadian clock. Cell 105:769–779.
- Marzluf, G. A. 1970. Genetic and metabolic controls for sulfate metabolism in *Neurospora crassa*: isolation and study of chromate-resistant and sulfate transport-negative mutants. J. Bacteriol. 102:716–721.
- 505. Marzluf, G. A. 1997. Genetic regulation of nitrogen metabolism in the fungi. Microbiol Mol. Biol. Rev. 61:17–32.
- Marzluf, G. A. 1997. Molecular genetics of sulfur assimilation in filamentous fungi and yeast. Annu. Rev. Microbiol. 51:73–96.
- 507. Mata, J., R. Lyne, G. Burns, and J. Bahler. 2002. The transcriptional program of meiosis and sporulation in fission yeast. Nat. Genet. 32:143–147.
- 508. Matsuura, S., H. Tauchi, A. Nakamura, N. Kondo, S. Sakamoto, S. Endo, D. Smeets, B. Solder, B. H. Belohradsky, V. M. Der Kaloustian, M. Oshimura, M. Isomura, Y. Nakamura, and K. Komatsu. 1998. Positional cloning of the gene for Nijmegen breakage syndrome. Nat. Genet. 19:179–181.
- 509. Mautino, M. R., and A. L. Rosa. 1998. Analysis of models involving enzymatic activities for the occurrence of C→T transition mutations during repeat-induced point mutation (RIP) in *Neurospora crassa*. J. Theor. Biol. 192:61–71.
- 510. Mazur, P., N. Morin, W. Baginsky, M. el-Sherbeini, J. A. Clemas, J. B. Nielsen, and F. Foor. 1995. Differential expression and function of two homologous subunits of yeast 1,3-beta-D-glucan synthase. Mol. Cell. Biol. 15:5671–5681.

- McBride, A. E., and P. A. Silver. 2001. State of the arg: protein methylation at arginine comes of age. Cell 106:5–8.
- McConnell, S. J., and M. P. Yaffe. 1992. Nuclear and mitochondrial inheritance in yeast depends on novel cytoplasmic structures defined by the MDM1 protein. J. Cell Biol. 118:385–395.
- 513. McGoldrick, C. A., C. Gruver, and G. S. May. 1995. *myoA* of *Aspergillus nidulans* encodes an essential myosin I required for secretion and polarized growth. J. Cell Biol. 128:577–587.
- 514. McKeon, T. A., M. Goodrich-Tanrikulu, J. T. Lin, and A. Stafford. 1997. Pathways for fatty acid elongation and desaturation in *Neurospora crassa*. Lipids 32:1–5.
- 515. McKim, K. S., J. B. Dahmus, and R. S. Hawley. 1996. Cloning of the Drosophila melanogaster meiotic recombination gene mei-218: a genetic and molecular analysis of interval 15E. Genetics 144:215–228.
- 516. Meesapyodsuk, D., D. W. Reed, C. K. Savile, P. H. Buist, U. A. Schafer, S. J. Ambrose, and P. S. Covello. 2000. Substrate specificity, regioselectivity and cryptoregiochemistry of plant and animal omega-3 fatty acid desaturases. Biochem. Soc. Trans. 28:632–635.
- 517. Mello, J. A., and G. Almouzni. 2001. The ins and outs of nucleosome assembly. Curr. Opin. Genet. Dev. 11:136–141.
- Melnick, M. B., C. Melnick, M. Lee, and D. O. Woodward. 1993. Structure and sequence of the calmodulin gene from *Neurospora crassa*. Biochim. Biophys. Acta 1171:334–336.
- 519. Meluh, P. B., and D. Koshland. 1995. Evidence that the MIF2 gene of Saccharomyces cerevisiae encodes a centromere protein with homology to the mammalian centromere protein CENP-C. Mol. Biol. Cell 6:793–807.
- 520. Meluh, P. B., P. Yang, L. Glowczewski, D. Koshland, and M. M. Smith. 1998. Cse4p is a component of the core centromere of *Saccharomyces cerevisiae*. Cell 94:607–613.
- 521. Meneghini, M. D., M. Wu, and H. D. Madhani. 2003. Conserved histone variant H2A.Z protects euchromatin from the ectopic spread of silent heterochromatin. Cell 112:725–736.
- 522. Merika, M., and S. H. Orkin. 1993. DNA-binding specificity of GATA family transcription factors. Mol. Cell. Biol. 13:3999–4010.
- 523. Merino, S. T., W. J. Cummings, S. N. Acharya, and M. E. Zolan. 2000. Replication-dependent early meiotic requirement for Spo11 and Rad50. Proc. Natl. Acad. Sci. USA 97:10477–10482.
- 524. Mette, M. F., W. Aufsatz, J. van Der Winden, M. A. Matzke, and A. J. Matzke. 2000. Transcriptional silencing and promoter methylation triggered by double-stranded RNA. EMBO J. 19:5194–5201.
- 525. Metzenberg, R. L., J. N. Stevens, E. U. Selker, and E. Morzycka-Wroblewska. 1985. Identification and chromosomal distribution of 5S rRNA genes in *Neurospora crassa*. Proc. Natl. Acad. Sci. USA 82:2067–2071.
- 526. Miao, V. P., M. Freitag, and E. U. Selker. 2000. Short TpA-rich segments of the zeta-eta region induce DNA methylation in *Neurospora crassa*. J. Mol. Biol. 300:249–273.
- 527. Miao, V. P. W., M. J. Singer, M. R. Rountree, and E. U. Selker. 1994. A targeted replacement system for identification of signals for de novo methylation in *Neurospora crassa*. Mol. Cell. Biol. 14:7059–7067.
- Mikolajczyk, S., and S. Brody. 1990. De novo fatty acid synthesis mediated by acyl-carrier protein in *Neurospora crassa* mitochondria. Eur. J. Biochem. 187:431–437.
- Milewski, S. 2002. Glucosamine-6-phosphate synthase—the multi-facets enzyme. Biochim. Biophys. Acta 1597:173–192.
- 530. Miller, R. K., K. K. Heller, L. Frisen, D. L. Wallack, D. Loayza, A. E. Gammie, and M. D. Rose. 1998. The kinesin-related proteins, Kip2p and Kip3p, function differently in nuclear migration in yeast. Mol. Biol. Cell 9:2051–2068.
- 531. Minehart, P. L., and B. Magasanik. 1984. Sequence and expression of *GLN3*, a positive regulator gene of *Saccharomyces cerevisiae* encoding a protein with a putative zinc finger DNA-binding domain. Mol. Cell. Biol. 11:6216–6228.
- 532. Miseta, A., R. Kellermayer, D. P. Aiello, L. Fu, and D. M. Bedwell. 1999. The vacuolar Ca²⁺/H⁺ exchanger Vex1p/Hum1p tightly controls cytosolic Ca²⁺ levels in *S. cerevisiae*. FEBS Lett. 451:132–136.
- Mishra, N. C. 1977. Genetics and biochemistry of morphogenesis in *Neurospora*. Adv. Genet. 19:341–405.
- Mitchell, A. P. 1994. Control of meiotic gene expression in *Saccharomyces cerevisiae*. Microbiol. Rev. 58:56–70.
- 535. Moazed, D. 2001. Common themes in mechanisms of gene silencing. Mol. Cell 8:489–498.
- 536. Montijn, R. C., E. Vink, W. H. Muller, A. J. Verkleij, H. Van Den Ende, B. Henrissat, and F. M. Klis. 1999. Localization of synthesis of beta1,6-glucan in *Saccharomyces cerevisiae*. J. Bacteriol. 181:7414–7420.
- 537. Mooney, J. L., and L. N. Yager. 1990. Light is required for conidiation in *Aspergillus nidulans*. Genes Dev. **4**:1473–1482.
- Morris, N. R. 2000. Nuclear migration. From fungi to the mammalian brain. J. Cell Biol. 148:1097–1101.
- 539. Morrow, G., Y. Inaguma, K. Kato, and R. M. Tanguay. 2000. The small heat shock protein Hsp22 of *Drosophila melanogaster* is a mitochondrial protein displaying oligomeric organization. J. Biol. Chem. 275:31204–31210.
- 540. Mösch, H. U. 2002. Pseudohyphal growth in yeast, p. 1-27. In H. D. Os-

iewacz (ed.), Molecular biology of fungal development. Marcel Dekker, Inc, New York, N.Y.

- 541. Moshkin, Y. M., J. A. Armstrong, R. K. Maeda, J. W. Tamkun, P. Verrijzer, J. A. Kennison, and F. Karch. 2002. Histone chaperone ASF1 cooperates with the Brahma chromatin-remodelling machinery. Genes Dev. 16:2621– 2626.
- 542. Muchardt, C., and M. Yaniv. 1999. The mammalian SWI/SNF complex and the control of cell growth. Semin. Cell Dev. Biol. 10:189–195.
- 543. Muldrow, T. A., A. M. Campbell, P. A. Weil, and D. T. Auble. 1999. MOT1 can activate basal transcription in vitro by regulating the distribution of TATA binding protein between promoter and nonpromoter sites. Mol. Cell. Biol. 19:2835–2845.
- 544. Mullen, J. R., V. Kaliraman, S. S. Ibrahim, and S. J. Brill. 2001. Requirement for three novel protein complexes in the absence of the Sgs1 DNA helicase in *Saccharomyces cerevisiae*. Genetics 157:103–118.
- 545. Muller, E. M., E. G. Locke, and K. W. Cunningham. 2001. Differential regulation of two Ca²⁺ influx systems by pheromone signaling in *Saccharomyces cerevisiae*. Genetics **159**:1527–1538.
- 546. Muller, J., C. M. Hart, N. J. Francis, M. L. Vargas, A. Sengupta, B. Wild, E. L. Miller, M. B. O'Connor, R. E. Kingston, and J. A. Simon. 2002. Histone methyltransferase activity of a *Drosophila* Polycomb group repressor complex. Cell 111:197–208.
- 547. Murayama, T., and T. Ishikawa. 1975. Characterization of *Neurospora crassa* mutants deficient in glucosephosphate isomerase. J. Bacteriol. 122: 54–58.
- Murayama, T., and T. Ishikawa. 1973. Mutation in *Neurospora crassa* affecting some of the extracellular enzymes and several growth characteristics. J. Bacteriol. 115:796–804.
- Murray, A., and T. Hunt. 1993. The cell cycle. Oxford University Press, New York, N.Y.
- 550. Murray, A. W., and D. Marks. 2001. Can sequencing shed light on cell cycling? Nature 409:844–846.
- 551. Murre, C., P. S. McCaw, and D. Baltimore. 1989. A new DNA binding and dimerization motif in immunoglobulin enhancer binding, daughterless, MyoD, and myc proteins. Cell 56:777–783.
- 552. Musacchio, A., and K. G. Hardwick. 2002. The spindle checkpoint: structural insights into dynamic signalling. Nat. Rev. Mol. Cell Biol. 3:731–741.
- 553. Nagy, P. L., J. Griesenbeck, R. D. Kornberg, and M. L. Cleary. 2002. A trithorax-group complex purified from *Saccharomyces cerevisiae* is required for methylation of histone H3. Proc. Natl. Acad. Sci. USA 99:90–94.
- Nakagawa, T., and R. D. Kolodner. 2002. The MER3 DNA helicase catalyzes the unwinding of holliday junctions. J. Biol. Chem. 277:28019–28024.
- 555. Nakagawa, T., and H. Ogawa. 1999. The Saccharomyces cerevisiae MER3 gene, encoding a novel helicase-like protein, is required for crossover control in meiosis. EMBO J. 18:5714–5723.
- 556. Nakamura, T., S. Muraoka, R. Sanokawa, and N. Mori. 1998. N-Shc and Sck, two neuronally expressed She adapter homologs. Their differential regional expression in the brain and roles in neurotrophin and Src signaling. J. Biol. Chem. 273:6960–6967.
- 557. Nakamura, T., Y. Yamaguchi, and H. Sano. 1999. Four rice genes encoding cysteine synthase: isolation and differential responses to sulfur, nitrogen and light. Gene 229:155–161.
- 558. Nakaoka, H., D. M. Perez, K. J. Baek, T. Das, A. Husain, K. Misono, M. J. Im, and R. M. Graham. 1994. Gh: a GTP-binding protein with transglutaminase activity and receptor signaling function. Science 264:1593–1596.
- 559. Nakayama, J., J. C. Rice, B. D. Strahl, C. D. Allis, and S. I. Grewal. 2001. Role of histone H3 lysine 9 methylation in epigenetic control of heterochromatin assembly. Science 292:110–113.
- 560. Nan, X., H. H. Ng, C. A. Johnson, C. D. Laherty, B. M. Turner, R. N. Eisenman, and A. Bird. 1998. Transcriptional repression by the methyl-CpG-binding protein MeCP2 involves a histone deacetylase complex. Nature 393:386–389.
- Nasmyth, K. 1996. At the heart of the budding yeast cell cycle. Trends Genet. 12:405–412.
- Neer, E. J. 1995. Heterotrimeric G proteins: organizers of transmembrane signals. Cell 80:249–257.
- 563. Nelson, D. L., and M. M. Cox. 2000. Lehninger principles of biochemistry, 3rd ed. Worth Publishers, New York, N.Y.
- 564. Nelson, M. A. 1996. Mating systems in ascomycetes: a romp in the sac. Trends Genet. 12:69–74.
- 565. Nelson, M. A., S. Kang, E. L. Braun, M. E. Crawford, P. L. Dolan, P. M. Leonard, J. Mitchell, A. M. Armijo, L. Bean, E. Blueyes, T. Cushing, A. Errett, M. Fleharty, M. Gorman, K. Judson, R. Miller, J. Ortega, I. Pavlova, J. Perea, S. Todisco, R. Trujillo, J. Valentine, A. Wells, M. Werner-Washburne, D. O. Natvig, et al. 1997. Expressed sequences from conidial, mycelial, and sexual stages of *Neurospora crassa*. Fungal Genet. Biol 21: 348–363.
- Nelson, W. J. 2003. Adaptation of core mechanisms to generate cell polarity. Nature 422:766–774.
- Neupert, W. 1997. Protein import into mitochondria. Annu. Rev. Biochem. 66:863–917.
- 568. Ng, H. H., Q. Feng, H. Wang, H. Erdjument-Bromage, P. Tempst, Y. Zhang,

and K. Struhl. 2002. Lysine methylation within the globular domain of histone H3 by Dot1 is important for telomeric silencing and Sir protein association. Genes Dev. 16:1518–1527.

- Nicholson, M. L., and D. E. Laudenbach. 1995. Genes encoded on a cyanobacterial plasmid are transcriptionally regulated by sulfur availability and CysR. J. Bacteriol. 177:2143–2150.
- Novak, J. E., P. B. Ross-Macdonald, and G. S. Roeder. 2001. The budding yeast Msh4 protein functions in chromosome synapsis and the regulation of crossover distribution. Genetics 158:1013–1025.
- 571. Nover, L., K. Bharti, P. Doring, S. K. Mishra, A. Ganguli, and K. D. Scharf. 2001. Arabidopsis and the heat stress transcription factor world: how many heat stress transcription factors do we need? Cell Stress Chaperones 6:177– 189.
- 572. Nowak, S. J., and V. G. Corces. 2000. Phosphorylation of histone H3 correlates with transcriptionally active loci. Genes Dev. 14:3003–3013.
- 573. O'Connell, M. J., P. B. Meluh, M. D. Rose, and N. R. Morris. 1993. Suppression of the *bimC4* mitotic spindle defect by deletion of *klpA*, a gene encoding a KAR3-related kinesin-like protein in *Aspergillus nidulans*. J. Cell. Biol. **120**:153–162.
- 574. O'Connell, M. J., A. H. Osmani, N. R. Morris, and S. A. Osmani. 1992. An extra copy of nimEcyclinB elevates pre-MPF levels and partially suppresses mutation of nimTcdc25 in *Aspergillus nidulans*. EMBO J. 11:2139–2149.
- 575. Orbach, M. J., W. P. Schneider, and C. Yanofsky. 1988. Cloning of methylated transforming DNA from *Neurospora crassa* in *Escherichia coli*. Mol. Cell. Biol. 8:2211–2213.
- 576. O'Rourke, R., S. Renault, W. Mo, and C. P. Selitrennikoff. 2003. Neurospora crassa FKS protein binds to the (1,3)β-glucan synthase substrate, UDPglucose. Curr. Microbiol. 46:408–412.
- 577. Ortiz, J., O. Stemmann, S. Rank, and J. Lechner. 1999. A putative protein complex consisting of Ctf19, Mcm21, and Okp1 represents a missing link in the budding yeast kinetochore. Genes Dev. 13:1140–1155.
- Osherov, N., R. A. Yamashita, Y. S. Chung, and G. S. May. 1998. Structural requirements for in vivo myosin I function in *Aspergillus nidulans*. J. Biol. Chem. 273:27017–27025.
- 579. Osiewacz, H. D. (ed.). 2002. Molecular biology of fungal development. Marcel Dekker, Inc., New York, NY.
- 580. Ouspenski, I. I., S. J. Elledge, and B. R. Brinkley. 1999. New yeast genes important for chromosome integrity and segregation identified by dosage effects on genome stability. Nucleic Acids Res. 27:3001–3008.
- 581. Owen, D. J., P. Ornaghi, J. C. Yang, N. Lowe, P. R. Evans, P. Ballario, D. Neuhaus, P. Filetici, and A. A. Travers. 2000. The structural basis for the recognition of acetylated histone H4 by the bromodomain of histone acetyl-transferase gen5p. EMBO J. 19:6141–6149.
- 582. Owens-Grillo, J. K., K. Hoffmann, K. A. Hutchison, A. W. Yem, M. R. Deibel, Jr., R. E. Handschumacher, and W. B. Pratt. 1995. The cyclosporin A-binding immunophilin CyP-40 and the FK506-binding immunophilin hsp56 bind to a common site on hsp90 and exist in independent cytosolic heterocomplexes with the untransformed glucocorticoid receptor. J. Biol. Chem. 270:20479–20484.
- 583. Paidhungat, M., and S. Garrett. 1997. A homolog of mammalian, voltagegated calcium channels mediates yeast pheromone-stimulated Ca²⁺ uptake and exacerbates the cdc1(Ts) growth defect. Mol. Cell. Biol. 17:6339–6347.
- Paietta, J. V. 1989. Molecular cloning and regulatory analysis of the arylsulfatase structural gene of *Neurospora crassa*. Mol. Cell. Biol. 9:3630–3637.
- 585. Paietta, J. V. 1992. Production of the CYS3 regulator, a bZIP DNA-binding protein, is sufficient to induce sulfur gene expression in *Neurospora crassa*. Mol. Cell. Biol. 12:1568–1577.
- 586. Paietta, J. V. 2004. Regulation of sulfur metabolism in mycelial fungi, p. 369–383. *In* R. Brambl and G. A. Marzluf (ed.), The mycota: biochemistry and molecular biology, in press. Springer-Verlag KG, Berlin, Germany.
- 587. Palacios, F., J. K. Schweitzer, R. L. Boshans, and C. D'Souza-Schorey. 2002. ARF6-GTP recruits Nm23-H1 to facilitate dynamin-mediated endocytosis during adherens junctions disassembly. Nat. Cell Biol. 4:929–936.
- 588. Paltauf, F., S. D. Kohlwein, and S. A. Henry. 1992. Regulation and compartmentalization of lipid synthesis in yeast, p. 415–500. *In E. W. Jones*, J. R. Pringle, and J. R. Broach (ed.), The molecular and cellular biology of the yeast *Saccharomyces*: gene expression, vol. 2. Cold Spring Laboratory Press, Cold Spring Harbor, N.Y.
- Paluh, J. L., and C. Yanofsky. 1991. Characterization of *Neurospora* CPC1, a bZIP DNA-binding protein that does not require aligned heptad leucines for dimerization. Mol. Cell. Biol. 11:935–944.
- 590. Pan, H. G., B. Feng, and G. A. Marzluf. 1997. Two distinct protein-protein interactions between the NIT2 and NMR regulatory proteins are required to establish nitrogen metabolite repression in *Neurospora crassa*. Mol. Microbiol. 26:721–729.
- 591. Pan, T., and J. E. Coleman. 1990. GAL4 transcription factor is not a "zinc finger" but forms a Zn(II)2Cys6 binuclear cluster. Proc. Natl. Acad. Sci. USA 87:2077-2081.
- 592. Panaretou, B., and P. W. Piper. 1992. The plasma membrane of yeast acquires a novel heat-shock protein (hsp30) and displays a decline in proton-pumping ATPase levels in response to both heat shock and the entry to stationary phase. Eur. J. Biochem. 206:635–640.

- 593. Pandey, R., A. Muller, C. A. Napoli, D. A. Selinger, C. S. Pikaard, E. J. Richards, J. Bender, D. W. Mount, and R. A. Jorgensen. 2002. Analysis of histone acetyltransferase and histone deacetylase families of *Arabidopsis thaliana* suggests functional diversification of chromatin modification among multicellular eukaryotes. Nucleic Acids Res. 30:5036–5055.
- 594. Pandit, N. N., and V. E. A. Russo. 1992. Reversible inactivation of a foreign gene, hph, during the asexual cycle in *Neurospora crassa* transformants. Mol. Gen. Genet. 234:412–422.
- 595. Papadaki, P., V. Pizon, B. Onken, and E. C. Chang. 2002. Two ras pathways in fission yeast are differentially regulated by two ras guanine nucleotide exchange factors. Mol. Cell. Biol. 22:4598–4606.
- 596. Paques, F., and J. E. Haber. 1999. Multiple pathways of recombination induced by double-strand breaks in *Saccharomyces cerevisiae*. Microbiol. Mol. Biol. Rev. 63:349–404.
- 597. Park, G., C. Xue, L. Zheng, and J. R. Xu. 2002. Mst12 regulates infectious growth but not appressorium formations in the rice blast fungus *Magnaporthe grisea*. Mol. Plant-Microbe Interact. 15:183–192.
- Park, H. O., J. Chant, and I. Herskowitz. 1993. BUD2 encodes a GTPaseactivating protein for Bud1/Rsr1 necessary for proper bud-site selection in yeast. Nature 365:269–274.
- 599. Park, I. C., H. Horiuchi, C. W. Hwang, W. H. Yeh, A. Ohta, J. C. Ryu, and M. Takagi. 1999. Isolation of csm1 encoding a class V chitin synthase with a myosin motor-like domain from the rice blast fungus. *Pyricularia oryzae*. FEMS Microbiol. Lett. **170**:131–139.
- Parkinson, J. S., and E. C. Kofoid. 1992. Communication modules in bacterial signaling proteins. Annu. Rev. Genet. 26:71–112.
- Parsell, D. A., A. S. Kowal, M. A. Singer, and S. Lindquist. 1994. Protein disaggregation mediated by heat-shock protein Hsp104. Nature 372:475– 478.
- 602. Parsons, R. B., D. Sampson, C. C. Huggins, R. H. Waring, A. C. Williams, and D. B. Ramsden. 2001. Renal localisation of rat cysteine dioxygenase. Nephron 88:340–346.
- Partridge, J. F., B. Borgstrom, and R. C. Allshire. 2000. Distinct protein interaction domains and protein spreading in a complex centromere. Genes Dev. 14:783–791.
- 604. Paul, A., S. Wilson, C. M. Belham, C. J. M. Robinson, P. H. Scott, G. W. Gould, and R. Pelvin. 1997. Stress-activated protein kinases: activation, regulation and function. Cell. Signaling 9:403–410.
- 605. Paulsen, I. T., M. K. Sliwinski, B. Nelissen, A. Goffeau, and M. H. Saier, Jr. 1998. Unified inventory of established and putative transporters encoded within the complete genome of *Saccharomyces cerevisiae*. FEBS Lett. 430: 116–125.
- 606. Pavesi, A. 1999. Relationships between transcriptional and translational control of gene expression in *Saccharomyces cerevisiae*: a multiple regression analysis. J. Mol. Evol. 48:133–141.
- 607. Payen, A. 1843. Extrain d'un rapport addresse' a' M. Le Marechal Duc de Dalmatie, Ministre de la Guerre, President du Conseil, sur une alteration extraordinaire du pain du munition. Ann. Chim. Phys. 3rd Ser. 9:5–21.
- Pereira, S. L., R. A. Grayling, R. Lurz, and J. N. Reeve. 1997. Archaeal nucleosomes. Proc. Natl. Acad. Sci. USA 94:12633–12637.
- 609. Perier, F., C. M. Radeke, K. F. Raab-Graham, and C. A. Vandenberg. 1995. Expression of a putative ATPase suppresses the growth defect of a yeast potassium transport mutant: identification of a mammalian member of the Clp/HSP104 family. Gene 152:157–163.
- Perkins, D. D. 1991. The first published scientific study of *Neurospora*, including a description of photoinduction of carotenoids. Fungal Genet. Newsl. 38:64–65.
- 611. Perkins, D. D., R. L. Metzenberg, N. B. Raju, E. U. Selker, and E. G. Barry. 1986. Reversal of a Neurospora translocation by crossing over involving displaced rDNA, and methylation of the rDNA segments that result from recombination. Genetics 114:791–817.
- 612. Perkins, D. D., A. Radford, and M. S. Sachs. 2001. The *Neurospora* compendium. Chromosomal loci. Academic Press, Inc., San Diego, Calif.
- 613. Petersen, J., and I. M. Hagan. 2003. S. pombe Aurora kinase/survivin is required for chromosome condensation and the spindle checkpoint attachment response. Curr. Biol. 13:590–597.
- Peyssonnaux, C., and A. Eychene. 2001. The Raf/MEK/ERK pathway: new concepts of activation. Biol. Cell. 93:53–62.
- Philley, M. L., and C. Staben. 1994. Functional analyses of the *Neurospora* crassa MT a-1 mating type polypeptide. Genetics 137:715–722.
- 616. Pimentel, B. E., R. Moreno-Sanchez, and C. Cervantes. 2002. Efflux of chromate by *Pseudomonas aeruginosa* cells expressing the ChrA protein. FEMS Microbiol. Lett. 212:249–254.
- Plesofsky, N., and R. Brambl. 1999. Glucose metabolism in *Neurospora* is altered by heat shock and by disruption of HSP30. Biochim. Biophys. Acta 1449:73–82.
- 618. Plesofsky, N., N. Gardner, R. Lill, and R. Brambl. 1999. Disruption of the gene for Hsp30, an alpha-crystallin-related heat shock protein of *Neuro-spora crassa*, causes defects in import of proteins into mitochondria. Biol. Chem. 380:1231–1236.
- 619. Plesofsky-Vig, N., and R. Brambl. 1998. Characterization of an 88-kDa heat

shock protein of Neurospora crassa that interacts with Hsp30. J. Biol. Chem. 273:11335-11341.

- 620. Plesofsky-Vig, N., and R. Brambl. 1995. Disruption of the gene for hsp30, an alpha-crystallin-related heat shock protein of Neurospora crassa, causes defects in thermotolerance. Proc. Natl. Acad. Sci. USA 92:5032-5036.
- 621. Plesofsky-Vig, N., and R. Brambl. 1990. Gene sequence and analysis of hsp30, a small heat shock protein of Neurospora crassa which associates with mitochondria. J. Biol. Chem. 265:15432-15440.
- 622. Poggeler, S., and U. Kuck. 2001. Identification of transcriptionally expressed pheromone receptor genes in filamentous ascomycetes. Gene 280: 9 - 17.
- 623. Pollard, T. D. 2001. Genomics, the cytoskeleton and motility. Nature 409: 842-843
- 624. Posas, F., and H. Saito. 1998. Activation of the yeast SSK2 MAP kinase kinase kinase by the SSK1 two-component response regulator. EMBO J. 17:1385–1394.
- 625. Posas, F., M. Takekawa, and H. Saito. 1998. Signal transduction by MAP kinase cascades in budding yeast. Curr. Opin. Microbiol. 1:175-182
- 626. Posas, F., S. M. Wurgler-Murphy, T. Maeda, E. A. Witten, T. C. Thai, and H. Saito. 1996. Yeast HOG1 MAP kinase cascade is regulated by a multistep phosphorelay mechanism in the SLN1-YPD1-SSK1 "two-component" osmosensor. Cell 86:865-875.
- 627. Pott, G. B., T. K. Miller, J. A. Bartlett, J. S. Palas, and C. P. Selitrennikoff. 2000. The isolation of FOS-1, a gene encoding a putative two-component histidine kinase from Aspergillus fumigatus. Fungal Genet. Biol. 31:55-67.
- Prakash, S., and L. Prakash. 2002. Translesion DNA synthesis in eu-628. karyotes: a one- or two-polymerase affair. Genes Dev. 16:1872-1883.
- 629. Preuss, U., G. Landsberg, and K. H. Scheidtmann. 2003. Novel mitosisspecific phosphorylation of histone H3 at Thr11 mediated by Dlk/ZIP kinase. Nucleic Acids Res. 31:878-885.
- 630. Prodromou, C., B. Panaretou, S. Chohan, G. Siligardi, R. O'Brien, J. E. Ladbury, S. M. Roe, P. W. Piper, and L. H. Pearl. 2000. The ATPase cycle of Hsp90 drives a molecular "clamp' via transient dimerization of the N-terminal domains. EMBO J. 19:4383-4392.
- 631. Prokisch, H., O. Yarden, M. Dieminger, M. Tropschug, and I. B. Barthelmess. 1997. Impairment of calcineurin function in Neurospora crassa reveals its essential role in hyphal growth, morphology and maintenance of the apical Ca²⁺ gradient. Mol. Gen. Genet. 256:104-114.
- 632. Pruyne, D., and A. Bretscher. 2000. Polarization of cell growth in yeast. J. Cell Sci. 113:571-585.
- 633. Pruyne, D., and A. Bretscher. 2000. Polarization of cell growth in yeast. I. Establishment and maintenance of polarity states. J. Cell Sci. 113:365-375.
- 634. Pruyne, D., M. Evangelista, C. Yang, E. Bi, S. Zigmond, A. Bretscher, and C. Boone. 2002. Role of formins in actin assembly: nucleation and barbedend association. Science 297:612-615.
- 635. Pu, R. T., G. Xu, L. Wu, J. Vierula, K. O'Donnell, X. S. Ye, and S. A. Osmani. 1995. Isolation of a functional homolog of the cell cycle-specific NIMA protein kinase of Aspergillus nidulans and functional analysis of conserved residues. J. Biol. Chem. 270:18110-18116.
- 636. Punt, P. J., B. Seiboth, X. O. Weenink, C. van Zeijl, M. Lenders, C. Konetschny, A. F. Ram, R. Montijn, C. P. Kubicek, and C. A. van den Hondel. 2001. Identification and characterization of a family of secretionrelated small GTPase-encoding genes from the filamentous fungus Aspergillus niger: a putative SEC4 homologue is not essential for growth. Mol. Microbiol. 41:513-525.
- 637. Raitt, D. C., A. L. Johnson, A. M. Erkine, K. Makino, B. Morgan, D. S. Gross, and L. H. Johnston. 2000. The Skn7 response regulator of Saccharomyces cerevisiae interacts with Hsf1 in vivo and is required for the induction of heat shock genes by oxidative stress. Mol. Biol. Cell 11:2335-2347.
- 638. Raju, N. B. 1992. Genetic control of the sexual cycle in Neurospora. Mycol. Res. 96:241-262.
- 639. Ramakrishnan, V. 1995. The histone fold: evolutionary questions. Proc Natl Acad Sci USA 92:11328-11330
- 640. Ramon, A., M. I. Muro-Pastor, C. Scazzocchio, and R. Gonzalez. 2000. Deletion of the unique gene encoding a typical histone H1 has no apparent phenotype in Aspergillus nidulans. Mol. Microbiol. 35:223-233.
- 641. Rea, S., F. Eisenhaber, D. O'Carroll, B. D. Strahl, Z. W. Sun, M. Schmid, S. Opravil, K. Mechtler, C. P. Ponting, C. D. Allis, and T. Jenuwein. 2000. Regulation of chromatin structure by site-specific histone H3 methyltransferases. Nature 406:593-599.
- 642. Read, N. D. 1994. Cellular nature and multicellular morphogenesis in higher fungi, p. 251–269. In D. S. Ingram and A. Hudson (ed.), Shape and form in plants and fungi. Academic Press, Ltd., London, United Kingdom.
- 643. Read, N. D., and P. C. Hickey. 2001. The vesicle trafficking network and tip growth in fungal hyphae, p. 137–146. In A. Geitmann, M. Cresti, and I. B. Heath (ed.), Cell biology of plant and fungal tip growth. IOS Press, Amsterdam, The Netherlands.
- 644. Read, N. D., and E. R. Kalkman. 2003. Does endocytosis occur in fungal hyphae? Fungal Genet. Biol. 39:199-203.
- 645. Refojo, D., A. C. Liberman, D. Giacomini, A. Carbia Nagashima, M. Graciarena, C. Echenique, M. Paez Pereda, G. Stalla, F. Holsboer, and E. Arzt. 2003. Integrating systemic information at the molecular level: cross-talk

between steroid receptors and cytokine signaling on different target cells. Ann. N. Y. Acad. Sci. 992:196-204.

- 646. Rehling, P., N. Wiedemann, N. Pfanner, and K. N. Truscott. 2001. The mitochondrial import machinery for preproteins. Crit. Rev. Biochem. Mol. Biol. 36:291-336.
- 647. Reichmann, J. L., J. Heard, G. Martin, L. Reuber, C.-Z. Jiang, J. Keddie, L. Adam, O. Pineda, O. J. Ratliffe, R. R. Samaha, R. Creelman, M. Pilgrim, P. Broun, J. Z. Zhang, D. Ghandehari, B. K. Sherman, and G.-L. Yu. 2000. Arabidopsis transcription factors: genome-wide comparative analysis among eukaryotes. Science 290:2105-2110.
- 648. Requena, N., C. Alberti-Segui, E. Winzenburg, C. Horn, M. Schliwa, P. Philippsen, R. Liese, and R. Fischer. 2001. Genetic evidence for a microtubule-destabilizing effect of conventional kinesin and analysis of its consequences for the control of nuclear distribution in Aspergillus nidulans. Mol. Microbiol. 42:121-132.
- 649. Reuther, G. W., and C. J. Der. 2000. The Ras branch of small GTPases: Ras family members don't fall far from the tree. Curr. Opin. Cell Biol. 12:157-165.
- 650. Richard, I., O. Broux, V. Allamand, F. Fougerousse, N. Chiannikulchai, N. Bourg, L. Brenguier, C. Deváusd, P. Pasturaud, C. Roudaut, D. Hillaire, M. R. Passobueno, M. Zatz, J. A. Tischfield, M. Fardeau, C. E. Jackson, D. Cohen, and J. S. Beckmann. 1995. Mutations in the proteolytic enzyme calpain-3 cause limb-girdle muscular-dystrophy type-2A. Cell 81:27-40.
- 651. Richards, E. J., and S. C. Elgin. 2002. Epigenetic codes for heterochromatin formation and silencing: rounding up the usual suspects. Cell 108:489-500
- 652. Richardson, H., D. J. Lew, M. Henze, K. Sugimoto, and S. I. Reed. 1992. Cyclin-B homologs in Saccharomyces cerevisiae function in S phase and in G2. Genes Dev. 6:2021-2034.
- 653. Ridgway, P., and G. Almouzni. 2000. CAF-1 and the inheritance of chromatin states: at the crossroads of DNA replication and repair. J. Cell Sci. 113:2647-2658
- 654. Ridley, A. J. 2001. Rho family proteins: coordinating cell responses. Trends Cell Biol. 11:471-477.
- 655. Rieder, C. L., A. Schultz, R. Cole, and G. Sluder. 1994. Anaphase onset in vertebrate somatic cells is controlled by a checkpoint that monitors sister kinetochore attachment to the spindle. J. Cell Biol. 127:1301-1310.
- 656. Riquelme, M., C. G. Reynaga-Pena, G. Gierz, and S. Bartnicki-Garcia. 1998. What determines growth direction in fungal hyphae? Fungal Genet. Biol. 24:101-109.
- 657. Robertson, K. D., S. Ait-Si-Ali, T. Yokochi, P. A. Wade, P. L. Jones, and A. P. Wolffe. 2000. DNMT1 forms a complex with rb, E2F1 and HDAC1 and represses transcription from E2F-responsive promoters. Nat. Genet. **25:**338–342.
- 658. Robzyk, K., J. Recht, and M. A. Osley. 2000. Rad6-dependent ubiquitination of histone H2B in yeast. Science 287:501-504.
- 659. Rockmill, B., J. A. Engebrecht, H. Scherthan, J. Loidl, and G. S. Roeder. 1995. The yeast MER2 gene is required for chromosome synapsis and the initiation of meiotic recombination. Genetics 141:49-59.
- 660. Roguev, A., D. Schaft, A. Shevchenko, W. W. Pijnappel, M. Wilm, R. Aasland, and A. F. Stewart. 2001. The Saccharomyces cerevisiae Set1 complex includes an Ash2 homologue and methylates histone 3 lysine 4. EMBO I 20:7137-7148
- 661. Roh, D. H., B. Bowers, H. Riezman, and E. Cabib. 2002. Rholp mutations specific for regulation of beta $(1\rightarrow 3)$ glucan synthesis and the order of assembly of the yeast cell wall. Mol Microbiol 44:1167-1183.
- 662. Romano, N., and G. Macino. 1992. Quelling: transient inactivation of gene expression in Neurospora crassa by transformation with homologous sequences. Mol. Microbiol. 6:3343-3353.
- 663. Rose, M. D., L. M. Misra, and J. P. Vogel. 1989. KAR2, a karyogamy gene, is the yeast homolog of the mammalian BiP/GRP78 gene. Cell 57:1211-1221
- 664. Rountree, M. R., and E. U. Selker. 1997. DNA methylation inhibits elongation but not initiation of transcription in Neurospora crassa. Genes Dev. 11:2383-2395
- 665. Rowley, N., C. Prip-Buus, B. Westermann, C. Brown, E. Schwarz, B. Barrell, and W. Neupert. 1994. Mdj1p, a novel chaperone of the DnaJ family, is involved in mitochondrial biogenesis and protein folding. Cell 77:249-259
- 666. Roychowdhury, H. S., D. Wong, and M. Kapoor. 1992. hsp80 of Neurospora crassa: cDNA cloning, gene mapping, and studies of mRNA accumulation under stress. Biochem. Cell Biol. 70:1356-1367.
- 667. Ruhf, M. L., A. Braun, O. Papoulas, J. W. Tamkun, N. Randsholt, and M. Meister. 2001. The domino gene of Drosophila encodes novel members of the SWI2/SNF2 family of DNA-dependent ATPases, which contribute to the silencing of homeotic genes. Development **128**:1429–1441. 668. **Ruiz-Herrera**, J. (ed.). 1992. Fungal cell wall: structure, synthesis and as-
- sembly. CRC Press, Inc., Boca Raton, Fla.
- 669 Russell, P. J., K. D. Rodland, E. M. Rachlin, and J. A. McCloskey. 1987. Differential DNA methylation during the vegetative life cycle of Neurospora crassa. J. Bacteriol. 169:2902-2905.
- 670. Sachowics, R., S. Farlow, and L. S. B. Goldstein. 1999. Cloning and expres-

sion of kinesins from the thermophilic fungus *Thermomyces lanuginosus*. Protein Sci. 8:2705–2710.

- 671. Sadler, I., A. Chiang, T. Kurihara, J. Rothblatt, J. Way, and P. Silver. 1989. A yeast gene important for protein assembly into the endoplasmic reticulum and the nucleus has homology to DnaJ, an *Escherichia coli* heat shock protein. J. Cell Biol. 109:2665–2675.
- 672. Saitou, N., and M. Nei. 1987. The neighbor-joining method: a new method for reconstructing phylogenetic trees. Mol. Biol. Evol. 4:406–425.
- 673. Sakai, W., C. Ishii, and H. Inoue. 2002. The upr-1 gene encodes a catalytic subunit of the DNA polymerase zeta which is involved in damage-induced mutagenesis in *Neurospora crassa*. Mol. Genet. Genomics 267:401–408.
- 674. Sancar, G. B. 2000. Enzymatic photoreactivation: 50 years and counting. Mutat. Res. 451:25–37.
- 675. Sanchez, Y., and S. L. Lindquist. 1990. HSP104 required for induced thermotolerance. Science 248:1112–1115.
- 676. Sanders, D., J. Pelloux, C. Brownlee, and J. F. Harper. 2001. Calcium at the crossroads of signaling. Plant Cell 14(Suppl.):S401–S417.
- 677. Santos, B., and M. Snyder. 2000. Sbe2p and sbe22p, two homologous Golgi proteins involved in yeast cell wall formation. Mol. Biol. Cell 11:435–452.
- 678. Santos, B., and M. Snyder. 1997. Targeting of chitin synthase 3 to polarized growth sites in yeast requires Chs5p and Myo2p. J. Cell Biol. 136:95–110.
- 679. Santos, J. L., and K. Shiozaki. 2001. Fungal histidine kinases. Science STKE 2001:RE1.
- Sargent, M. L., and D. O. Woodward. 1969. Gene-enzyme relationships in Neurospora invertase. J. Bacteriol. 97:544–549.
- 681. Saris, N., H. Holkeri, R. A. Craven, C. J. Stirling, and M. Makarow. 1997. The Hsp70 homologue Lhs1p is involved in a novel function of the yeast endoplasmic reticulum, refolding and stabilization of heat-denatured protein aggregates. J. Cell Biol. 137:813–824.
- 682. Sass, P., J. Field, J. Nikawa, T. Toda, and M. Wigler. 1986. Cloning and characterization of the high-affinity cAMP phosphodiesterase of *Saccharomyces cerevisiae*. Proc. Natl. Acad. Sci. USA 83:9303–9307.
- 683. Schaeffer, H. J., A. D. Catling, S. T. Eblen, L. S. Collier, A. Krauss, and M. J. Weber. 1998. MP1: a MEK binding partner that enhances enzymatic activation of the MAP kinase cascade. Science 281:1668–1671.
- Schafer, D. A., and T. A. Schroer. 1999. Actin-related proteins. Annu. Rev. Cell Dev. Biol. 15:341–363.
- 685. Schjerling, P., and S. Holmberg. 1996. Comparative amino acid sequence analysis of the C6 zinc cluster family of transcriptional regulators. Nucleic Acids Res. 24:4599–4607.
- Schlecht, U., and M. Primig. 2003. Mining meiosis and gametogenesis with DNA microarrays. Reproduction 125:447–456.
- 687. Schmid, K. J., and D. Tautz. 1997. A screen for fast evolving genes from Drosophila. Proc. Natl. Acad. Sci. USA 94:9746–9750.
- Schmidt, A., and M. N. Hall. 1998. Signaling to the actin cytoskeleton. Annu. Rev. Cell Dev. Biol. 14:305–338.
- 689. Schmitz, H.-P., and J. J. Heinisch. 2003. Evolution, biochemistry and genetics of protein kinase C in fungi. Curr. Genet. 43:245–254.
- 690. Schotta, G., A. Ebert, V. Krauss, A. Fischer, J. Hoffmann, S. Rea, T. Jenuwein, R. Dorn, and G. Reuter. 2002. Central role of *Drosophila* SU (VAR)3–9 in histone H3-K9 methylation and heterochromatic gene silencing. EMBO J. 21:1121–1131.
- 691. Schreiber, S. L., and B. E. Bernstein. 2002. Signaling network model of chromatin. Cell 111:771–778.
- 692. Schroeder, A., H. Inoue, and M. S. Sachs. 1998. DNA repair in *Neurospora*, p. 503–538. *In* J. A. Nickoloff and M. F. Hoekstra (ed.), DNA repair in prokaryotes and lower eukaryotes, vol. 1. Humana Press, Totowa, N.J.
- 693. Schroer, T. A. 1996. Structure and function of dynactin. Semin. Cell Biol. 7:321–328.
- 694. Schuller, H. J., A. Hahn, F. Troster, A. Schutz, and E. Schweizer. 1992. Coordinate genetic control of yeast fatty acid synthase genes FAS1 and FAS2 by an upstream activation site common to genes involved in membrane lipid biosynthesis. EMBO J. 11:107–114.
- 695. Schumacher, M. M., C. S. Enderlin, and C. P. Selitrennikoff. 1997. The osmotic-1 locus of *Neurospora crassa* encodes a putative histidine kinase similar to osmosensors of bacteria and yeast. Curr. Microbiol. 34:340–347.
- 696. Schurter, B. T., S. S. Koh, D. Chen, G. J. Bunick, J. M. Harp, B. L. Hanson, A. Henschen-Edman, D. R. Mackay, M. R. Stallcup, and D. W. Aswad. 2001. Methylation of histone H3 by coactivator-associated arginine methyltransferase 1. Biochemistry 40:5747–5756.
- 697. Schwerdtfeger, C., and H. Linden. 2001. Blue light adaptation and desensitization of light signal transduction in *Neurospora crassa*. Mol. Microbiol. 39:1080–1087.
- Schwerdtfeger, C., and H. Linden. 2003. VIVID is a flavoprotein and serves as a fungal blue light photoreceptor for photoadaptation. EMBO J. 22: 4846–4855.
- 699. Sebald, W., P. Friedl, H. U. Schairer, and J. Hoppe. 1982. Structure and genetics of the H⁺-conducting F0 portion of the ATP synthase. Ann. N.Y. Acad. Sci. 402:28–44.
- 700. Seiler, S., J. Kirchner, C. Horn, A. Kallipolitou, G. Woehlke, and M. Schliwa. 2000. Cargo binding and regulatory sites in the tail of fungal conventional kinesin. Nat. Cell Biol. 2:333–338.

- 701. Seiler, S., F. E. Nargang, G. Steinberg, and M. Schliwa. 1997. Kinesin is essential for cell morphogenesis and polarized secretion in *Neurospora crassa*. EMBO J. 16:3025–3034.
- Seiler, S., M. Plamann, and M. Schliwa. 1999. Kinesin and dynein mutants provide novel insights into the roles of vesicle traffic during cell morphogenesis in *Neurospora*. Curr. Biol. 9:779–785.
- 703. Sekelsky, J. J., K. S. McKim, G. M. Chin, and R. S. Hawley. 1995. The Drosophila meiotic recombination gene mei-9 encodes a homologue of the yeast excision repair protein Rad1. Genetics 141:619–627.
- 704. Selitrennikoff, C. P., L. Alex, T. K. Miller, K. V. Clemons, M. I. Simon, and D. A. Stevens. 2001. COS-1, a putative two-component histidine kinase of *Candida albicans*, is an in vivo virulence factor. Med. Mycol. 39:69–74.
- Selitrennikoff, C. P., and M. Nakata. 2003. New cell wall targets for antifungal drugs. Curr. Opin. Investig. Drugs 4:200–205.
- Selker, E. U. 1990. DNA methylation and chromatin structure: a view from below. Trends Biochem. Sci. 15:103–107.
- 707. Selker, E. U. 1997. Epigenetic phenomena in filamentous fungi: useful paradigms or repeat-induced confusion? Trends Genet. 13:296–301.
- Selker, E. U. 1990. Premeiotic instability of repeated sequences in *Neurospora crassa*. Annu. Rev. Genet. 24:579–613.
- 709. Selker, E. U. 2002. Repeat-induced gene silencing in fungi. Adv. Genet. 46:439–450.
- Selker, E. U. 1991. Repeat-induced point mutation (RIP) and DNA methylation, p. 258–265. *In* J. W. Bennet and L. Lasure (ed.), More gene manipulations in fungi. Academic Press, Inc., New York, N.Y.
- 711. Selker, E. U., E. B. Cambareri, B. C. Jensen, and K. R. Haack. 1987. Rearrangement of duplicated DNA in specialized cells of *Neurospora*. Cell 51:741–752.
- 712. Selker, E. U., M. Freitag, G. O. Kothe, B. S. Margolin, M. R. Rountree, C. D. Allis, and H. Tamaru. 2002. Induction and maintenance of nonsymmetrical DNA methylation in *Neurospora* Proc. Natl. Acad. Sci. USA 99(Suppl. 4):16485–16490.
- Selker, E. U., and P. W. Garrett. 1988. DNA sequence duplications trigger gene inactivation in *Neurospora crassa*. Proc. Natl. Acad. Sci. USA 85:6870– 6874.
- 714. Selker, E. U., B. C. Jensen, and G. A. Richardson. 1987. A portable signal causing faithful DNA methylation de novo in *Neurospora crassa*. Science 238:48–53.
- 715. Selker, E. U., G. A. Richardson, P. W. Garrett-Engele, M. J. Singer, and V. Miao. 1993. Dissection of the signal for DNA methylation in the zeta-eta region of *Neurospora*. Cold Spring Harbor Symp. Quant. Biol. 58:323–329.
- 716. Selker, E. U., and J. N. Stevens. 1985. DNA methylation at asymmetric sites is associated with numerous transition mutations. Proc. Natl. Acad. Sci. USA 82:8114–8118.
- 717. Selker, E. U., N. A. Tountas, S. H. Cross, B. S. Margolin, J. G. Murphy, A. P. Bird, and M. Freitag. 2003. The methylated component of the *Neurospora crassa* genome. Nature 422:893.
- Sellers, J. R. 2000. Myosins: a diverse superfamily. Biochim. Biophys. Acta 1496:3–22.
- 719. Sells, M. A., and J. Chernoff. 1997. Emerging from the Pak: the p21activated protein kinase family. Trends Cell Biol. 7:162–167.
- 720. Shahinian, S., and H. Bussey. 2000. β-1,6-Glucan synthesis in Saccharomyces cerevisiae. Mol. Microbiol. 35:477–489.
- 721. Shahinian, S., G. J. Dijkgraaf, A. M. Sdicu, D. Y. Thomas, C. A. Jakob, M. Aebi, and H. Bussey. 1998. Involvement of protein N-glycosyl chain glucosylation and processing in the biosynthesis of cell wall beta-1,6-glucan of *Saccharomyces cerevisiae*. Genetics 149:843–856.
- 722. Shannon, M., A. T. Hamilton, L. Gordon, E. Branscomb, and L. Stubbs. 2003. Differential expansion of zinc-finger transcription factor loci in homologous human and mouse gene clusters. Genet. Res 6A:1097–1110.
- 723. Shaw, B. D., and H. C. Hoch. 2001. Biology of the fungal cell, p. 73–89. *In* R. J. Howard and N. A. R. Gow (ed.), The mycota VIII. Springer-Verlag KG, Berlin, Germany.
- 724. Shaw, N. M., and R. W. Harding. 1987. Intracellular and extracellular cyclic nucleotides in wild-type and white collar mutant strains of *Neurospora crassa*. Plant Physiol. 83:377–383.
- Shear, C. L., and B. O. Dodge. 1927. Life histories and heterothallism of the red bread-mold fungi of the *Monila sitophila* group. J. Agric. Res. 34:1019– 1042.
- 726. Shen, W. C., J. Wieser, T. H. Adams, and D. J. Ebbole. 1998. The Neurospora rca-1 gene complements an Aspergillus flbD sporulation mutant but has no identifiable role in Neurospora sporulation. Genetics 148:1031–1041.
- 727. Shen, X., and M. A. Gorovsky. 1996. Linker histone H1 regulates specific gene expression but not global transcription in vivo. Cell 86:475–483.
- Shen, X., G. Mizuguchi, A. Hamiche, and C. Wu. 2000. A chromatin remodelling complex involved in transcription and DNA processing. Nature 406:541–544.
- 729. Shen, X., L. Yu, J. W. Weir, and M. A. Gorovsky. 1995. Linker histones are not essential and affect chromatin condensation in vivo. Cell 82:47–56.
- 730. Shimura, M., Y. Ito, C. Ishii, H. Yajima, H. Linden, T. Harashima, A. Yasui, and H. Inoue. 1999. Characterization of a *Neurospora crassa* photol-

yase-deficient mutant generated by repeat induced point mutation of the phr gene. Fungal Genet. Biol. 28:12–20.

- 731. Shinohara, M. L., J. J. Loros, and J. C. Dunlap. 1998. Glyceraldehyde-3phosphate dehydrogenase is regulated on a daily basis by the circadian clock. J. Biol. Chem. 273:446–452.
- 732. Shiu, P. K., N. B. Raju, D. Zickler, and R. L. Metzenberg. 2001. Meiotic silencing by unpaired DNA. Cell 107:905–916.
- 733. Shrode, L. B., Z. A. Lewis, L. D. White, D. Bell-Pedersen, and D. J. Ebbole. 2001. vvd is required for light adaptation of conidiation-specific genes of *Neurospora crassa*. but not circadian conidiation. Fungal Genet. Biol. 32: 169–181.
- 734. Sierra, D. A., D. J. Gilbert, D. Householder, N. V. Grishin, K. Yu, P. Ukidwe, S. A. Barker, W. He, T. G. Wensel, G. Otero, G. Brown, N. G. Copeland, N. A. Jenkins, and T. M. Wilkie. 2002. Evolution of the regulators of G-protein signaling multigene family in mouse and human. Genomics 79:177–185.
- 735. Sietsma, J. H., A. Beth Din, V. Ziv, K. A. Sjollema, and O. Yarden. 1996. The localization of chitin synthase in membranous vesicles (chitosomes) in *Neurospora crassa*. Microbiology 142:1591–1596.
- Silberstein, S., G. Schlenstedt, P. A. Silver, and R. Gilmore. 1998. A role for the DnaJ homologue Scj1p in protein folding in the yeast endoplasmic reticulum. J. Cell Biol. 143:921–933.
- 737. Silverman-Gavrila, L. B., and R. R. Lew. 2002. An IP(3)-activated Ca²⁺ channel regulates fungal tip growth. J. Cell Sci. 115:5013–5025.
- 738. Silverman-Gavrila, L. B., and R. R. Lew. 2001. Regulation of the tip-high [Ca²⁺] gradient in growing hyphae of the fungus *Neurospora crassa*. Eur. J. Cell Biol. 80:379–390.
- 739. Simons, J. F., M. Ebersold, and A. Helenius. 1998. Cell wall 1,6-beta-glucan synthesis in Saccharomyces cerevisiae depends on ER glucosidases I and II, and the molecular chaperone BiP/Kar2p. EMBO J. 17:396–405.
- 740. Singer, M. S., A. Kahana, A. J. Wolf, L. L. Meisinger, S. E. Peterson, C. Goggin, M. Mahowald, and D. E. Gottschling. 1998. Identification of high-copy disruptors of telomeric silencing in saccharomyces cerevisiae. Genetics 150:613–632.
- 741. Sizemore, S. T., and J. V. Paietta. 2002. Cloning and Characterization of scon-3⁺, a new member of the *Neurospora crassa* sulfur regulatory system. Eukaryot. Cell 1:875–883.
- 742. Smith, A. V., and G. S. Roeder. 1997. The yeast Red1 protein localizes to the cores of meiotic chromosomes. J. Cell Biol. 136:957–967.
- 743. Smith, K. N., A. Penkner, K. Ohta, F. Klein, and A. Nicolas. 2001. B-type cyclins CLB5 and CLB6 control the initiation of recombination and synaptonemal complex formation in yeast meiosis. Curr. Biol. 11:88–97.
- 744. Smith, S. E., and D. J. Read. 1997. Mycorrhizal symbiosis. Academic Press, Inc., New York, N.Y.
- 745. Solscheid, B., and M. Tropschug. 2000. A novel type of FKBP in the secretory pathway of *Neurospora crassa*. FEBS Lett. 480:118–122.
- 746. Sondermann, H., A. K. Ho, L. L. Listenberger, K. Siegers, I. Moarefi, S. R. Wente, F. U. Hartl, and J. C. Young. 2002. Prediction of novel Bag-1 homologs based on structure/function analysis identifies Snl1p as an Hsp70 co-chaperone in *Saccharomyces cerevisiae*. J. Biol. Chem. 277:33220–33227.
- 747. Sonenberg, N., J. W. B. Hershey, and M. B. Mathews. 2000. Translational control of gene expression. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y.
- 748. Sorger, P. K., and H. C. Nelson. 1989. Trimerization of a yeast transcriptional activator via a coiled-coil motif. Cell 59:807–813.
- 749. Soshi, T., Y. Sakuraba, E. Kafer, and H. Inoue. 1996. The mus-8 gene of Neurospora crassa encodes a structural and functional homolog of the Rad6 protein of Saccharomyces cerevisiae. Curr. Genet. 30:224–231.
- 750. Spotswood, H. T., and B. M. Turner. 2002. An increasingly complex code. J. Clin. Investig. 110:577–582.
- Springer, M. L. 1993. Genetic control of fungal differentiation: the three sporulation pathways of *Neurospora crassa*. Bioessays 15:365–374.
- 752. Springer, M. L., and C. Yanofsky. 1989. A morphological and genetic analysis of conidiophore development in *Neurospora crassa*. Genes Dev. 3:559–571.
- 753. Stambrough, M., D. W. Rowen, and B. Magasanik. 1995. Role of GATA factors Gln3p and Nil1p of *Saccharomyces cerevisiae* in the expression of nitrogen- regulated genes. Proc. Natl. Acad. Sci. USA 92:9450–9454.
- 754. Stark, M. J. 1996. Yeast protein serine/threonine phosphatases: multiple roles and diverse regulation. Yeast 12:1647–1675.
- 755. Steinberg, G. 1997. A kinesin-like mechanoenzyme from the zygomycete syncephalastrum racemosum shares biochemical similarities with conventional kinesin from *Neurospora crassa*. Eur. J. Cell Biol. 73:124–131.
- Steinberg, G., and M. Schliwa. 1995. The *Neurospora* organelle motor: a distant relative of conventional kinesin with unconventional properties. Mol. Biol. Cell 6:1605–1618.
- 757. Steinberg, G., M. Schliwa, C. Lehmler, M. Bolker, R. Kahmann, and J. R. McIntosh. 1998. Kinesin from the plant pathogenic fungus ustilago maydis is involved in vacuole formation and cytoplasmic migration. J. Cell Sci. 111:2235–2246.
- Stock, A. M., V. L. Robinson, and P. N. Goudreau. 2000. Two-component signal transduction. Annu. Rev. Biochem. 69:183–215.

- 759. Stoler, S., K. C. Keith, K. E. Curnick, and M. Fitzgerald-Hayes. 1995. A mutation in CSE4, an essential gene encoding a novel chromatin-associated protein in yeast, causes chromosome nondisjunction and cell cycle arrest at mitosis. Genes Dev. 9:573–586.
- 760. Stone, P. J., A. J. Makoff, J. H. Parish, and A. Radford. 1993. Cloning and sequence analysis of the glucoamylase gene of *Neurospora crassa*. Curr. Genet 24:205–211.
- Strahl, B. D., and C. D. Allis. 2000. The language of covalent histone modifications. Nature 403:41–45.
- 762. Strahl, B. D., P. A. Grant, S. D. Briggs, Z. W. Sun, J. R. Bone, J. A. Caldwell, S. Mollah, R. G. Cook, J. Shabanowitz, D. F. Hunt, and C. D. Allis. 2002. Set2 is a nucleosomal histone H3-selective methyltransferase that mediates transcriptional repression. Mol. Cell. Biol. 22:1298–1306.
- 763. Strub, A., K. Rottgers, and W. Voos. 2002. The Hsp70 peptide-binding domain determines the interaction of the ATPase domain with Tim44 in mitochondria. EMBO J. 21:2626–2635.
- 764. Sullivan, D. S., S. Biggins, and M. D. Rose. 1998. The yeast centrin, cdc31p, and the interacting protein kinase, Kic1p, are required for cell integrity. J. Cell Biol. 143:751–765.
- 765. Sun, Z. W., and C. D. Allis. 2002. Ubiquitination of histone H2B regulates H3 methylation and gene silencing in yeast. Nature 418:104–108.
- 766. Szoor, B., V. Dombradi, P. Gergely, and Z. Feher. 1997. Purification and characterization of the catalytic subunit of protein phosphatase 1 from *Neurospora crassa*. Acta Biol. Hung. 48:289–302.
- 767. Szoor, B., Z. Feher, E. Bako, F. Erdodi, G. Szabo, P. Gergely, and V. Dombradi. 1995. Isolation and characterization of the catalytic subunit of protein phosphatase 2A from *Neurospora crassa*. Comp. Biochem. Physiol. Ser. B 112:515–522.
- 768. Szoor, B., Z. Feher, T. Zeke, P. Gergely, E. Yatzkan, O. Yarden, and V. Dombradi. 1998. pzl-1 encodes a novel protein phosphatase-Z-like Ser/Thr protein phosphatase in *Neurospora crassa*. Biochim. Biophys. Acta 1388: 260–266.
- 769. Tabara, H., M. Sarkissian, W. G. Kelly, J. Fleenor, A. Grishok, L. Timmons, A. Fire, and C. C. Mello. 1999. The *rde-1* gene, RNA interference, and transposon silencing in *C. elegans*. Cell **99**:123–132.
- 770. Tachibana, M., K. Sugimoto, T. Fukushima, and Y. Shinkai. 2001. Set domain-containing protein, G9a, is a novel lysine-preferring mammalian histone methyltransferase with hyperactivity and specific selectivity to lysines 9 and 27 of histone H3. J. Biol. Chem. 276:25309–25317.
- 771. Tachibana, M., K. Sugimoto, M. Nozaki, J. Ueda, T. Ohta, M. Ohki, M. Fukuda, N. Takeda, H. Niida, H. Kato, and Y. Shinkai. 2002. G9a histone methyltransferase plays a dominant role in euchromatic histone H3 lysine 9 methylation and is essential for early embryogenesis. Genes Dev. 16:1779–1791.
- 772. Takahashi, K., E. S. Chen, and M. Yanagida. 2000. Requirement of Mis6 centromere connector for localizing a CENP-A-like protein in fission yeast. Science 288:2215–2219.
- 773. Takai, Y., T. Sasaki, and T. Matozaki. 2001. Small GTP-binding proteins. Physiol. Rev. 81:153–208.
- 774. Takayama, S., Z. Xie, and J. C. Reed. 1999. An evolutionarily conserved family of Hsp70/Hsc70 molecular chaperone regulators. J. Biol. Chem. 274:781–786.
- 775. Taleb, F., and A. Radford. 1995. The cellulase complex of *Neurospora* crassa: cbh-1 cloning, sequencing and homologies. Gene 161:137–138.
- 776. Tamaru, H., and E. U. Selker. 2001. A histone H3 methyltransferase controls DNA methylation in *Neurospora crassa*. Nature 414:277–283.
- 777. Tamaru, H., and E. U. Selker. 2003. Synthesis of signals for de novo DNA methylation in *Neurospora crassa*. Mol. Cell. Biol. 23:2379–2394.
- 778. Tamaru, H., X. Zhang, D. McMillen, P. B. Singh, J. Nakayama, S. I. Grewal, C. D. Allis, X. Cheng, and E. U. Selker. 2003. Trimethylated lysine 9 of histone H3 is a mark for DNA methylation in *Neurospora crassa*. Nat. Genet. 34:75–79.
- 779. Tamaskovic, R., S. J. Bichsel, and B. A. Hemmings. 2003. NDR family of AGC kinases—essential regulators of the cell cycle and morphogenesis. FEBS Lett. 546:73–80.
- 780. Tanabe, M., N. Sasai, K. Nagata, X. D. Liu, P. C. Liu, D. J. Thiele, and A. Nakai. 1999. The mammalian *HSF4* gene generates both an activator and a repressor of heat shock genes by alternative splicing. J. Biol. Chem. 274: 27845–27856.
- 781. Tang, J., J. D. Gary, S. Clarke, and H. R. Herschman. 1998. PRMT 3, a type I protein arginine N-methyltransferase that differs from PRMT1 in its oligomerization, subcellular localization, substrate specificity, and regulation. J. Biol. Chem. 273:16935–16945.
- 782. Tanton, L. T., C. E. Nargang, K. E. Kessler, Q. Li, and F. E. Nargang. 2003. Alternative oxidase expression in *Neurospora crassa*. Fungal Genet. Biol. 39:176–190.
- 783. Taylor, B. L., and I. B. Zhulin. 1999. PAS domains: internal sensors of oxygen, redox potential, and light. Microbiol. Mol. Biol. Rev. 63:479–506.
- 784. Taylor, S. W., E. Fahy, and S. S. Ghosh. 2003. Global organellar proteomics. Trends Biotechnol. 21:82–88.
- 785. Taylor, S. W., E. Fahy, B. Zhang, G. M. Glenn, D. E. Warnock, S. Wiley, A. N. Murphy, S. P. Gaucher, R. A. Capaldi, B. W. Gibson, and S. S. Ghosh.

2003. Characterization of the human heart mitochondrial proteome. Nat. Biotechnol. **21:**281–286.

- 786. Teather, R. M., and P. J. Wood. 1982. Use of Congo red-polysaccharide interactions in enumeration and characterization of cellulolytic bacteria from the bovine rumen. Appl. Environ. Microbiol. 43:777–780.
- 787. Tellez de Inon, M. T., and H. N. Torres. 1973. Regulation of glycogen phosphorylase a phosphatase in *Neurospora crassa*. Biochim. Biophys. Acta 297:399–412.
- Thomas, D., and Y. Surdin-Kerjan. 1997. Metabolism of sulfur amino acids in Saccharomyces cerevisiae. Microbiol. Mol. Biol. Rev. 61:503–532.
- 789. Thorne, A. W., P. Sautiere, G. Briand, and C. Crane-Robinson. 1987. The structure of ubiquitinated histone H2B. EMBO J. 6:1005–1010.
- 790. Toda, T., S. Cameron, P. Sass, M. Zoller, and M. Wigler. 1987. Three different genes in *S. cerevisiae* encode the catalytic subunits of the cAMPdependent protein kinase. Cell 50:277–287.
- 791. Toda, T., I. Uno, T. Ishikawa, S. Powers, T. Kataoka, D. Broek, S. Cameron, J. Broach, K. Matsumoto, and M. Wigler. 1985. In yeast, *RAS* proteins are controlling elements of adenylate cyclase. Cell 40:27–36.
- 792. Tomita, H., T. Soshi, and H. Inoue. 1993. The Neurospora uvs-2 gene encodes a protein which has homology to yeast RAD18, with uique zinc finger motifs. Mol. Gen. Genet. 238:225–233.
- 793. Tomomori, C., T. Tanaka, R. Dutta, H. Park, S. K. Saha, Y. Zhu, R. Ishima, D. Liu, K. I. Tong, H. Kurokawa, H. Qian, M. Inouye, and M. Ikura. 1999. Solution structure of the homodimeric core domain of *Escherichia coli* histidine kinase EnvZ. Nat. Struct. Biol. 6:729–734.
- 794. Tong, J. K., C. A. Hassig, G. R. Schnitzler, R. E. Kingston, and S. L. Schreiber. 1998. Chromatin deacetylation by an ATP-dependent nucleosome remodelling complex. Nature 395:917–921.
- 795. Torralba, S., and I. B. Health. 2002. Analysis of three separate probes suggests the absence of endocytosis in *Neurospora crassa* hyphae. Fungal Genet. Biol. 37:221–232.
- 796. Tran, H. G., D. J. Steger, V. R. Iyer, and A. D. Johnson. 2000. The chromo domain protein chd1p from budding yeast is an ATP-dependent chromatinmodifying factor. EMBO J. 19:2323–2331.
- 797. Trilla, J. A., A. Duran, and C. Roncero. 1999. Chs7p, a new protein involved in the control of protein export from the endoplasmic reticulum that is specifically engaged in the regulation of chitin synthesis in *Saccharomyces cerevisiae*. J. Cell Biol. 145:1153–1163.
- 798. Trinci, A. P. J. 1984. Regulation of hyphal branching and hyphal orientation, p. 23–52. *In* D. H. Jennings and A. D. M. Rayner (ed.), The ecology and physiology of the fungal mycelium. Cambridge University Press, Cambridge, United Kingdom.
- Trinci, A. P. J., and P. Saunders. 1977. Tip growth of fungal hyphae. J. Gen Microbiol. 103:243–248.
- 800. Trinci, A. P. J., M. G. Wiebe, and G. D. Robson. 1994. The mycelium as an integrated entity, p. 173–193. *In J. G. H. Wessels and F. Meinhardt (ed.)*, The mycota: growth, differentiation and sexuality, vol. 1. Springer-Verlag KG, Berlin, Germany.
- 801. Tropschug, M., D. W. Nicholson, F. U. Hartl, H. Kohler, N. Pfanner, E. Wachter, and W. Neupert. 1988. Cyclosporin A-binding protein (cyclophilin) of *Neurospora crassa*. One gene codes for both the cytosolic and mitochondrial forms. J. Biol. Chem. 263:14433–14440.
- 802. Tropschug, M., E. Wachter, S. Mayer, E. R. Schonbrunner, and F. X. Schmid. 1990. Isolation and sequence of an FK506-binding protein from *N. crassa* which catalyses protein folding. Nature 346:674–677.
- 803. Trotter, E. W., C. M. Kao, L. Berenfeld, D. Botstein, G. A. Petsko, and J. V. Gray. 2002. Misfolded proteins are competent to mediate a subset of the responses to heat shock in *Saccharomyces cerevisiae*. J. Biol. Chem. 277: 44817–44825.
- 804. Tsai, S.-F., D. I. K. Martin, L. I. Zon, A. D. D'Andrea, G. G. Wong, and S. H. Orkin. 1989. Cloning of cDNA for the major DNA-binding protein of the erythroid lineage through expression in mammalian cells. Nature 328: 827–830.
- 805. Tsuji, G., S. Fugii, S. Tsuge, T. Shiraishi, and Y. Kubo. 2003. The Colletotrichum lagenarium Ste12-like gene CST1 is essential for appressorium penetration. Mol. Plant-Microbe Interact. 16:215–225.
- Tung, K. S., and G. S. Roeder. 1998. Meiotic chromosome morphology and behavior in zip1 mutants of *Saccharomyces cerevisiae*. Genetics 149:817– 832.
- 807. Reference deleted.
- Turian, G., and D. E. Bianchi. 1972. Conidiation in *Neurospora*. Bot. Rev. 38:119–154.
- Turner, B. C., D. D. Perkins, and A. Fairfield. 2001. Neurospora from natural populations: a global study. Fungal Genet. Biol. 32:67–92.
- 810. Turner, B. M. 2002. Cellular memory and the histone code. Cell 111:285–291.
- Turner, B. M. 2000. Histone acetylation and an epigenetic code. Bioessays 22:836–845.
- 812. Turner, G., and S. D. Harris. 1997. Genetic control of polarized growth and branching in filamentous fungi, p. 229–260. *In* N. A. R. Gow, G. D. Robson, and G. M. Gadd (ed.), The fungal colony. Cambridge University Press, Cambridge, United Kingdom.

- 813. Tyler, J. K., C. R. Adams, S. R. Chen, R. Kobayashi, R. T. Kamakaka, and J. T. Kadonaga. 1999. The RCAF complex mediates chromatin assembly during DNA replication and repair. Nature 402:555–560.
- 814. Tzagoloff, A., G. Macino, and W. Sebald. 1979. Mitochondrial genes and translation products. Annu. Rev. Biochem. 48:419–441.
- 815. Uria-Nickelsen, M. R., E. R. Leadbetter, and W. Godchaux, 3rd. 1993. Sulfonate-sulfur assimilation by yeasts resembles that of bacteria. FEMS Microbiol. Lett. 114:73–77.
- Urrutia, A. O., and L. D. Hurst. 2001. Codon usage bias covaries with expression breadth and the rate of synonymous evolution in humans, but this is not evidence for selection. Genetics 159:1191–1199.
- 817. Ushinsky, S. C., H. Bussey, A. A. Ahmed, Y. Wang, J. Friesen, B. A. Williams, and R. K. Storms. 1997. Histone H1 in Saccharomyces cerevisiae. Yeast 13:151–161.
- Vallee, R. B., N. E. Faulkner, and C. Y. Tai. 2000. The role of cytoplasmic dynein in the human brain developmental disease lissencephaly. Biochim. Biophys. Acta. Mol. Cell Res. 1496:89–98.
- Vallim, M. A., K. Y. Miller, and B. L. Miller. 2000. Aspergillus SteA (Sterile 12-like) is a homeodomain-C2/H2-Zn⁺² finger transcription factor required for sexual reproduction. Mol. Microbiol. 36:290–301.
- Van Aelst, L., and C. D'Souza-Schorey. 1997. Rho GTPases and signaling networks. Genes Dev. 11:2295–2322.
- 821. van Biesen, T., L. M. Luttrell, B. E. Hawes, and R. J. Lefkowitz. 1996. Mitogenic signaling via G protein-coupled receptors. Endocr. Rev. 17:698– 714.
- 822. Vance, J. E. 1998. Eukaryotic lipid-biosynthetic enzymes: the same but not the same. Trends Biochem. Sci. 23:423–428.
- 823. van den Boogaart, P., J. Samallo, S. Van Dijck, and E. Agsteribbe. 1982. Structural and functional analyses of the genes for subunit II of cytochrome aa3 and for a dicyclohexylcarbodiimide-binding protein in *Neurospora crassa* mitochondrial DNA, p. 375–380. *In* P. Slonimski, P. Borst, and G. Attardi (ed.), Mitochondrial genes. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y.
- 824. van der Ploeg, J. R., M. A. Weiss, E. Saller, H. Nashimoto, N. Saito, M. A. Kertesz, and T. Leisinger. 1996. Identification of sulfate starvation-regulated genes in *Escherichia coli*: a gene cluster involved in the utilization of taurine as a sulfur source. J. Bacteriol. 178:5438–5446.
- 825. van der Westhuizen, J., J. Kock, A. Botha, and P. Botes. 1994. The distribution of the ω3- and ω6-series of cellular long-chain fatty acids in fungi. Syst. Appl. Microbiol. 17:327–345.
- 826. van Holde, K. C. 1989. Chromatin. Springer-Verlag KG, Berlin, Germany.
- 827. Van Hooser, A. A., M. A. Mancini, C. D. Allis, K. F. Sullivan, and B. R. Brinkley. 1999. The mammalian centromere: structural domains and the attenuation of chromatin modeling. FASEB J. 13(Suppl. 2):S216–S220.
- Vanlerberghe, G., and L. McIntosh. 1996. Signals regulating the expression of the nuclear gene encoding alternative oxidase of plant mitochondria. Plant Physiol. 111:589–595.
- 829. Vasara, T., L. Salusjarvi, M. Raudaskoski, S. Keranen, M. Penttila, and M. Saloheimo. 2001. Interactions of the *Trichoderma reesei rho3* with the secretory pathway in yeast and *T. reesei*. Mol. Microbiol. 42:1349–1361.
- Verhey, K. J., D. L. Lizotte, T. Abramson, L. Barenboim, B. J. Schnapp, and T. A. Rapoport. 1998. Light chain-dependent regulation of Kinesin's interaction with microtubules. J. Cell Biol. 143:1053–1066.
- Verhey, K. J., and T. A. Rapoport. 2001. Kinesin carries the signal. Trends Biochem. Sci. 26:545–550.
- Vinson, C., M. Myakishev, A. Acharya, A. A. Mir, J. R. Moll, and M. Bonovich. 2002. Classification of human B-ZIP proteins based on dimerization properties. Mol. Cell. Biol. 22:6321–6335.
- Vinson, C. R., P. B. Sigler, and S. L. McKnight. 1989. A scissors-grip model for DNA recognition by a family of leucine zipper proteins. Science 246: 911–916.
- 834. Virginia, M., C. L. Appleyard, W. L. McPheat, and M. J. Stark. 2000. A novel "two-component" protein containing histidine kinase and response regulator domains required for sporulation in *Aspergillus nidulans*. Curr. Genet. 37:364–372.
- 835. Voisard, C., J. Wang, J. L. McEvoy, P. Xu, and S. Leong. 1993. urbs, a gene regulating siderophore biosynthesis in Ustilago maydis, encodes a protein similar to the erythroid transcription factor GATA-1. Mol. Cell. Biol. 13: 7091–7100.
- 836. Voisine, C., Y. C. Cheng, M. Ohlson, B. Schilke, K. Hoff, H. Beinert, J. Marszalek, and E. A. Craig. 2001. Jac1, a mitochondrial J-type chaperone, is involved in the biogenesis of Fe/S clusters in *Saccharomyces cerevisiae*. Proc. Natl. Acad. Sci. USA 98:1483–1488.
- 837. Volpe, T. A., C. Kidner, I. M. Hall, G. Teng, S. I. Grewal, and R. A. Martienssen. 2002. Regulation of heterochromatic silencing and histone H3 lysine-9 methylation by RNAi. Science 297:1833–1837.
- Von Heijne, G. 1986. Mitochondrial targeting sequences may form amphiphilic helices. EMBO J. 5:1335–1342.
- 839. Wade, P. A., A. Gegonne, P. L. Jones, E. Ballestar, F. Aubry, and A. P. Wolffe. 1999. Mi-2 complex couples DNA methylation to chromatin remodelling and histone deacetylation. Nat. Genet. 23:62–66.
- 840. Wade, P. A., P. L. Jones, D. Vermaak, and A. P. Wolffe. 1998. A multiple

subunit Mi-2 histone deacetylase from *Xenopus laevis* cofractionates with an associated Snf2 superfamily ATPase. Curr. Biol. 8:843–846.

- 841. Wainright, M. 1992. An introduction to fungal biotechnology. John Wiley & Sons, Inc., New York, N.Y.
- 842. Wallis, J. G., J. L. Watts, and J. Browse. 2002. Polyunsaturated fatty acid synthesis: what will they think of next? Trends Biochem. Sci. 27:467.
- 843. Wang, H., Z. Q. Huang, L. Xia, Q. Feng, H. Erdjument-Bromage, B. D. Strahl, S. D. Briggs, C. D. Allis, J. Wong, P. Tempst, and Y. Zhang. 2001. Methylation of histone H4 at arginine 3 facilitating transcriptional activation by nuclear hormone receptor. Science 293:853–857.
- 844. Wang, X. Y., X. Chen, H. J. Oh, E. Repasky, L. Kazim, and J. Subjeck. 2000. Characterization of native interaction of hsp110 with hsp25 and hsc70. FEBS Lett. 465:98–102.
- 845. Watters, M. K., T. A. Randall, B. S. Margolin, E. U. Selker, and D. R. Stadler. 1999. Action of repeat-induced point mutation on both strands of a duplex and on tandem duplications of various sizes in *Neurospora*. Genetics 153:705–714.
- 846. Wedlich-Soldner, R., A. Straube, M. W. Friedrich, and G. Steinberg. 2002. A balance of KIF1A-like kinesin and dynein organizes early endosomes in the fungus *Ustilago maydis*. EMBO J. 21:2946–2957.
- 847. Welton, R. M., and C. S. Hoffman. 2000. Glucose monitoring in fission yeast via the Gpa2 galpha, the git5 Gbeta and the git3 putative glucose receptor. Genetics 156:513–521.
- Wendland, J., and P. Philippsen. 2001. Cell polarity and hyphal morphogenesis are controlled by multiple rho-protein modules in the filamentous ascomycete Ashbya gossypii. Genetics 157:601–610.
- 849. Wendland, J., and P. Philippsen. 2000. Determination of cell polarity in germinated spores and hyphal tips of the filamentous ascomycete *Ashbya* gossypii requires a rhoGAP homolog. J. Cell Sci. 113:1611–1621.
- Wenzel, R. P., and M. A. Pfaller. 1991. Candida species: emerging hospital bloodstream pathogens. Infect. Control Hosp. Epidemiol. 12:523–524.
- Wessels, J. G. H. 1986. Cell wall synthesis in apical hyphal growth. Int. Rev. Cytol. 104:7–79.
- Westwood, J. T., and C. Wu. 1993. Activation of *Drosophila* heat shock factor: conformational change associated with a monomer-to-trimer transition. Mol. Cell. Biol. 13:3481–3486.
- 853. Wigge, P. A., and J. V. Kilmartin. 2001. The Ndc80p complex from Saccharomyces cerevisiae contains conserved centromere components and has a function in chromosome segregation. J. Cell Biol. 152:349–360.
- 854. Williams, R. M., M. Primig, B. K. Washburn, E. A. Winzele, M. Bellis, C. Sarrauste de Menthiere, R. W. Davis, and R. E. Esposito. 2002. The Ume6 regulon coordinates metabolic and meiotic gene expression in yeast. Proc. Natl. Acad. Sci. USA 99:13431–13436.
- Wilson, T. E., U. Grawunder, and M. R. Lieber. 1997. Yeast DNA ligase IV mediates non-homologous DNA end joining. Nature 388:495–498.
- 856. Win, T. Z., D. P. Mulvihill, and J. S. Hyams. 2002. Take five: a myosin class act in fission yeast. Cell Motil. Cytoskeleton 51:53–56.
- 857. Wolanin, P. M., P. A. Thomason, and J. B. Stock. 2002. Histidine protein kinases: key signal transducers outside the animal kingdom. Genome Biol. 3:3013.1–3013.8.
- Wolffe, A. P., and M. A. Matzke. 1999. Epigenetics: regulation through repression. Science 286:481–486.
- 859. Woodage, T., M. A. Basrai, A. D. Baxevanis, P. Hieter, and F. S. Collins. 1997. Characterization of the CHD family of proteins. Proc. Natl. Acad. Sci. USA 94:11472–11477.
- 860. Woudt, L. P., A. Pastink, A. E. Kempers-Veenstra, A. E. M. Jansen, W. H. Mager, and R. J. Planta. 1983. The genes coding for histone H3 and H4 in *Neurospora crassa* are unique and contain intervening sequences. Nucleic Acids Res. 11:5347–5361.
- 861. Wu, K., J. H. Dawe, and J. P. Aris. 2000. Expression and subcellular localization of a membrane protein related to Hsp30p in *Saccharomyces cerevisiae*. Biochim. Biophys. Acta 1463:477–482.
- 862. Wu, Q. D., T. M. Sandrock, B. G. Turgeon, O. C. Yoder, S. G. Wirsel, and J. R. Aist. 1998. A fungal kinesin required for organelle motility, hyphal growth, and morphogenesis. Mol. Biol. Cell 9:89–101.
- Xiang, X., and N. R. Morris. 1999. Hyphal tip growth and nuclear migration. Curr. Opin. Microbiol. 2:636–640.
- 864. Xu, J. R., C. J. Staiger, and J. E. Hamer. 1998. Inactivation of the mitogenactivated protein kinase Mps1 from the rice blast fungus prevents penetration of host cells but allows activation of plant defense responses. Proc. Natl. Acad. Sci. USA 95:12713–12718.
- 865. Xu, Z., A. L. Horwich, and P. B. Sigler. 1997. The crystal structure of the asymmetric GroEL-GroES-(ADP)7 chaperonin complex. Nature 388:741– 750.
- 866. Xue, Y., M. Batlle, and J. P. Hirsch. 1998. *GPR1* encodes a putative G protein-coupled receptor that associates with the Gpa2p $G\alpha$ subunit and functions in a Ras-independent pathway. EMBO J. 17:1996–2007.
- 867. Yajima, H., H. Inoue, A. Oikawa, and i. A. Yasu. 1991. Cloning and functional characterization of a eucaryotic DNA photolyase gene from *Neuro*spora crassa. Nucleic Acids Res. 19:5359–5362.
- 868. Yajima, H., M. Takao, S. Yasuhira, J. H. Zhao, C. Ishii, H. Inoue, and A.

Yasui. 1995. A eukaryotic gene encoding an endonuclease that specifically repairs DNA damaged by ultraviolet light. EMBO J. 14:2393–2399.

- 869. Yamashiro, C. T., D. J. Ebbole, B.-K. Lee, R. E. Brown, Bourland C., L. Madi, and C. Yanofsky. 1996. Characterization of *rco-1* of *Neurospora crassa*, a pleiotropic gene affecting growth and development that encodes a homolog of Tup1 of *Saccharomyces cerevisiae*. Mol. Cell. Biol. 16:6218–6228.
- Yamashita, R. A., and G. S. May. 1998. Constitutive activation of endocytosis by mutation of *myoA*, the myosin I gene of *Aspergillus nidulans*. J. Biol. Chem. 273:14644–14648.
- Yan, W., and E. A. Craig. 1999. The glycine-phenylalanine-rich region determines the specificity of the yeast Hsp40 Sis1. Mol. Cell. Biol. 19:7751– 7758.
- 872. Yan, W., B. Schilke, C. Pfund, W. Walter, S. Kim, and E. A. Craig. 1998. Zuotin, a ribosome-associated DnaJ molecular chaperone. EMBO J. 17: 4809–4817.
- 873. Yang, Q., S. I. Poole, and K. A. Borkovich. 2002. A G-protein β subunit required for sexual and vegetative development and maintenance of normal Gα protein levels in *Neurospora crassa*. Eukaryot. Cell 1:378–390.
- 874. Yang, Y., P. Cheng, and Y. Liu. 2002. Regulation of the *Neurospora* circadian clock by casein kinase II. Genes Dev. 16:994–1006.
- 875. Yang, Y., P. Cheng, G. Zhi, and Y. Liu. 2001. Identification of a calcium/ calmodulin-dependent protein kinase that phosphorylates the *Neurospora* circadian clock protein FREQUENCY. J. Biol. Chem. 276:41064–41072.
- 876. Yarden, O., D. J. Ebbole, S. Freeman, R. J. Rodriguez, and M. B. Dickman. 2003. Fungal biology and agriculture: revisiting the field. Mol. Plant-Microbe Interact. 16:859–866.
- 877. Yarden, O., M. Plamann, D. J. Ebbole, and C. Yanofsky. 1992. cot-1, a gene required for hyphal elongation in *Neurospora crassa*, encodes a protein kinase. EMBO J. 11:2159–2166.
- Yarden, O., and C. Yanofsky. 1991. Chitin synthase 1 plays a major role in cell wall biogenesis in *Neurospora crassa*. Genes Dev. 5:2420–2430.
- 879. Yasuda, J., A. J. Whitmarsh, J. Cavanagh, M. Sharma, and R. J. Davis. 1999. The JIP group of mitogen-activated protein kinase scaffold proteins. Mol. Cell. Biol. 19:7245–7254.
- 880. Yasuhira, S., and A. Yasui. 2000. Alternative excision repair pathway of UV-damaged DNA in *Schizosaccharomyces pombe* operates both in nucleus and in mitochondria. J. Biol. Chem. 275:11824–11828.
- Yatzkan, E., V. Dombrádi, and O. Yarden. 1999. Detection of a protein phosphatase 2A holoenzyme in *Neurospora crassa*. Fungal Genet. Newsl. 46:32–33.
- 882. Yatzkan, E., B. Szoor, Z. Feher, V. Dombradi, and O. Yarden. 1998. Protein phosphatase 2A is involved in hyphal growth of *Neurospora crassa*. Mol. Gen. Genet. 259:523–531.
- 883. Yatzkan, E., and O. Yarden. 1999. The B regulatory subunit of protein phosphatase 2A is required for completion of macroconidiation and other developmental processes in *Neurospora crassa*. Mol. Microbiol. 31:197–209.
- 884. Yatzkan, E., and O. Yarden. 1995. Inactivation of a single-2A phosphoprotein phosphatase is lethal in *Neurospora crassa*. Curr. Genet. 28:458–466.
- Yatzkan, E., and O. Yarden. 1997. ppt-1, a *Neurospora crassa* PPT/PP5 subfamily serine/threonine protein phosphatase. Biochim. Biophys. Acta 1353:18–22.
- 886. Yazdi, M. T., A. Radford, J. N. Keen, and J. R. Woodward. 1990. Cellulase production by *Neurospora crassa*: purification and characterisation of cellulolytic enzymes. Enzyme Microb. Technol. 12:120–123.
- 887. Yonemasu, R., S. J. McCready, J. M. Murray, F. Osman, M. Takao, K. Yamamoto, A. R. Lehmann, and Y. A. 1997. Characterization of the alternative excision repair pathway of UV-damaged DNA in *Schizosaccharomyces pombe*. Nucleic Acids Res. 25:1553–1558.
- 888. Yoo, E. J., Y. H. Jin, Y. K. Jang, P. Bjerling, M. Tabish, S. H. Hong, K. Ekwall, and S. D. Park. 2000. Fission yeast *hrp1*, a chromodomain ATPase, is required for proper chromosome segregation and its overexpression interferes with chromatin condensation. Nucleic Acids Res. 28:2004–2011.
- 889. Yoon, J. H., P. M. Swiderski, B. E. Kaplan, M. Takao, A. Yasui, B. Shen, and G. P. Pfeifer. 1999. Processing of UV damage in vitro by FEN-1 proteins as part of an alternative DNA excision repair pathway. Biochemistry 38:4809–4817.
- 890. Young, J. C., and F. U. Hartl. 2000. Polypeptide release by Hsp90 involves ATP hydrolysis and is enhanced by the co-chaperone p23. EMBO J. 19: 5930–5940.
- 891. Yu, G., R. J. Deschenes, and J. S. Fassler. 1995. The essential transcription factor, Mcm1, is a downstream target of Sln1, a yeast "two-component" regulator. J. Biol. Chem. 270:8739–8743.
- 892. Yuan, G. H., Y. H. Fu, and G. A. Marzluf. 1991. nit-4, a pathway-specific regulatory gene of *Neurospora crassa*, encodes a protein with a putative binuclear zinc DNA-binding domain. Mol. Cell. Biol. 11:5735–5745.
- 893. Zapella, P. D., A. M. da Silva, J. C. da Costa Maia, and H. F. Terenzi. 1999. Serine/threonine protein phosphatases and a protein phosphatase 1 inhibitor from *Neurospora crassa*. Braz. J. Med. Biol. Res. 29:599–604.
- 894. Zatz, M., F. de Paula, A. Starling, and M. Vainzof. 2003. The 10 autosomal recessive limb-girdle muscular dystrophies. Neuromuscul. Disord. 13:532– 544.
- 895. Zeitlin, S. G., R. D. Shelby, and K. F. Sullivan. 2001. CENP-A is phosphorylated by Aurora B kinase and plays an unexpected role in completion of cytokinesis. J. Cell Biol. 155:1147–1157.
- 896. Zeke, T., E. Kokai, B. Szoor, E. Yatzkan, O. Yarden, K. Szirak, Z. Feher, P. Bagossi, P. Gergely, and V. Dombradi. 2003. Expression of protein phosphatase 1 during the asexual development of *Neurospora crassa*. Comp. Biochem. Physiol. Ser. B 134:161–170.
- 897. Zhang, Y., R. Lamm, C. Pillonel, S. Lam, and J. R. Xu. 2002. Osmoregulation and fungicide resistance: the *Neurospora crassa os-2* gene encodes a HOG1 mitogen-activated protein kinase homologue. Appl. Environ. Microbiol. 68:532–538.
- 898. Zhou, L., and G. A. Marzluf. 1999. Functional analysis of the two zinc fingers of SRE, a GATA-type factor that negatively regulates siderphore synthesis in *Neurospora crassa*. Biochemistry 38:4335–4341.
- 899. Zhou, L.-W., H. Haas, and G. A. Marzluf. 1998. Isolation and character-

ization of a new gene, *sre*, which encodes a GATA-type regulatory protein that controls iron transport in *Neurospora crassa*. Mol. Gen. Genet. **259**: 532–540.

- 900. Zhu, H., M. Nowrousian, D. Kupfer, H. V. Colot, G. Berrocal-Tito, H. Lai, D. Bell-Pedersen, B. A. Roe, J. J. Loros, and J. C. Dunlap. 2001. Analysis of expressed sequence tags from two starvation, time-of-day-specific libraries of *Neurospora crassa* reveals novel clock-controlled genes. Genetics 157:1057–1065.
- Ziman, M., J. S. Chuang, M. Tsung, S. Hamamoto, and R. Schekman. 1998. Chs6p-dependent anterograde transport of Chs3p from the chitosome to the plasma membrane in *Saccharomyces cerevisiae*. Mol. Biol. Cell 9:1565– 1576
- 902. Zimmerman, J. W., C. A. Specht, B. X. Cazares, and P. W. Robbins. 1996. The isolation of a Do1-P-Man synthase from *Ustilago maydis* that functions in *Saccharomyces cerevisiae*. Yeast 12:765–771.