

## Reconstitution of a *MEC1*-Independent Checkpoint in Yeast by Expression of a Novel Human *fork head* cDNA

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**A novel human cDNA, *CHES1* (checkpoint suppressor 1), has been isolated by suppression of the *mec1-1* checkpoint mutation in *Saccharomyces cerevisiae*. *CHES1* suppresses a number of DNA damage-activated checkpoint mutations in *S. cerevisiae*, including *mec1*, *rad9*, *rad24*, *dun1*, and *rad53*. *CHES1* suppression of sensitivity to DNA damage is specific for checkpoint-defective strains, in contrast to DNA repair-defective strains. Presence of *CHES1* but not a control vector resulted in G<sub>2</sub> delay after UV irradiation in checkpoint-defective strains, with kinetics, nuclear morphology, and cycloheximide resistance similar to those of a wild-type strain. *CHES1* can also suppress the lethality, UV sensitivity, and G<sub>2</sub> checkpoint defect of a *mec1* null mutation. In contrast to this activity, *CHES1* had no measurable effect on the replication checkpoint as assayed by hydroxyurea sensitivity of a *mec1* strain. Sequence analysis demonstrates that *CHES1* is a novel member of the *fork head*/Winged Helix family of transcription factors. Suppression of the checkpoint-defective phenotype requires a 200-amino-acid domain in the carboxy terminus of the protein which is distinct from the DNA binding site. Analysis of *CHES1* activity is most consistent with activation of an alternative *MEC1*-independent checkpoint pathway in budding yeast.**

Eukaryotes have a complex yet conserved response to DNA damage including changes in cell cycle kinetics (23) and transcriptional induction of multiple genes (17). Some of the genes responsible for these DNA damage-inducible cell cycle arrests or checkpoints have been identified (11). In at least one instance, the human checkpoint gene mutated in the disease ataxia telangiectasia, *ATM* (47), has sequence and functional homologs in many eukaryotes, including *Drosophila melanogaster* (*MEI41* [22]), *Saccharomyces cerevisiae* (*TEL1* and *MEC1/ESR1* [20, 29, 38]), and *Schizosaccharomyces pombe* (*rad3*, [50]). In contrast, other genes such as mammalian *p53*, a regulator of the damage-inducible G<sub>1</sub> checkpoint (31), do not appear to be conserved in lower eukaryotes. Germ line mutations in human checkpoint genes *ATM* and *p53* result in a predisposition to malignancy, perhaps due to the unstable nature of the genome which results when checkpoint function is absent (reviewed by Weinert and Lydall [61]).

Multiple checkpoint genes in both budding yeast (*RAD17*, *RAD24*, *RAD9*, *RAD53/MEC2/SAD1*, *MEC3*, and *POL2* [4, 23, 60, 62]) and fission yeast (*hus1*, *hus3*, *rad1*, *rad9*, *rad17*, *rad24*, *rad25*, *rad26*, *cds1*, and *chk1* [2, 3, 13, 16, 28, 37, 45, 52]) are required for normal checkpoint function. Examination of these mutant phenotypes reveals that certain gene products, e.g., Mec1 (60) and Rad3 (28), are required for response to both incomplete replication and DNA damage. Other mutant strains have specific defects in the response to DNA damage, e.g., *rad9*, *rad24* (60) of *S. cerevisiae* and *chk1* (56) and *rad27* (3) of *S. pombe*. The number of proteins required for establishment of cell cycle arrest after DNA damage suggest that a complex signal transduction pathway exists from the initial detection of DNA damage to cell cycle arrest and transcriptional induction in eukaryotic cells (7, 30).

Although substantial information has been obtained on the

control of the G<sub>1</sub> checkpoint in mammalian cells, human homologs of many of the genes responsible for the G<sub>2</sub> checkpoint in yeast have not yet been identified (reviewed by Hawley and Friend [24]). In particular, few of the gene products required for the G<sub>2</sub> arrest in mammalian cells after DNA damage have been elucidated. The human homolog of the *S. pombe rad9* checkpoint gene has been recently isolated by sequence homology, and expression of the cDNA results in partial suppression of a *rad9* mutant (34). We have developed a screen for the isolation of new human checkpoint cDNAs by demanding high-copy suppression of several checkpoint mutations in *S. cerevisiae*. In this way, human cDNA libraries can be effectively screened to obtain both homologous cDNAs and cDNAs which may function further downstream in the checkpoint pathway. This screen has resulted in the isolation of a novel member of the *fork head*/Winged Helix family of transcription factors. Expression of this clone results in reconstitution of the G<sub>2</sub> checkpoint in a *mec1*-independent manner.

### MATERIALS AND METHODS

**Strains.** The *S. cerevisiae* strains used for these experiments are described in Table 1. The source of each strain other than this laboratory is indicated.

**Yeast transformation.** Logarithmic cultures of the indicated strain were transformed by a modification of the method of Schiestl and Gietz (48) in which the DNA and 50% polyethylene glycol solution are added directly to the yeast in lithium acetate without any preincubation. Plasmid DNA from yeast was extracted by glass bead disruption (25) and transformed into *Escherichia coli* by electroporation (Bio-Rad, Hercules, Calif.).

**Libraries and DNAs.** The cDNA library and ADANS vector for screening in yeast were obtained from John Colicelli (8, 9). Adrenal and lymphocyte human libraries used to obtain longer clones were constructed by S. E. Plon and Joshua La Baer (MGH Cancer Center), respectively. Screening of the library in yeast was performed as described previously (42) by transformation of the recipient strain with purified library DNA. The transformed cultures were divided onto leucine-deficient plates and left at 23°C for 14 to 18 h, after which the plates were transferred to 30°C. Colonies were isolated after 4 to 6 days of growth. DNA was isolated as described above and retransformed into the recipient strain to determine plasmid dependence for suppression of the checkpoint phenotype. Sequencing of both strands of the *CHES1* cDNAs was performed both manually and on a Licor automated sequencer after subcloning into pBluescriptKS+.

**Survival determination after methyl methanesulfonate (MMS), hydroxyurea (HU), UV light, and X-ray exposure.** For all treatments, logarithmically growing

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TABLE 1. Yeast strains used in these experiments

Strain	Genotype	Source other than our lab
171-10-2	<i>MAT<math>\alpha</math> cdc9-8 mec1-1 leu2 ura3 ade2 ade3 trp1</i>	T. Weinert
121-1-1b	<i>MAT<math>\alpha</math> mec1-1 leu2 his3 his7 ura3</i>	T. Weinert
9085-1-7-3	<i>MAT<math>\alpha</math> leu2 his3 ura3</i>	
9085-1-8-3	<i>MAT<math>\alpha</math> cdc9-8 leu2 ura3 trp1</i>	
9085-1-10-4	<i>MAT<math>\alpha</math> cdc9-8 leu2 his3</i>	
9085-1-2-3	<i>MAT<math>\alpha</math> rad9::HIS3 leu2 ade2</i>	
10011-2-1	<i>MAT<math>\alpha</math> rad6 leu2 ura3</i>	
CRY1	<i>MAT<math>\alpha</math> can1-100 ade2 his3 leu2 trp1 ura3</i>	S. Elledge
CRY2	<i>MAT<math>\alpha</math> can1-100 ade2 his3 leu2 trp1 ura3</i>	S. Elledge
Y286	<i>MAT<math>\alpha</math> dun1-<math>\Delta</math>100::HIS3 can1-100 ade2 his3 leu2 trp1 ura3</i>	S. Elledge
Y301	<i>MAT<math>\alpha</math> rad53-21(sad1-1) can1-100 ade2 his3 leu2 trp1 ura3</i>	S. Elledge
Y438	<i>MAT<math>\alpha</math> rad9::HIS3 can1-100 ade2 his3 leu2 trp1 ura3</i>	S. Elledge
Y669	<i>MAT<math>\alpha</math> mec1::HIS3 can1-100 ade2 his3 leu2 trp1 ura3 + PWJ81 (containing the MEC1 gene and URA3 marker)</i>	S. Elledge
EMY62	<i>MAT<math>\alpha</math> rad52::URA3 his3 leu2 trp1 ura3</i>	D. Botstein
XKS255-1A	<i>MAT<math>\alpha</math> rad24-1 his3 leu2 lys2 trp1 ura3</i>	J. Game

cultures were obtained by inoculation of 10 to 20 ml of leucine-deficient medium with an aliquot of a stationary-phase culture and grown overnight. The next morning, cultures containing between 1 million and 5 million cells/ml were used for the experiments.

(i) **MMS.** Cultures were incubated at indicated times in the presence and absence of media containing 0.1% MMS (Sigma, St. Louis, Mo.) at 30°C. Following the incubation, cells were briefly sonicated and plated onto leucine-deficient plates, colonies were counted in 48 to 72 h, and percent survival was determined.

(ii) **UV irradiation.** Cells were diluted in leucine-deficient medium at different concentrations, briefly sonicated, and then plated. Plates were irradiated at indicated doses, using a Spectronics (Westbury, N.Y.) Spectrolinker UV cross-linker. Plates were incubated at 30°C for 48 to 72 h, colonies were counted, and percent survival compared to the unirradiated control was determined.

(iii) **X-ray irradiation.** Cultures were prepared as described for UV irradiation. Plates were exposed to various doses of radiation by using a Machlett OEG-60 X-ray tube operated at 50 kV and 20 mA, delivering a dose of 1 Gy/s.

(iv) **HU.** HU treatment was as described above for determination of MMS sensitivity, with 0.2 M HU (Sigma) added to the medium. In addition, transformants containing the indicated plasmids were streaked out onto solid medium containing 10 mM HU, and colony formation was assessed in 48 to 72 h.

**Doubling time.** Ten milliliters of medium was inoculated with 10 to 100  $\mu$ l of saturated culture, and cells were grown overnight until early log phase (approximately  $10^6$  cells/ml). Over the next 6 h, aliquots were removed, iced, sonicated, and counted on a Coulter (Miami, Fla.) Multisizer II Particle Counter. Linear regression was performed on the  $\log_{10}$  of cell number versus time to calculate doubling time and  $r^2$  value for each data set.

**$\alpha$ -Factor arrest.** Cultures were grown overnight in leucine-deficient pH 4.0 medium. After pelleting, cells were resuspended in medium with 2  $\mu$ M  $\alpha$ -factor and incubated until >95% of the cells were arrested as single buds by visual inspection under light microscopy. Cells were pelleted and resuspended in medium containing 20  $\mu$ g of pronase E per ml. At indicated times, aliquots were removed, fixed in 70% ethanol for 1 h, and stored. Propidium iodide staining was performed (27), and samples were analyzed on a Becton Dickinson (Franklin Lakes, N.J.) FACScan.

**Assay of  $\beta$ -galactosidase activity.** Transcriptional induction by *CHES1* in wild-type and *rad9* and *dun1* mutant strains was assayed by using a reporter construct containing the *RNR2* (ribonucleotide reductase 2) promoter upstream of  $\beta$ -galactosidase (pSE788) (12).  $\beta$ -Galactosidase levels were measured quantitatively in a chromogenic assay using yeast extracts and *o*-nitrophenyl- $\beta$ -D-galactoside (ONPG) as a substrate (44). The activity was then normalized to the amount of protein assayed in each lysate by Bio-Rad protein assay.  $\beta$ -Galactosidase activity was measured for wild-type (CRY1), *rad9* (Y438), and *dun1* (Y286) strains containing the pSE788 reporter and either ADANS or *CHES1* before and after exposure to 0.1% MMS for 4 h in liquid media.

**Suppression of a *mec1* $\Delta$  null strain.** A *mec1* $\Delta$  strain (Y669) containing the wild-type *MEC1* gene on a plasmid was transformed with ADANS, *CHES1*, or a longer cDNA containing the DNA binding domain [*CHES1*(db)]. Transformants were selected on leucine-deficient plates and then transferred to plates contain-

ing 5-fluoro-orotic acid (5-FOA; PCR Incorporated, Gainesville, Fla.) at 0.1% to select against the *URA3* plasmid containing the yeast *MEC1* gene. The loss of the *MEC1*-containing plasmid (in the colonies grown on 5-FOA) was verified by loss of a *MEC1*-specific PCR product by using two sets of primers designed in the 5' (GCTTTCCAGCTGCTTATATCGATC and GAAC TAGCAGCACTAGAA AATGCCGA) and 3' (TCTTTCCATGATTGCGCAAGAT and TGGGCTTA AGGAAGTTCGATACG) regions of the *MEC1* gene. *mec1* $\Delta$  clones containing *CHES1* were restreaked from a 5-FOA plate onto leucine-deficient medium and used for further assays.

**Effect of DNA damage on the release of G<sub>2</sub>-arrested cells.** Logarithmically growing cultures in leucine-deficient medium were transferred to 50% rich media (YM-1) for 3 h. Ten milliliters of this culture was pelleted by centrifugation at 2,000 rpm for 3 min and resuspended in 5 ml of rich medium containing nocodazole (10  $\mu$ g/ml; Sigma) in 1% dimethyl sulfoxide. After 2.5 to 3.0 h of incubation at 30°C, approximately 87 to 96% cells were arrested with a large bud and contained an undivided nucleus with DNA content in the neck of the large-budded cells, as monitored by staining with 4',6'-diamidino-2-phenylindole (DAPI; Sigma) as described previously (1). The cells were pelleted once again and plated in duplicate onto nocodazole (10  $\mu$ g/ml)-containing agar plates. One plate was UV irradiated at a dose of 70 J/m<sup>2</sup>, while the second plate served as a control. The cells were then immediately harvested from the plates and washed twice with medium to remove nocodazole and to recover cells into liquid medium. Samples were collected at 20-min intervals during recovery, briefly sonicated, fixed with 70% ethanol for 1 h, and resuspended in 10 mM Tris-1 mM EDTA buffer. Samples were counted for the fraction of large-budded cells as previously described (49) and for nuclear morphology after DAPI staining (1).

**Inhibition of protein synthesis.** Nocodazole-induced G<sub>2</sub>-arrested cells (described above) were exposed to cycloheximide (10  $\mu$ g/ml; Sigma) beginning 15 min prior and then continuously, during and after UV exposure as previously described (58). The inhibition of protein synthesis after cycloheximide treatment was determined by incorporation of <sup>35</sup>S-labeled methionine (ICN Radiochemicals, Costa Mesa, Calif.) into acid-precipitable materials (21, 34) and was found to be greater than 97% suppressed. Arrest after DNA damage was monitored by following the percentage of cells with a large-budded morphology with no visible septum between the bud and mother cell. In control experiments, we detected a decrease in large-budded cells after release from nocodazole in cells exposed to cycloheximide. Our results of monitor of cells in this fashion contrast with the inhibition of complete cytokinesis (as measured by cell number after treatment with glusulase) by cycloheximide exposure as reported by Burke and Church (6).

**Nucleotide sequence accession number.** The GenBank accession number for the *CHES1* cDNA sequence is U68723.

## RESULTS

**Isolation of a human cDNA which suppresses the *mec1-1* mutation.** The genetic screen to obtain potential human checkpoint cDNAs was based on previous observations (59, 62) that a temperature-sensitive allele of DNA ligase (*cdc9-8*) had a decrease in maximum permissive temperature from 30 to 23°C when coupled with mutations in either the *RAD9* or *MEC1* checkpoint gene. Therefore, suppression of either the *mec1* or *rad9* mutation by a human cDNA in the double-mutant strain should allow growth at 30°C. A human cDNA library derived from the U118 glioblastoma cell line cloned in the 2 $\mu$ m yeast expression vector ADANS was used to transform the double-mutant strains. The vector from this library, ADANS, contains a constitutive *ADH* promoter upstream of the human cDNA. Baseline experiments with the ADANS vector demonstrated growth at 30°C of approximately one colony per 10,000 and 3,000 transformants for *rad9* $\Delta$  *cdc9-8* and *mec1-1* *cdc9-8* strains, respectively.

Preliminary screening using this strategy resulted in the isolation of human *CDC34* (42), a cDNA which suppressed *mec1-1* *cdc9-8* for growth at 30°C but did not suppress any other *mec1-1* phenotype. In total, screening of approximately 300,000 library *rad9* $\Delta$  *cdc9-8* and 400,000 *mec1-1* *cdc9-8* transformant strains for growth at 30°C resulted in a total of 45 and 65 colonies, respectively. Only three of these transformants (transformants 6 [*CDC34*], 42, and 23) demonstrated plasmid-dependent growth at 30°C. Further analysis revealed a single clone, transformant 23 (*tx23*) of the *mec1-1* *cdc9-8* strain, which conferred plasmid-dependent growth at 30°C and suppressed other checkpoint phenotypes (as described below). As demonstrated in Fig. 1A, *tx23* confers significant colony growth

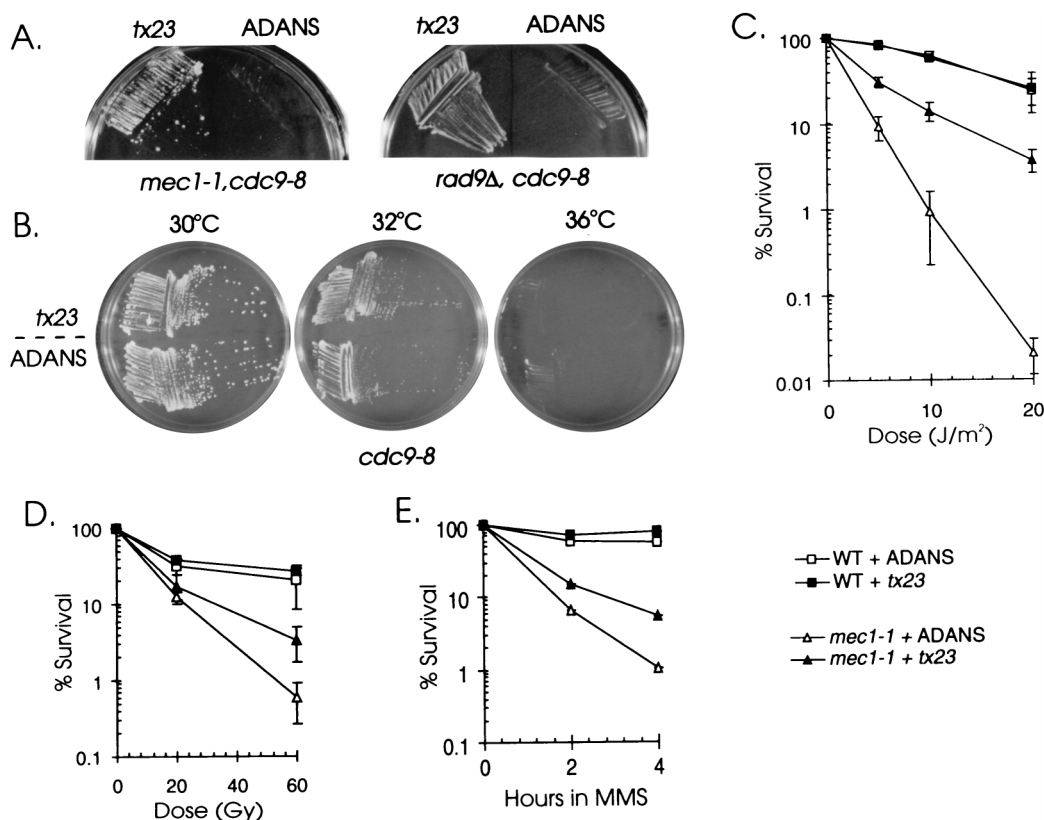


FIG. 1. Isolation of a human cDNA which suppresses multiple *mec1-1* phenotypes. (A) *tx23* rescues the temperature-sensitive lethality of *mec1-1 cdc9-8* (171-10-2) and *rad9Δ cdc9-8* (9085-1-8-3) double-mutant strains for growth at 30°C. (B) *tx23* does not rescue a *cdc9-8* (9085-1-10-4) strain for growth at 36°C. (C to E) Effects of UV light (C), X rays (D), and MMS (E) on the survival of wild-type (WT; 9085-1-7-3; □) and *mec1-1* (121-1-1b; △) yeast strains in the presence of an ADANS control vector (open symbols) or *tx23* cDNA (filled symbols). Graphical data shown in all figures are the averages and standard errors of means of two to four experiments. In some cases, e.g., panel E, the standard error of the mean is smaller than the icon size.

at 30°C for a *mec1 cdc9-8* strain compared to a library vector (ADANS) control. A similar suppression of the *rad9 cdc9-8* strain for growth at 30°C was also found (Fig. 1A). The cDNA did not directly suppress the *cdc9-8* mutation, as there was no change in the temperature-sensitive profile of a *MEC*<sup>+</sup> *RAD*<sup>+</sup> *cdc9-8* strain containing *tx23* (Fig. 1B).

Secondary assays for suppression of other DNA damage checkpoint phenotypes were then carried out. A common feature of strains containing mutations in DNA damage checkpoint genes is sensitivity to radiation-induced damage. In all screens of sensitivity to damage, exponentially growing fresh cultures in leucine-deficient medium were used. Transformation of a *mec1* strain with *tx23* resulted in 185-fold increase in survival after exposure to 20 J of UV radiation per m<sup>2</sup> (Fig. 1C). Similarly, survival after exposure to 60 Gy of ionizing radiation of both *mec1-1* (Fig. 1D) and *rad9Δ* (data not shown) strains was increased 10-fold by *tx23* compared with ADANS vector controls. There was no effect on survival of a wild-type strain containing *tx23* to either UV or ionizing radiation.

*mec1* mutant strains have also been shown to exhibit defects in regulation of S phase after MMS and HU exposure. While treatment of *mec1* strains with MMS results in a S-phase arrest due to DNA damage (41), HU treatment is thought to trigger the S-phase checkpoint due to detection of stalled replication forks (60). Exposure of a *mec1-1* strain containing *tx23* to either HU or MMS revealed disparate results. *tx23* expression resulted in a fivefold increase in survival of *mec1-1* strains to MMS treatment (Fig. 1E). This difference was seen in multiple

independent experiments. However, expression of *tx23* in this same strain had no measurable effect on HU sensitivity assayed both by survival of an exponential culture after 3 h of exposure to 0.2 M HU (0.17% for ADANS and 0.065% for *tx23* [Fig. 2A]) and by growth of wild-type and *mec1-1* mutant strains on plates containing 10 mM HU (Fig. 2B).

These experiments demonstrate that *tx23* is able to suppress multiple phenotypes in both the *mec1-1* and *rad9Δ* checkpoint-deficient strains. Given this activity, we named this cDNA *CHES1* (for checkpoint suppressor 1). To ascertain whether this effect is specific for the A364A strain background used in the *mec1-1* and *rad9Δ* strains described above, we tested the effect of *CHES1* on the same *rad9::HIS3* deletion allele in strain Y438, which is in a W303 strain background. *CHES1* expression in this background resulted in a 30-fold increase in survival of a *rad9* strain containing *CHES1* compared with an ADANS control when exposed to 20 J of UV radiation per m<sup>2</sup> (Fig. 3A). The result of these secondary screens demonstrate that *CHES1* suppression of *mec1-1* and *rad9Δ* strains is specific for the response of yeast to DNA damage and has significant activity in several strain backgrounds.

***CHES1* suppresses multiple checkpoint mutations.** Additional radiation-sensitive strains of *S. cerevisiae* were tested for suppression of UV sensitivity by *CHES1*. *RAD24* is in the same epistasis group as *RAD9*. However, Lydall and Weinert (36) presented data suggesting that *RAD24* plays a distinct role early in processing of DNA damage. Expression of *CHES1* also suppresses a *rad24-1* mutant strain when exposed to UV dam-

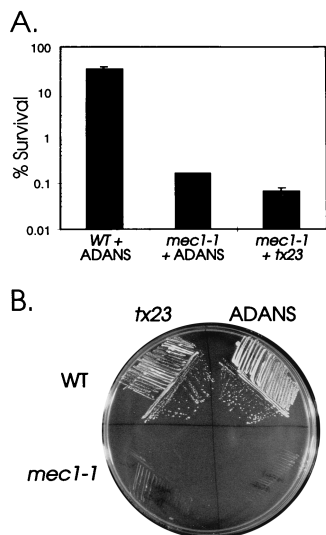


FIG. 2. *tx23* expression does not alter sensitivity of a *mec1-1* strain to HU. (A) Survival of wild-type (WT; 9085-1-7-3) and *mec1-1* (121-1-1b) strains containing ADANS control or *tx23* to 3 h of exposure to 0.2 M HU. (B) Survival upon exposure of transformants of wild-type and *mec1-1* strains to solid medium containing 10 mM HU.

age to a reproducible but lesser extent (fourfold) than it suppresses *rad9Δ* (Fig. 3B).

The protein encoded by *RAD53* (also isolated as *SPK1* [54], *SAD1* [4], and *MEC2* [60]) is an essential protein kinase which is required for both cell cycle arrest and transcriptional induction after DNA damage. Requirement for Rad53 activity has been placed downstream of Mec1 due to *RAD53* suppression of a *mec1* null allele and *MEC1*-dependent phosphorylation of Rad53 protein (46). There was a reproducible threefold suppression of the moderate UV sensitivity of a *rad53-21* strain (*sad1-1* allele) by expression of *CHES1* cDNA (Fig. 3C).

*DUN1* encodes another protein kinase which is required for the transcriptional induction program after DNA damage (64). Sensitivity of *dun1Δ* strains to UV radiation is much greater than that of *rad53-21* strains, and this sensitivity was suppressed over 40-fold in multiple experiments by the presence of *CHES1* (Fig. 3D). A *dun1Δ* strain containing *CHES1* has resistance to UV radiation of 20 J/m<sup>2</sup> equivalent to that of an isogenic wild-type strain containing the ADANS control vector.

In contrast to the suppression of multiple checkpoint mutant strains, expression of *CHES1* did not suppress the sensitivity of strains mutant in two other radiation-sensitive epistasis groups required for DNA repair (43). There was no suppression of strains mutant in either *rad6* (Fig. 3E) or *rad52* (data not shown) for sensitivity to UV radiation.

The result of these secondary assays are summarized in Table 2. We have detected significant suppression of multiple aspects of the checkpoint response to DNA damage but not replication delays or repair defects by expression of *CHES1* in *S. cerevisiae*. This suppression results in increased resistance of the mutant yeast to DNA damage but does not change the response of wild-type yeast. The highest degree of suppression by *CHES1* is for *mec1-1*, *rad9Δ*, and *dun1Δ* strains exposed to UV radiation.

***CHES1* does not act by induction of DNA damage-responsive genes or constitutive effects on cell cycle kinetics.** We next explored the mechanism responsible for the increased survival of these mutant strains when *CHES1* is present. We considered three possibilities: (i) expression of *CHES1* results in constitutive induction of the DNA damage response genes, (ii) *CHES1* has a nonspecific effect on cell cycle kinetics, for example, slowing of the cycle, which makes the yeast strains more resistant; and (iii) *CHES1* results in reconstitution of the normal cell cycle arrest which accompanies DNA damage.

Given the finding that *CHES1* had significant suppression of a *dun1* mutant strain, we examined transcriptional induction after DNA damage. A reporter construct (pSE788) containing an *RNR2* promoter upstream of β-galactosidase was intro-

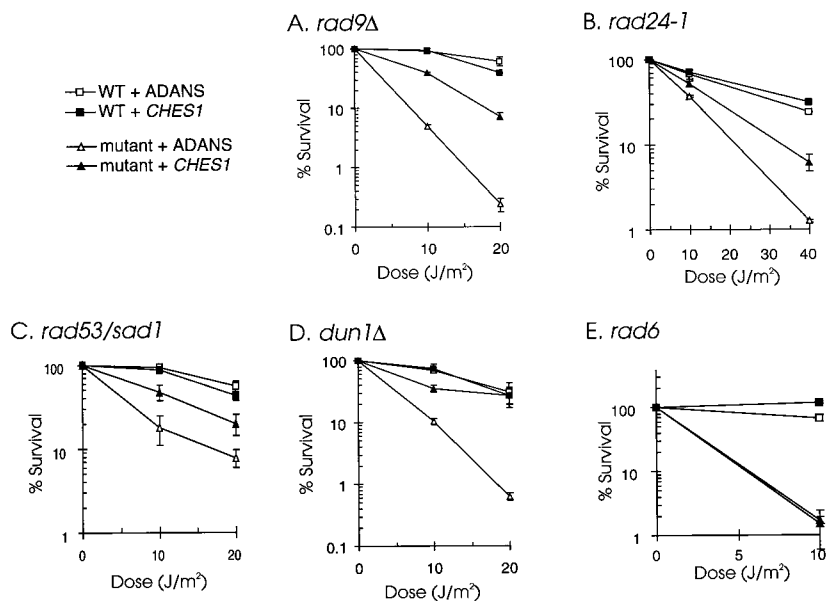


FIG. 3. *CHES1* suppresses the UV sensitivity of multiple-checkpoint-deficient but not repair-deficient strains. In all panels, the wild-type (WT) strain (CRY1 except for panel D, which is CRY2 to maintain the same mating type) is depicted by □; the mutant strains are indicated by △. (A) *rad9Δ* (Y438); (B) *rad24* (XRS255-1A); (C) *rad53-21* (Y301); (D) *dun1Δ* (Y286); (E) *rad6* (10011-2-1). Open symbols, ADANS control; closed symbols, *CHES1* cDNA.

TABLE 2. Summary of *CHES1* suppression of mutant phenotypes<sup>a</sup>

Relevant genotype	<i>cdc9-8</i> checkpoint	Ionizing radiation	UV	MMS	HU
<i>mec1-1</i>	+++	++	+++	+	-
<i>rad9Δ</i>	+++	++	+++		
<i>dun1Δ</i>			+++		
<i>rad24-1</i>			++		
<i>rad53-21 (sad1)</i>			+		
<i>rad52Δ</i>			-		
<i>rad6</i>			-		

<sup>a</sup> The symbols refer to the average degree of suppression of lethality compared with an ADANS control of the treatments indicated from analysis of between two and four independent experiments. -, no detectable suppression; +, 3- to 5-fold; ++, >5-fold; +++, >40-fold.

duced into these strains. This construct had previously been shown to be responsive to DNA damage in yeast (12), and this induction is lost in *dun1Δ* strains (64). Quantitative  $\beta$ -galactosidase activity was determined for wild-type (CRY1), *rad9Δ* (Y438), and *dun1Δ* (Y286) isogenic strains containing the reporter and either the ADANS or *CHES1* plasmid before and after exposure to MMS (Fig. 4A) and UV radiation (data not shown). There was no significant  $\beta$ -galactosidase induction of the exponential cultures prior to damage with either ADANS or *CHES1*. After damage, there was a small increase in induction of a wild-type strain carrying *CHES1* as opposed to ADANS but no increase in induction of either the *rad9Δ* or *dun1Δ* strain. Thus, rescue of the *rad9* and *dun1* mutations does not appear to be due to induction of this DNA damage-sensitive promoter either constitutively or after DNA damage.

To explore the effect of *CHES1* on cell cycle kinetics, we carried out quantitative assays to determine the doubling time of wild-type, *rad9Δ*, and *mec1-1* strains containing either ADANS or *CHES1*. In multiple assays, there were no significant differences between the doubling times of exponential cultures (Table 3). Second, we saw no differences in the profile of propidium iodide-stained cultures of exponential cells or cells after release from arrest in G<sub>1</sub> by  $\alpha$ -factor (Fig. 4B and C). The *rad9Δ* cultures exited G<sub>1</sub> and progressed through S phase to G<sub>2</sub> with the same kinetics whether containing *CHES1* (Fig. 4C) or ADANS (Fig. 4B). Third, wild-type *mec1-1*, *rad9Δ*, and *dun1Δ* strains were synchronized in G<sub>2</sub> by nocodazole, and transit through the cycle upon release of G<sub>2</sub> synchronization was monitored by following the percentage of large-budded yeast with the nucleus in the neck after staining with DAPI. As shown in Fig. 5, undamaged cells of wild-type, *mec1-1*, *rad9Δ*, or *dun1Δ* strains containing either ADANS or *CHES1* when released from nocodazole block had superimposable exit from G<sub>2</sub>. To summarize, on all three measures, exponential growth, release from G<sub>1</sub> arrest by  $\alpha$ -factor, and release from G<sub>2</sub> arrest by nocodazole, there was no significant difference in cultures containing ADANS or *CHES1* compared with the large differences seen in survival after DNA damage in these strains.

***CHES1* restores the G<sub>2</sub> delay of *mec1-1* and *rad9Δ* strains damaged by UV irradiation.** Detailed assay of the damage-inducible checkpoint was studied by synchronization of cells in G<sub>2</sub> by addition of nocodazole (Fig. 5). These synchronized cells were then exposed to UV radiation, and the G<sub>2</sub> block was released by removal of nocodazole. Transit through the cycle was monitored by morphology and staining with DAPI. As expected (Fig. 5A), wild-type strains show significant delay after UV radiation with either ADANS or *CHES1*. In contrast, a *mec1-1* mutant strain (Fig. 5B) demonstrates complete loss of G<sub>2</sub> delay after UV radiation with the ADANS control vector.

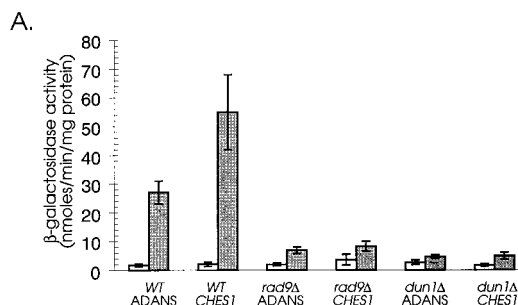
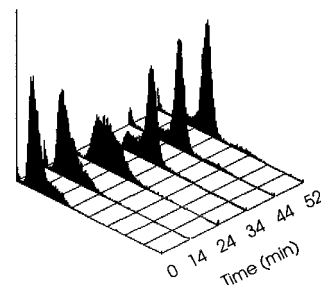
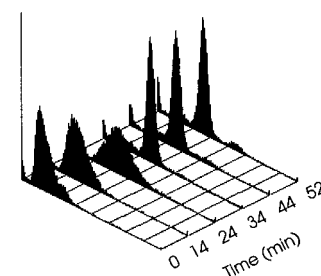
B. *rad9Δ* + ADANSC. *rad9Δ* + *CHES1*

FIG. 4. Lack of transcriptional induction or cell cycle slowing by *CHES1*. (A) A reporter construct containing the *RNR2* promoter upstream of  $\beta$ -galactosidase was introduced in *rad9Δ* and *dun1Δ* strains.  $\beta$ -Galactosidase activity was measured for wild-type (WT; CRY1), *rad9Δ* (Y438), and *dun1Δ* (Y286) strains containing the reporter and either ADANS or *CHES1* cDNA after exposure to control medium (open bars) or medium containing 0.1% MMS (shaded bars) for 4 h. (B and C) An exponential culture of a *rad9Δ* (Y438) strain containing ADANS (B) or *CHES1* (C) was arrested with  $\alpha$ -factor and then released into medium without  $\alpha$ -factor. Samples were taken, fixed, stained with propidium iodide, and analyzed as described in the text.

Introduction of *CHES1* results in reconstitution of the G<sub>2</sub> delay, yielding a profile of G<sub>2</sub> arrest similar to that of the wild type. The same experiment was performed with wild-type, *rad9Δ*, and *dun1Δ* strains in a W303 background. *CHES1* also reconstituted the G<sub>2</sub> delay in *rad9Δ* strains to wild-type levels

TABLE 3. Doubling times of strains containing the *CHES1* cDNA or ADANS control

Strain	Plasmid	Doubling time (min) <sup>a</sup>
CRY1 (wild type)	ADANS	105
	<i>CHES1</i>	103
121-1-1b ( <i>mec1-1</i> )	ADANS	126
	<i>CHES1</i>	128
Y438 ( <i>rad9Δ</i> )	ADANS	101
	<i>CHES1</i>	109

<sup>a</sup> Based on experiments with  $r^2 > 0.975$ .

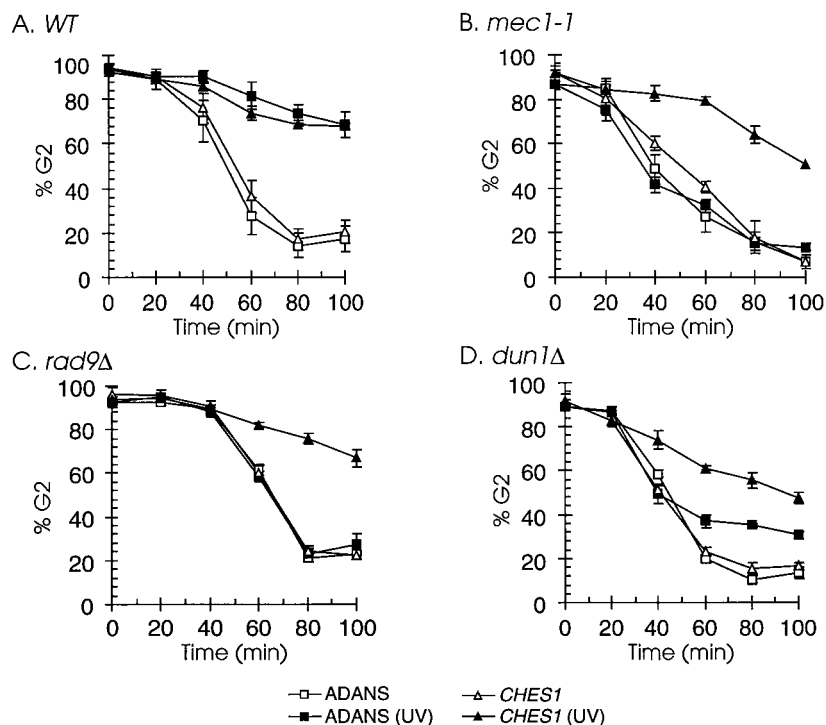


FIG. 5. Restoration of the DNA damage-induced  $G_2$  checkpoint by *CHES1*. Nocodazole-arrested cultures were plated, UV irradiated, and then released into medium. Timed samples were fixed and analyzed as described in the text. Each panel represent analysis of one strain. (A) Wild-type (WT; *CRY1*); (B) *mec1-1* (121-1-1b); (C) *rad9* $\Delta$  (Y438); (D) *dun1* $\Delta$  (Y286).  $\square$ , ADANS control;  $\triangle$ , *CHES1* cDNA; filled symbols, UV-irradiated cells; open symbols, unirradiated controls.

(Fig. 5C) after UV irradiation. The *dun1* $\Delta$  strain containing the ADANS control had an intermediate loss of checkpoint function, with only 35% of the cells remaining in  $G_2$  80 min after damage (Fig. 5D). Addition of *CHES1* resulted in an increase in  $G_2$  delay in the *dun1* $\Delta$  strain to near wild-type levels. Thus, our analysis demonstrates no significant change in cell cycle control of undamaged cultures but substantial normalization of the  $G_2$  checkpoint by *CHES1* upon damage of cells deficient in *Mec1*, *Rad9*, or *Dun1* function.

***CHES1* encodes a truncated *fork head*/Winged Helix cDNA.** Analysis of the *CHES1* cDNA revealed a 1.6-kb message (Fig. 6A) with 600 bp of open reading frame placed in frame with the start site of translation and first 13 amino acids of the yeast *ADH* gene found in the ADANS vector. The remainder of the message was a 1.1-kb 3' untranslated region (UTR) containing a CA dinucleotide repeat (see below). The cDNA was primed off of a short internal run of poly(A) in the 3' UTR sequence. Analysis of the putative protein encoded by *CHES1* revealed a small but significant region of homology to HTLF (human T-cell leukemia enhancer factor), a member of the *fork head*/Winged Helix family (33). This possibility was further explored by obtaining longer cDNAs encoding an entire open reading frame (Fig. 6A) from human adrenal and T-lymphocyte cDNA libraries. This longer cDNA encodes a protein of 491 amino acids with predicted molecular mass of 54 kDa (Fig. 6B and C). Truncation of the cDNA to produce the original *tx23* clone occurred due to the presence of an internal *NotI* site in the cDNA.

The chromosomal location of *CHES1* was determined by use of the CA repeat in the 3' UTR of the message. This repeat was found to be highly polymorphic in human DNA. Initial mapping determined linkage of this marker (named *CCCI*) with 14q32, which was further refined to the region between

14q24.3 and 14q31 (15). Flanking markers include D14S67 and AFM343.

Protein sequence analysis (5) with the open reading frame encoded by the longer *CHES1* cDNA revealed a high degree of homology to the DNA binding region of *fork head*/Winged Helix proteins. However, the DNA binding domain is not found in the original *CHES1* cDNA, *tx23* (Fig. 6B and C). An extensive analysis of 72 members of this family by both phylogenetic (14) and multiple alignment (53) methods centered on the shared DNA binding domain demonstrated that *CHES1* belonged to a subfamily of *fork head*/Winged Helix proteins including HTLF, rodent WHN (40), and FKHR (18, 51). The alignment with HTLF, the closest relative of *CHES1*, is shown in Fig. 6C. Overall, there are 51% identical and 69% conserved residues between HTLF and *CHES1*. In addition to the highly conserved DNA binding domain, there are significant regions of homology between these two proteins both upstream and downstream of this motif, including portions encoded by the original truncated *tx23* cDNA.

**Reconstitution of the  $G_2$  checkpoint does not require new protein synthesis.** Weinert and Hartwell previously demonstrated that the *RAD9*-dependent  $G_2$  checkpoint did not require new protein synthesis (58). Although *CHES1* is a member of a family of transcription factors, the lack of the DNA binding domain in the truncated *CHES1* cDNA suggested that the *CHES1*-induced checkpoint would also be independent of new protein synthesis. UV irradiation of nocodazole-synchronized cultures was repeated with cycloheximide added before, during, and after irradiation. As shown in Fig. 7, *CHES1* reconstituted a  $G_2$  delay in a *mec1-1* strain in the presence of cycloheximide with similar kinetics as in cells irradiated in the absence of cycloheximide.



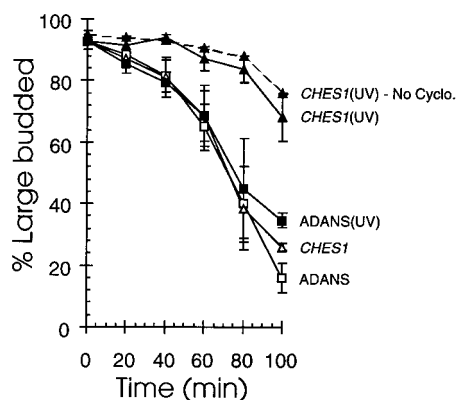


FIG. 7. Restoration of the *CHES1*-mediated  $G_2$  delay of a *mec1-1* strain does not require protein synthesis. Nocodazole-arrested  $G_2$  cells containing either ADANS ( $\square$ ) or *CHES1* cDNA ( $\Delta$ ) were exposed to the protein synthesis inhibitor cycloheximide (Cyclo.; 10  $\mu$ g/ml) for 15 min before, during and after UV exposure (filled symbols). Open symbols, unirradiated controls. The broken line represents data from irradiated samples in the absence of cycloheximide.

***CHES1* suppresses the lethality of a *mec1* null mutation.** In addition to the multiple checkpoint phenotypes of a *mec1-1* strain, the *MEC1/ESR1* gene was also identified as being essential for growth in the absence of DNA damage (29). The essential requirement for *MEC1* has not been well characterized. We expressed both the *CHES1* and *CHES1(db)* cDNAs in a *mec1 $\Delta$*  strain which contained the wild-type *MEC1* gene on a *URA3*-marked plasmid. Subsequently these transformants were plated on 5-FOA to select against the presence of the *URA3* plasmid. Expression of *CHES1* allowed numerous 5-FOA colonies to grow, in contrast to the lack of growth of colonies containing the ADANS control vector (Fig. 8A). Colonies containing the longer cDNA were visible about 1 day earlier than those containing the truncated *CHES1* cDNA. *CHES1*-containing colonies grown on 5-FOA were verified for the loss of the *MEC1*-bearing plasmid by using a robust PCR assay specific for the 5' and 3' ends of the *MEC1* gene. Of five *mec1 $\Delta$*  *CHES1* colonies picked from the 5-FOA plate, we were unable to amplify either *MEC1* fragment (data not shown), confirming loss of the *MEC1* plasmid. The *mec1 $\Delta$*  cells containing *CHES1* grow well with colonies first visible about 1 day later than the *mec1 $\Delta$*  cells containing the *MEC1* gene. Thus, expression of *CHES1* is sufficient to overcome the essential requirement for *MEC1* function during growth. *mec1 $\Delta$*  null mutants expressing *CHES1* demonstrated survival after UV irradiation (Fig. 8B) and a  $G_2$  checkpoint response (Fig. 8C) similar to those of *mec1-1* cells expressing *CHES1*. This result clearly demonstrates that a checkpoint can be activated by *CHES1* in the complete absence of *MEC1* function.

## DISCUSSION

**Isolation of a human suppressor of multiple checkpoint mutations.** In this paper, we report the isolation of a human cDNA, *CHES1*, which can substitute for several genes known to regulate the cell cycle response to DNA damage in yeast. The suppression of these mutant strains results in at least 10-fold increase in survival to several different forms of DNA damage. The greatest suppression (between 40- and 185-fold) is the UV sensitivity of *mec1-1*, *rad9 $\Delta$* , and *dun1 $\Delta$*  strains. There is no suppression of UV sensitivity of the repair-deficient strains *rad6* and *rad52*. This suppression of the checkpoint strains is accompanied by reconstitution of a wild-type

$G_2$  arrest after DNA damage in yeast strains deficient in this arrest due to mutations in the *MEC1*, *RAD9*, *RAD24*, *RAD53*, and *DUN1* checkpoint genes. Despite mutation in the known  $G_2$  checkpoint pathway, the characteristics of this  $G_2$  arrest are similar to those of the wild type in time course, nuclear morphology, and cycloheximide resistance.

*mec1-1* strains remain very sensitive to HU upon expression of *CHES1*. HU sensitivity is reported to be due to loss of the checkpoint which senses incomplete replication (39). Although we were not able to detect suppression of HU sensitivity of a *mec1-1* strain, the truncated *CHES1* and longer *CHES1(db)* cDNAs are able to suppress the lethality of a *mec1 $\Delta$*  strain. This result suggests that the essential requirement for *MEC1* may not be due the ability of Mec1 protein to sense replication delays.

In contrast to the ability of *CHES1* to reconstitute the  $G_2$  checkpoint, expression of this clone does not restore the loss of transcriptional induction of an *RNR2* promoter after DNA damage in these mutants. We have found that in addition to a defect in transcriptional induction, the *dun1 $\Delta$*  mutation results

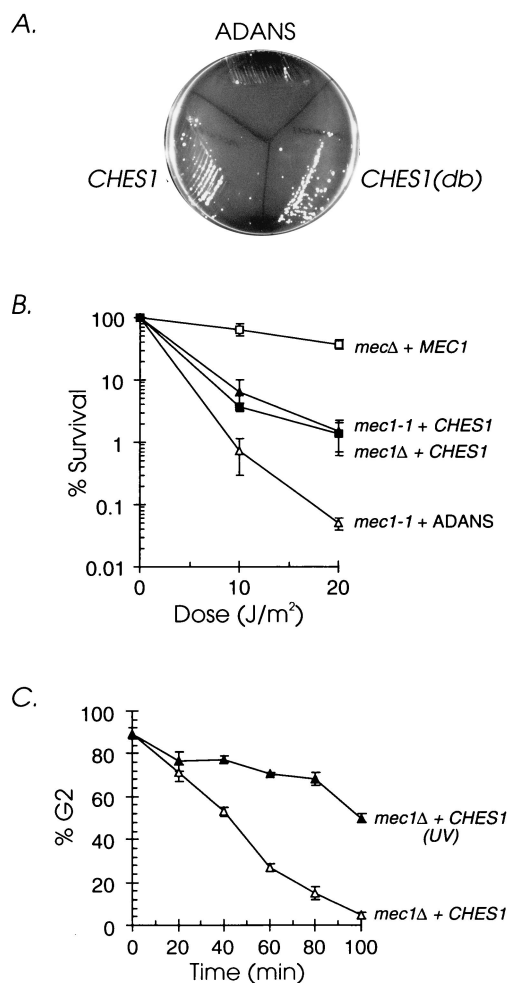


FIG. 8. *CHES1* suppresses a *mec1 $\Delta$*  null mutant. (A) Expression of *CHES1* and *CHES1(db)* but not ADANS suppressed the lethality of a *mec1 $\Delta$*  (Y669) strain, allowing growth of colonies on plates containing 0.1% 5-FOA, which selects against the *URA3*-marked plasmid containing the *S. cerevisiae* *MEC1* gene. (B) *CHES1* expression results in increased survival after UV irradiation.  $\square$ , *mec1 $\Delta$*  strain (Y669);  $\Delta$ , *mec1-1* strain (121-1-1b); open symbols, ADANS; filled symbols, *CHES1*. (C) *CHES1* restores a  $G_2$  checkpoint to a *mec1 $\Delta$*  strain after UV irradiation.



in a partial G<sub>2</sub> checkpoint defect after UV irradiation. This checkpoint defect of a *dun1Δ* strain was not previously detected by Zhou and Elledge in assays using similar irradiation doses and strains (64). Restoration of this delay may explain the suppression of lethality in a *dun1Δ* strain by *CHES1* despite the lack of restoration of transcriptional induction.

***CHES1* encodes a novel fork head/Winged Helix protein.** Analysis of the *CHES1* cDNA reveals that this message encodes a novel member of the *fork head*/Winged Helix family of transcription factors and not a protein homologous to a known checkpoint gene. Similarly, Davey and Beach (10) identified a novel human cDNA, *RACH2*, in a screen for complementation of the *S. pombe rad1-1* mutation. We have no evidence that transcriptional induction of new genes is required for restoration of the checkpoint defect. First, only 200 amino acids in the carboxy terminus not including the conserved DNA binding motif are required for restoration of the checkpoint; second, the *RNR2* damage-inducible promoter is not activated by *CHES1*; and third, the G<sub>2</sub> checkpoint is restored in the presence of cycloheximide. Inclusion of a longer cDNA containing the DNA binding domain resulted in increased suppression of the checkpoint defect in *rad9* and *rad24* strains to wild-type levels (data not shown). However, this increased suppression may be a combination of the cell cycle slowing observed in cultures containing the longer *CHES1* cDNA (40a) and restoration of the DNA damage-induced G<sub>2</sub> checkpoint or the result of other factors such as increased stability of the protein.

The *fork head*/Winged Helix family was originally identified through studies of the *Drosophila* region-specific homeotic mutant *fork head* (57) and in studies of transcription factors which play a role in mammalian liver-specific gene (*HNF-3A*) expression (32). Members of this large family share a DNA binding domain which encodes a Winged Helix motif (57) but may have little homology outside this region. Although *S. cerevisiae* contains several genes with conserved *fork head*/Winged Helix motifs, there are no predicted proteins with significant homology to the carboxy terminus of *CHES1*. *CHES1* also does not contain the *fork head*-associated domain first identified by short regions of homology outside the DNA binding domain in a subset of *fork head*-encoded proteins and many proteins involved in signal transduction, including Rad53 and Dun1 (26).

Sequence alignment and phylogenetic analysis of the *fork head*/Winged Helix family reveal that *CHES1* is most closely related to the mammalian genes *HTLF* (33), *FKHR* (18, 51), and murine *WHN* (40). *HTLF* shows significant homology to *CHES1* in the portion of the protein-coding region contained in the truncated *CHES1* cDNA. *HTLF* was first isolated in a screen for factors which bind to the long terminal repeat of the retrovirus human T-cell leukemia virus type 1. Further studies are needed to determine if *CHES1* or *HTLF* plays a role in the induction of viruses upon DNA damage in mammalian cells.

**Evidence for an alternative *MEC1*-independent checkpoint pathway in budding yeast.** The DNA damage-inducible checkpoint pathway in yeast can be divided into multiple steps including the sensor of damage and transduction of that signal, with subsequent cell cycle arrest and transcriptional induction (Fig. 9). Surprisingly, introduction of *CHES1* suppresses mutations which have been implicated in processing of damage (*rad9* and *rad24*) and in transduction of the signal (*mec1*, *rad53*, and *dun1*). The suppression of null alleles of *MEC1*, *RAD9*, and *DUN1* demonstrates that *CHES1* activity is not due to stabilization of the mutant gene products. Instead, our results suggest that in yeast *CHES1* must either act late in the pathway of response to damage after transcriptional induction or activate an alternative *MEC1*-independent pathway which

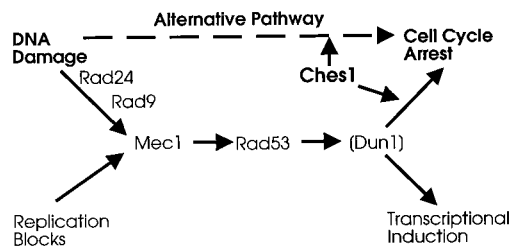


FIG. 9. A model for possible mechanisms of Ches1 action in cell cycle control and cellular response to DNA damage. Our data suggests that Dun1 may regulate both cell cycle arrest and transcriptional induction steps.

results in G<sub>2</sub> arrest after damage. Examples of other alternative pathways in yeast include suppression of *rad6* UV sensitivity by overexpression of the  $\alpha1/a2$  repressor (63). This suppression results from use of a *RAD52*-dependent recombination pathway. Similarly, there are rare survivors of *est1* mutations which escape senescence by maintaining telomere length through a parallel recombination pathway (35).

Our data support the alternate checkpoint pathway model due to the lack of detectable suppression of the *mec1* replication checkpoint (HU sensitivity) which would be expected for a protein acting downstream of *MEC1*. Activation of an alternative checkpoint pathway may not yield a phenotypic difference in wild-type strains due to an intact *MEC1*-dependent checkpoint pathway. Thus, expression of *CHES1* cDNAs in a wild-type strain would not be expected to yield a detectable difference in survival. The gene products required for the alternative checkpoint pathway can be further defined by identifying mutations which lose *CHES1*-dependent suppression of DNA damage checkpoints and proteins that interact with *CHES1* in yeast after DNA damage occurs.

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