Thesis for the Master's degree in Molecular Biosciences
Main field of study in Molecular Biology

The importance of Gcn2 and Rad3 in cell-cycle regulation of the G2/M transition

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60 study points

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Abstract

Previously the existence of a Gcn2-dependent G1/S checkpoint which delays entry into S phase in response to ultraviolet (UVC) radiation had been shown. Moreover, Gcn2 is active in S phase and G2 phase after UVC irradiation (Krohn et al., 2008). Therefore we investigated whether Gcn2 also affects cell-cycle progression from G2 into mitosis. The classic checkpoint pathway is known to arrest cell-cycle progression in response to DNA damage in G2. The kinase Rad3 plays a central role in this checkpoint since mutants with a deletion of rad3Δ are checkpoint deficient after treatment with ionizing radiation. We therefore reasoned that if Gcn2 has a minor effect on cell-cycle progression after UVC irradiation, this could best be seen in a rad3Δ checkpoint mutant.

To explore this hypothesis, a rad3Δgcn2Δ double mutant was made to study cell-cycle progression of synchronous cells after UVC irradiation in G2. Cell-cycle progression was monitored using a new flow cytometric method that conveniently measures the frequency of binuclear cells. The kinetics of entry into mitosis after UVC irradiation of rad3Δ and rad3Δgcn2Δ cells were the same. We conclude that Gcn2 does not play a significant role in regulation of G2/M after UVC irradiation. Interestingly, while investigating the role of Gcn2 in the G2/M transition, we found that UVC-treated rad3Δ cells delay mitosis in the same way as wild type cells do. Despite lacking the rad3 gene they elicit a checkpoint response to prevent entry into mitosis. This might indicate that other damage pathways exist. However, further studies are needed to characterize the checkpoint response in the absence of Rad3 and to uncover possible other players in the response to UV irradiation.
Contents

1. General introduction 5

2. Theoretical background 7
   2.1. *Schizosaccharomyces pombe* as a model organism 7
   2.2. DNA damage ............................................. 7
   2.3. The cell cycle in fission yeast .......................... 8
   2.4. Cell-cycle checkpoints .................................. 10
       2.4.1. The DNA-damage checkpoint ....................... 10
       2.4.2. The G1/S checkpoint ............................... 11
   2.5. Gcn2, cell-cycle regulation and cancer ................ 13

3. Aim of study 15

4. Materials 17
   4.1. Yeast strains ............................................ 17
   4.2. Enzymes .................................................. 17
   4.3. PCR primers ............................................. 18
   4.4. Antibodies ............................................... 18
   4.5. Molecular weight standards ............................ 19
   4.6. Chemicals and reagents ................................ 19
   4.7. Solutions ................................................ 20
       4.7.1. Growth media and agar plates .................... 20
       4.7.2. Buffers and other solutions ...................... 20

5. Methods 23
   5.1. Yeast media and cell handling .......................... 23
   5.2. Performing genetic crosses ............................. 24
   5.3. Genomic Mini prep in *S. pombe* ....................... 25
   5.4. Polymerase Chain Reaction (PCR) ....................... 26
   5.5. Synchronization methods ............................... 28
## Contents

5.6. UVC irradiation ............................................................... 30
5.7. Flow cytometry ................................................................. 30
5.8. Cell staining ................................................................. 33
5.9. Immunoblotting .............................................................. 36

6. Results ................................................................. 41
   6.1. Making \textit{gcn2 rad3} double mutants .............................. 41
   6.1.1. Polymerase Chain Reaction (PCR) of \textit{rad3} and \textit{gcn2} .... 43
   6.2. Measuring cell-cycle progression using \textit{cdc10} block and release ...... 45
     6.2.1. Cell-cycle analysis by flow cytometry ............................ 45
     6.2.2. Synchronization and timing of irradiation ......................... 46
     6.2.3. Entry into mitosis after UVC irradiation .......................... 49
     6.2.4. Cell-cycle progression monitored by fluorescent microscopy .... 50
   6.3. Measuring cell-cycle progression using a selection method for synchroniz-  
   6.3.1. Cell-cycle progression monitored by flow cytometry ............ 52
   6.4. The role of Rad3 after UVC irradiation ............................ 53
     6.4.1. Confirming the \textit{rad3}\textsuperscript{Δ} strain ...................... 53
     6.4.2. Cell-cycle progression of \textit{rad3}\textsuperscript{Δ} monitored by immunoblotting  55

7. Discussion .............................................................. 57

Appendix:

A. Molecular weight standards .................................................. 63

Bibliography .............................................................. 65
Chapter 1.

General introduction

Dysfunction of cell cycle checkpoints and DNA repair enzymes is known to be a characteristic of cancer cells (Krauss, 2008). Growth and proliferation of normal cells is carefully regulated by extracellular and intracellular signals. If those signal pathways are impaired, cells might escape the normal constraints on cell growth (increase in cell mass) and proliferation (increase in cell number) and thus cancer might develop. As cell division and replication are essential processes in cell proliferation, accumulation of changes in genes related to the cell-cycle or DNA repair is a major cause of cancer. Therefore the study of cell-cycle regulatory mechanisms and checkpoints are particularly relevant both to understand the processes leading to cancer development and as potential therapeutic targets.

Our group studies the G1/S transition using fission yeast (*Schizosaccharomyces pombe*) as a model organism. Tvegard *et al.* (2007) have identified a novel Gcn2-dependent checkpoint that regulates entry into S phase after treatment with ultraviolet (UVC) light. Furthermore, upon UVC irradiation the translation initiation factor eIF2α, a known target of Gcn2, becomes phosphorylated in G1, S and G2 phase (Krohn *et al.*, 2008). Phosphorylation of eIF2α mediated by Gcn2 is known as a stress response to amino acid deficiency leading to downregulation of global protein synthesis (Hinnebusch, 2005) but had not been linked to an UVC induced cell-cycle arrest before. Since at the G1/S transition Gcn2 is essential both for the translational downregulation and for the cell-cycle delay and in G2 Gcn2 is activated and downregulates translation (Krohn *et al.*, 2008), the question arises whether Gcn2 might also regulate the cell cycle at the G2/M transition. This is the subject of this master thesis.
Chapter 2.

Theoretical background

2.1. *Schizosaccharomyces pombe* as a model organism

Model organisms are simple organisms that allow to study and manipulate genes in order to understand fundamental mechanisms of cellular function. The choice of model organism often depends on the question being asked. Within animals, fungi, bacteria and plants various species are used as model organisms. Among fungi, *Schizosaccharomyces pombe*, is a very popular model used to study basic cellular mechanisms and their regulation. Failure of the corresponding pathways in human cells might lead to diseases. Therefore understanding the basic mechanisms in yeast is highly relevant for understanding the human diseases.

The fission yeast *S. pombe* is 3-4 µm in diameter and 7-15 µm in length. It has a typical eukaryotic cell cycle with G1, S, G2 and M phase and divides by binary fission (Moreno *et al.*, 1991). Cells are rod-shaped and have a simple structure where morphological changes can easily be observed. Sequencing of the genome by Wood *et al.* (2002) revealed that it has a size of 13.8 Mb coding for ~5000 genes distributed on three chromosomes. As fission yeast is not very specialized in evolutionary terms it is thought to represent genuine eukaryotic features which makes it an excellent model organism to study human disease genes; especially those involved in DNA damage and repair, cell-cycle control and recombination.

2.2. DNA damage

All living cells have to deal with DNA damage either caused by sun light and other naturally occurring DNA damaging agents or replication errors. Replication errors arise spontaneously when incorrect nucleotides are incorporated during replication. Normally the replication machinery removes these misincorporations
through its proofreading mechanism, but some errors might remain undetected. Mutagenic agents are another source of DNA damage. They include both chemicals and radiation (Brown, 2006). Base analogs, intercalating agents and alkylating agents belong to the group of chemical mutagens. Ultraviolet (UVC) and ionizing radiation (IR) are types of physical mutagens. UVC light induces covalent bonds between adjacent pyrimidines. The resulting dimers pose obstacles to the replicative DNA polymerase. IR is the most hazardous kind of radiation causing double strand breaks by attacking the DNA backbone (Watson et al., 2008).

Both spontaneous mutations (e.g. replication errors) and mutations caused by mutagens have to be repaired to prevent loss of genetic information. Therefore DNA damage activates a checkpoint which inhibits cell-cycle progression, which in turn allows time for DNA repair. Alternatively, if the damage is too severe to be repaired, programmed cell death is triggered to eliminate the cells with unreparable damage.

2.3. The cell cycle in fission yeast

The cell cycle is a series of events allowing cell duplication and division. Its goal is to produce two identical daughter cells. The cycle is divided into four different phases: G1 phase, S (synthesis) phase, G2 phase and M (mitosis) phase, where G1/S/G2 corresponds to interphase.

In S phase the nuclear DNA is replicated which leads to the doubling of genetic information. During M phase, the process of nuclear division, chromosomes are partitioned into two daughter cells. The remaining phases G1 and G2 are gaps between mitosis and S phase. They provide time for the cell to grow and to double their mass of proteins and organelles. Under typical laboratory growth conditions the cell cycle of fission yeast has, in contrast to the human cell cycle, a short G1 and a long G2 phase. Cytokinesis occurs just after mitosis, as in human cells. However, because of the short G1, completion of cytokinesis in fission yeast coincides with the end of S phase under standard laboratory growth conditions.

The cell cycle is highly coordinated so that each stage occurs in a defined order and at the right time point. Key regulators are Ser/Thr-specific protein kinases
that trigger the transition between each cell-cycle phase. These protein kinases, also called cyclin-dependent kinases (CDK), phosphorylate and activate other enzymes and proteins important during the cell cycle. Since they are essential for the major cell-cycle transitions, their activity is tightly controlled. The cells employ three major strategies to regulate CDK activity.

- In order to be active CDKs have to complex with cyclins. The expression level of the different cyclins rises and falls through the cell cycle, with different cyclins being expressed in different cell-cycle phases. This ensures that the right CDK activity is provided at the right time to the right level. Cdc2 is the only CDK in fission yeast. It associates with four different cyclins. The cyclin Cig2 complexes with Cdc2 in early G1 and remains bound until the end of S phase, Cig1 and Puc1 are associated with Cdc2 in late G1. The CDK-cyclin complex Cdc2-Cdc13 is present in S, G2 and M phase (Moser & Russel, 2000; Krohn et al., 2008).

- Phosphorylation of the kinase component. Phosphorylation of the T-loop, a flexible region of the kinase, leads to a conformational change so that the loop no longer blocks the catalytic pocket of the kinase. This phosphorylation stimulates the kinase activity enabling the substrate to access the active site. On the other hand phosphorylations on other sites of the catalytic pocket can decrease substrate affinity. In fission yeast phosphorylation of the kinase Cdc2 at tyrosine 15 (Y15) inhibits its activity.

- CDK inhibitors (CDI) bind to the CDK and thereby rearrange the catalytic site to inhibit its function (Alberts et al., 2008; Pollard et al., 2007).

**Cell-cycle regulation at the G2/M transition**

At the beginning of G2 Cdc2 is already associated with Cdc13, the cyclin needed for entry into mitosis. To prevent cells from going into mitosis at this early stage the Cdc2-Cdc13 complex is kept inactive by an inhibitory phosphorylation on Cdc2 Y15. The phosphorylation is maintained by the kinases Mik1 and Wee1 during G2 until cells are ready for mitosis. At the G2/M transition the inhibitory phosphate is removed by Cdc25 which activates Cdc2 and triggers entry into mitosis (figure 2.1).
2.4. Cell-cycle checkpoints

Checkpoints are control mechanisms that actively halt cell-cycle progression (Kastan & Bartek, 2004). In fission yeast several checkpoint pathways monitor the status of the DNA and arrest the cell cycle in response to DNA damage or inhibition of DNA replication. They include mechanisms to inhibit mitosis when the DNA is damaged (the G2/M checkpoint) or S phase is perturbed (the S/M checkpoint) as well as a mechanism to inhibit ongoing DNA replication when the DNA is damaged (the intra-S checkpoint). Other checkpoints relevant for this work are the G1/S checkpoint (see below) and the spindle assembly checkpoint. The spindle assembly checkpoint regulates transition from metaphase to anaphase and monitors tension and/or occupancy at the kinetochores (Lew & Burke, 2003). In the DNA-damage checkpoints pathways consist of sensor, mediator and effector components that transduce the checkpoint signal from DNA damage to an appropriate cell cycle response. At the end of the signaling cascade the CDK is inactivated by an inhibitory phosphorylation which leads to a cell-cycle arrest.

2.4.1. The DNA-damage checkpoint

In a log phase culture of fission yeast most cells will reside within G2 phase as it is the longest cell cycle phase. Therefore the G2/M transition is the major control point in fission yeast (Humphrey, 2000).

The first event in the checkpoint response is damage recognition. In the G2/M checkpoint this is executed by the rad checkpoint genes. There are two complexes that bind to damaged DNA independently of each other. This includes...
the checkpoint proteins Rad3-Rad26 and the Rad9-Rad1-Hus1 complex, also called the checkpoint sliding clamp (CSC) due to its homology to the PCNA, the sliding clamp for the polymerase in DNA replication (O’Connell et al., 2000). The CSC is loaded onto chromatin by yet another checkpoint rad protein, Rad17. Binding of Rad3 to damaged DNA induces phosphorylation of both Rad26 and the CSC. The mediator Crb2 serves as an adaptor protein between sensor and effector kinases (Kilkenny et al., 2008). Crb2 is required for Rad3/CSC-dependent phosphorylation of the checkpoint effector kinase, Chk1, which also undergoes autophosphorylation and thereby amplifies the DNA damage checkpoint signal (Humphrey, 2000). Chk1 inactivates Cdc25 by an inhibitory phosphorylation and increases the activity of Wee1 by phosphorylation. Both phosphorylations lead to the accumulation of Cdc2 in its Y15-phosphorylated, inactive form and consequently to a cell cycle arrest (Figure 2.2).

Figure 2.2: The checkpoint response for the DNA damage checkpoint at the G2/M transition.

2.4.2. The G1/S checkpoint

The G1/S checkpoint in *S. pombe* is dependent on the kinase Gcn2 (Tvegard et al., 2007). The Gcn2-dependent checkpoint can be activated by UVC, methyl methanesulfonate (MMS) and H₂O₂ and leads to inhibition of an early step in the preparation for S phase and thereby to a delay at initiation of DNA replication.
The only known substrate for Gcn2 is eIF2α, which is phosphorylated in response to UVC, MMS and H₂O₂ (Krohn et al., 2008). Phosphorylation of eIF2α leads to a downregulation of general translation, although some selected mRNAs are still translated (Krauss, 2008). It is as yet unclear whether there is a causal relationship between translation downregulation and cell-cycle delay. It is conceivable that Gcn2 phosphorylates an unknown substrate that is required for entry into S phase. It is also possible that eIF2α phosphorylation and/or downregulation of translation is directly responsible for the cell-cycle delay, presumably by translational regulation of a protein required for entry into S phase.

It is not yet known what the signal is that activates Gcn2. UVC and the other agents cause DNA damage, as well as damage to RNA and proteins. It is not clear whether the signal derives from DNA damage or damage to other macromolecules. Gcn2 is known for its role in translation downregulation after starvation. In that case, it is activated by the accumulating uncharged tRNAs. It is not immediately obvious how UVC, MMS and H₂O₂ might lead to the accumulation of uncharged tRNAs (figure 2.3).

![Figure 2.3: The G1/S checkpoint model (Krohn et al., 2008).](image-url)
2.5. Gcn2, cell-cycle regulation and cancer

Cell growth and cell-cycle progression are tightly coupled processes (Fingar & Blenis, 2004; Petersen & Nurse, 2007). The cell has to increase in cell mass through biosynthesis of macromolecules to maintain cell size at division. Otherwise cells would get smaller with every cell cycle. Therefore cell growth requires a sufficient nutrient supply. As reviewed in Hinnebusch (2005) Gcn2 is a kinase which senses the availability of nutrients (amino acids) and is activated by uncharged tRNAs. It has also been reported that Gcn2 is activated by glucose deprivation. In this case activation is indirect and due to the fact that amino acids are used as a secondary energy source (Wek & Staschke, 2010).

It has been shown by Deng et al. (2002) that Gcn2 is activated by UVC light also in human cells, but it is not clear yet whether it is important for the regulation of G1/S in human cells. Given the observation that the activity of Gcn2 is important for the survival of cancer cells (Wek & Staschke, 2010; Ye et al., 2010), it is all the more important to understand whether and how Gcn2 might regulate the cell cycle.
Chapter 3.

Aim of study

A novel checkpoint which is dependent on Gcn2 and inhibits entry into S phase after UVC irradiation in G1 was described by Tvegard et al. (2007). It could also be shown that Gcn2 is activated after UVC irradiation in G2 (Krohn et al., 2008). This observation led to the question whether activation of Gcn2 after UVC irradiation in G2 has any effect on cell-cycle progression from G2 phase into mitosis. The aim of this study is to investigate the role of Gcn2 in the G2/M transition.
Chapter 4.

Materials

4.1. Yeast strains

The following S. pombe strains were used in this study.

Table 4.1: \textit{Schizosaccharomyces pombe}

<table>
<thead>
<tr>
<th>Strain</th>
<th>Referred to as</th>
<th>Genotype</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>489</td>
<td>wild type</td>
<td>cdc10-M17 h-</td>
<td>lab collection</td>
</tr>
<tr>
<td>1137</td>
<td>\textit{gen2}Δ</td>
<td>\textit{gen2::ura4+ cdc10-M17 ura4-D18 h+}</td>
<td>lab collection</td>
</tr>
<tr>
<td>1153</td>
<td>\textit{rad3}Δ</td>
<td>\textit{rad3::ura4+ leu1-32 ura4-D18 ade6-704 h-}</td>
<td>lab collection</td>
</tr>
<tr>
<td>1353</td>
<td>\textit{rad3}Δ</td>
<td>\textit{rad3::ura4+ cdc10-M17 ura4-D18 h+}</td>
<td>lab collection</td>
</tr>
<tr>
<td>1638</td>
<td>\textit{gen2}Δ \textit{rad3}Δ</td>
<td>\textit{gen2::ura4+ rad3::ura4+ cdc10-M17 ura4-D18}</td>
<td>this study</td>
</tr>
</tbody>
</table>

4.2. Enzymes

Enzymes presented in table 4.2 were used for PCR, crossing of yeast strains, isolation of DNA and flow cytometry.

Table 4.2: List of enzymes and their suppliers.

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Supplier</th>
</tr>
</thead>
<tbody>
<tr>
<td>High Fidelity Polymerase</td>
<td>Roche</td>
</tr>
<tr>
<td>Ribonuclease A</td>
<td>Sigma-Aldrich</td>
</tr>
<tr>
<td>S.H.P/H.P.J Helix Pomatia Juice (helicase)</td>
<td>BioSepra</td>
</tr>
<tr>
<td>Zymolase 20T</td>
<td>MP Biomedicals</td>
</tr>
</tbody>
</table>
4.3. PCR primers

Primers listed in table 4.3 were used for amplification of *gcn2* or *rad3*.

**Table 4.3.:** List of primers and primer sequences. Primers were delivered by Invitrogen.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primer</th>
<th>Direction</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>gcn2</em></td>
<td><em>SP</em>/BamHI_ Gcn2_ ORF</td>
<td>forward</td>
<td>5’ TAGCGGATCCTAAAAGCTGTTCATGGAATGC 3’</td>
</tr>
<tr>
<td></td>
<td><em>ASP</em>/XmaI_ Gcn2_ ORF</td>
<td>revers</td>
<td>5’ TAGCCCCGGGTTTAAGTCATAAGATAAA 3’</td>
</tr>
<tr>
<td><em>rad3</em></td>
<td>Rad3_sense</td>
<td>forward</td>
<td>5’ TATTTTTGACGTTTTGGACAGG 3’</td>
</tr>
<tr>
<td></td>
<td>inura4rev2</td>
<td>revers</td>
<td>5’ CCAGGATATGGTAGAAAAAC 3’</td>
</tr>
</tbody>
</table>

4.4. Antibodies

Primary and secondary antibodies used for immunoblotting and cell staining are presented in table 4.4 and 4.5.

**Table 4.4.:** List of primary antibodies and their suppliers.

<table>
<thead>
<tr>
<th>Primary antibody</th>
<th>Supplier</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anti-cdc2-P antibody, rabbit</td>
<td></td>
</tr>
<tr>
<td>Anti-tubulin antibody, mouse</td>
<td></td>
</tr>
<tr>
<td>TAT1 antibody, mouse</td>
<td></td>
</tr>
</tbody>
</table>

**Table 4.5.:** List of secondary antibodies and their suppliers.

<table>
<thead>
<tr>
<th>Secondary antibody</th>
<th>Supplier</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cy3-linked antibody, Sheep anti-mouse, IgG</td>
<td>GE Healthcare</td>
</tr>
<tr>
<td>alkaline phosphatase-linked antibody, Goat anti-mouse, IgG + IgM (H+L)</td>
<td>GE Healthcare</td>
</tr>
<tr>
<td>alkaline phosphatase-linked antibody, Goat anti-rabbit, IgG + IgM (H+L)</td>
<td>GE Healthcare</td>
</tr>
</tbody>
</table>
4.5. **Molecular weight standards**

Molecular weight standards were used as a reference point to estimate DNA or protein size on an agarose or polyacrylamid gel. They are listed in table 4.6 and 4.7.

**Table 4.6.:** DNA standard and supplier.

<table>
<thead>
<tr>
<th>DNA molecular weight standard</th>
<th>Range</th>
<th>Supplier</th>
</tr>
</thead>
<tbody>
<tr>
<td>O’GeneRuler™ 1kb DNA ladder</td>
<td>0.25-10 kb</td>
<td>Fermentas</td>
</tr>
</tbody>
</table>

**Table 4.7.:** Protein standard and supplier.

<table>
<thead>
<tr>
<th>Protein molecular weight standard</th>
<th>Range</th>
<th>Supplier</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dual Color Precision Plus Protein Standard</td>
<td>10-250 kDa</td>
<td>Bio-Rad</td>
</tr>
</tbody>
</table>

4.6. **Chemicals and reagents**

Chemicals and reagents used are presented in table 4.8.

**Table 4.8.:** List of chemicals and their suppliers.

<table>
<thead>
<tr>
<th>Chemical</th>
<th>Supplier</th>
</tr>
</thead>
<tbody>
<tr>
<td>DAPI (4,6-diamidino-2-phenylindole)</td>
<td>Kemetyl Norge AS</td>
</tr>
<tr>
<td>Ethanol</td>
<td>Merck</td>
</tr>
<tr>
<td>Methanol</td>
<td></td>
</tr>
<tr>
<td>Phloxin B</td>
<td></td>
</tr>
<tr>
<td>Sytox green</td>
<td>Invitrogen</td>
</tr>
<tr>
<td>Trichloroacetic acid (TCA)</td>
<td></td>
</tr>
</tbody>
</table>
4.7. Solutions

4.7.1. Growth media and agar plates

Formulations of growth media and agar plates are detailed in table 4.9.

**Table 4.9.:** Yeast media and recipes.

<table>
<thead>
<tr>
<th>Media</th>
<th>Recipe</th>
</tr>
</thead>
<tbody>
<tr>
<td>YE 50 medium</td>
<td>17.625 g YE 50 (Sunrise Science products # 2009-500) to 500 ml H₂O, sterile filtrated</td>
</tr>
<tr>
<td>YE 50 agar</td>
<td>26.125 g agar powder (Sunrise Science products # 2010-500) to 500 ml H₂O, autoclaved at 121°C for 15 min</td>
</tr>
<tr>
<td>YE 50 agar with phloxin B</td>
<td>YE 50 agar supplemented with 5 mg/liter phloxin B</td>
</tr>
<tr>
<td>YE 50 agar with hydroxyurea (HU)</td>
<td>YE 50 agar supplemented with 15 mM HU</td>
</tr>
<tr>
<td>EMM medium</td>
<td>15.9 g EMM (Sunrise Science products # 2005-500) to 500 ml H₂O, sterile filtrated</td>
</tr>
<tr>
<td>EMM agar</td>
<td>24.4 g EMM agar (Sunrise Science products # 2006-500) to 500 ml H₂O.</td>
</tr>
</tbody>
</table>

4.7.2. Buffers and other solutions

Formulations of buffers and other solutions used in this study are presented in table 4.10.

**Table 4.10.:** Solutions and recipes.

<table>
<thead>
<tr>
<th>Solution</th>
<th>Recipe</th>
</tr>
</thead>
<tbody>
<tr>
<td>Agarose gel solution, 1%</td>
<td>1 % agarose</td>
</tr>
<tr>
<td></td>
<td>1 x TAE buffer</td>
</tr>
<tr>
<td>Citrate/phosphate pH 5.6</td>
<td>7.1 g/l Na₂HPO₄</td>
</tr>
<tr>
<td></td>
<td>11.5 g/l citric acid</td>
</tr>
<tr>
<td>Citrate/phosphate/EDTA/sorbitol</td>
<td>50 mM citrate/phosphate pH 5.6</td>
</tr>
<tr>
<td></td>
<td>40 mM EDTA pH 8.0</td>
</tr>
</tbody>
</table>
1 M sorbitol

**DTT, 1M**
5g DTT
32.4 ml 0.01M NaAc

**EDTA, pH 8.0 (0.5 M)**
146.12 g/L EDTA
NaOH to pH 8.0

**Electrophoresis buffer, 10x**
30.2 g/L tris
144 g/L glycine
1% (w/v) SDS

**Lactose solution, 10%**
10 g lactose
100 ml EMM

**Lactose solution, 40 %**
40 g lactose
100 ml EMM

**Laemmli buffer, 2x**
1.25 ml 1 M Tris HCl, pH 6.8
2.0 ml Glycerol
4.0 ml 10 % SDS solution
0.5 ml 0.1% (w/v) bromophenol blue
1 M DTT (final concentration 0.2 M)
deionized H₂O to 10 ml

**LiAc/TE, 10x**
1 M lithium acetate
1x TE

**PEG/LiAc/TE**
40% PEG 4000
1x LiAc/TE

**PEM**
100 mM PIPES pH 6.9
1 mM EGTA
1 mM MgSO₄

**PEMBAL**
PEM
0.1 M L-lysine
1% BSA (globulin free)
0.1% NaN₃

**Potassium acetate, 5 M pH 5**
118 g KAc
46 ml acetic acid
with acetic acid adjusted to pH 5
### Materials

<table>
<thead>
<tr>
<th>Buffer Type</th>
<th>Components</th>
</tr>
</thead>
<tbody>
<tr>
<td>SDS, 20%</td>
<td>10 g SDS</td>
</tr>
<tr>
<td></td>
<td>50 ml H₂O</td>
</tr>
<tr>
<td>TBS-Tween</td>
<td>20 mM Tris HCl, pH 7.5</td>
</tr>
<tr>
<td></td>
<td>8 g/L NaCl</td>
</tr>
<tr>
<td></td>
<td>0.05% (v/v) Tween-20</td>
</tr>
<tr>
<td>TE, 10x pH 7.5</td>
<td>0.1 M Tris HCl, pH 8.0</td>
</tr>
<tr>
<td></td>
<td>0.01 M EDTA pH 8.0</td>
</tr>
<tr>
<td></td>
<td>with HCl adjusted to pH 7.5</td>
</tr>
<tr>
<td>Transfer buffer</td>
<td>50 mM Tris base</td>
</tr>
<tr>
<td></td>
<td>380 mM glycine</td>
</tr>
<tr>
<td></td>
<td>0.1% (w/v) SDS</td>
</tr>
<tr>
<td></td>
<td>20% (v/v) methanol</td>
</tr>
</tbody>
</table>
Chapter 5.

Methods

5.1. Yeast media and cell handling

All media formulations are detailed in chapter 4.

Yeast extract medium (YE)
Yeast extract medium is a complete, rich but poorly defined medium which provides optimal growth conditions. When testing different genotypes YE medium was supplemented with phloxin B, a stain used to distinguish between dead and living yeast cells. Yeast cells growing on solid media containing phloxin B will range in color from light pink for living colonies to dark red for dead colonies (Kucsera et al., 2000; Forsburg, 2003b). Hydroxy urea (HU) added to YE medium facilitated identification of HU sensitive mutants. HU inhibits the ribonuclease reductase and thereby depletes the supply of nucleotides. The checkpoint rad mutants are particularly sensitive to HU. They have a defective G2 arrest and will continue through the cell cycle despite incomplete replication due to the lack of nucleotides in S phase.

Edinburgh minimal medium (EMM)
A minimal medium contains only essential nutrient sources, salts and vitamins. In EMM all components are known and defined. It does not include any supplements and will therefore inhibit growth of mutants with auxotrophies (e.g. adenine, uracil). EMM was used when stable physiological conditions were needed as in experiments for cell-cycle progression.

Malt extract agar (MEA)
MEA is a complete, meagre, poorly defined medium. Cells growing on MEA soon run out of nutrients and, if cells of the opposite mating type are present, enter the
sexual differentiation pathway. MEA was used for conjugation and sporulation.

**Storage of fission yeast**

Yeast strains can be stored at -80°C. For this purpose a cell culture was grown to late-logarithmic phase and then mixed with 50% sterile glycerol in a 1:1 ratio. To reisolate cells, a small amount of the frozen glycerol stock is plated onto YE medium and incubated at 25°C.

**Growth of liquid cultures**

For physiological experiments (e.g. UVC irradiation) a fresh colony was inoculated in 10 ml YE medium and incubated at 25°C until the next day (preculture). Cells from the preculture were transferred to liquid EMM and grown in log phase for 2 days at 25°C. Exponential growth of the culture is obtained using the following equation for dilution from the preculture:

\[
V_{\text{preculture}} = \frac{V_{\text{total}} \cdot OD_{\text{desired}}}{2^n \cdot OD_{\text{preculture}}} \quad \text{and} \quad n = \frac{t}{t_D}
\]

where \( V_{\text{preculture}} \) is the volume of the preculture used to inoculate, \( V_{\text{total}} \) is the total volume of the new culture, \( OD_{\text{desired}} \) is the desired \( OD_{595} \) at a given time and \( OD_{\text{preculture}} \) is the OD measured for the preculture. The number of generations \( n \) is defined by the time cells are grown in liquid medium \( t \), divided by the doubling time of a given strain \( t_D \).

**5.2. Performing genetic crosses**

In fission yeast conjugation and sporulation occur only when cells are starved for nutrients (Moreno et al., 1991). MEA plates are usually used for genetic crossing. The two strains that are going to be crossed have to be of opposite mating type, h+ and h-. For crossing, a colony from each of the freshly plated strains was mixed and dissolved in 500 µl H₂O. 15 µl of the solution was then spotted onto MEA plates and left to dry. The cross was incubated at 25°C for 2 days.

When asci could be seen under a light microscope (Zeiss, Germany), a colony with asci was resuspended in 500 µl H₂O. Then 5 µl helicase (Helix pomatia juice) was added and the mixture was incubated for 4 hours at 32°C. The digestive juice of *Helix pomatia* is used to break down the cell wall of the asci and to kill vegetative
cells (Moreno et al., 1991). Spores were plated out on YE agar and incubated at 25 °C until colonies formed.

### 5.3. Genomic Mini prep in *S. pombe*

For isolation of DNA from *S. pombe* a cell culture was grown to saturation in YE. Samples were prepared as follows:

1. The cell culture was centrifuged at 1100 x g for 5 min.
2. The pellet was washed once with ice cold water.
3. 1 ml citrate/phosphate/EDTA/sorbitol containing 2,5 mg zymolase was added for enzymatic digestion of the cell wall.
4. The suspension was transferred to an Eppendorf tube and incubated at 36 °C for 60 min. Spheroplasting was checked by adding 1 µl 10% SDS to 9 µl of the digested cells on a microscope slide.
5. Cells were then pelleted at 16 000 x g for 10 seconds The supernatant was removed and pellet resuspended in 500 µl TE.
6. 25 µl 20% SDS was added, the tube inverted several times and incubated at 65 °C for 1 h.
7. 175 µl 5 M potassium acetate (pH 5) was added, vortexed and kept on ice for 15 min.
8. After that the solution was centrifuged at 16 000 x g for 10 min at 4 °C.
9. The supernatant was carefully transferred to an other Eppendorf tube
10. 500 µl ice-cold isopropanol was added followed by mixing and another centrifugation step; 16 000 x g for 10 min at 4 °C.
11. The supernatant was removed and the pellet was washed with 500 µl 70% EtOH.
12. The pellet was air-dried over night and then resuspended in 40 µl TE
13. To confirm successful DNA extraction, 5 µl sample was analyzed by gel electrophoresis.
Purified DNA was stored at -20 °C.

5.4. Polymerase Chain Reaction (PCR)

Polymerase chain reaction is a technique used to amplify a specific DNA sequence from genomic DNA in a simple enzymatic reaction. The major components in a PCR are a heat-stable DNA polymerase, 2 oligonucleotide primers, a DNA template and deoxynucleotide triphosphates (dNTPs), in an appropriate buffer that ensures optimal enzyme activity. For the generation of primers it is important to know the sequence of interest as each primer is complementary to the respective end of the DNA sequence. In a PCR one amplification cycle consists of 3 steps: DNA denaturation, primer annealing and DNA synthesis.

In the first step (denaturation), DNA is separated into single strands by heat (94-95 °C). Then the temperature is lowered to allow annealing of the two primers to their complementary sequences in the template. Finally, primers are elongated by DNA polymerase synthesizing new DNA duplexes. A PCR usually consists of a series of 30-40 cycles which results in an exponential amplification in the amount of the chosen DNA sequence (Fletcher et al., 2007; Berg et al., 2007). For all PCR reactions the High Fidelity PCR Master kit was used.

Polymerase Chain Reaction of *gcn2*

Primers used for amplification of *gcn2* were SP/BamHI *gcn2* ORF and ASP/XmaI *gcn2* ORF (see Chapter 4 - Materials: table 4.3). The primers are sequence-specific for *gcn2* and give PCR products for both *gcn2* deletion and *gcn2* wild type.
5.4 Polymerase Chain Reaction (PCR)

Figure 5.1: Primers used for amplification of gc2.

The reaction was run on a thermal cycler (Mastercycler® gradient, Eppendorf, Hamburg, Germany). PCR conditions are outlined in figure 5.2. The PCR product was analyzed by gel electrophoresis.

Figure 5.2: PCR conditions for gc2 amplification.

Polymerase Chain Reaction of rad3

Primers used to confirm the rad3 deletion were Rad3_sense and inura4rev2 (see Chapter 4 - Materials: table 4.3). Rad3_sense is specific for the rad3 gene, inura4rev2 anneals to the ura4-marker. Therefore only the rad3 gene carrying the ura4-marker will give a PCR product. There is no product expected for wild type rad3.

The reaction was run on a thermal cycler (Mastercycler® gradient, Eppendorf, Hamburg, Germany). PCR conditions are outlined in figure 5.4. The PCR product
5.5. Synchronization methods

Synchronization is an important tool when analyzing cell-cycle progression since cells in a synchronous culture are dividing at the same time (Walker, 1999). There are several different synchronization methods to promote synchrony in an otherwise randomly dividing yeast population. In this study two different methods for synchronization were used.

The induction method of synchronization

One particular induction method is block and release of cdc mutants. In this study, synchrony was achieved using temperature sensitive \( cdc10-M17 \) mutants. In \( S.\ pombe \), Cdc10 is a transcription factor required for expression of transcripts in G1. Incubation of \( cdc10^{ts} \)-cells at the restrictive growth temperature (36°C) blocks cell-cycle progression at the G1 stage (Sabatinos & Forsburg, 2010). Releasing the cells from temperature block results in a synchronous G1 culture.

In the current experiments, cells were grown to an OD=0.15 in minimal medium.
at 25°C, then shifted to 36°C for 4h to arrest them in G1, then returned to 25°C to release them into the cell cycle (figure 5.5).

**Figure 5.5.:** Synchronization by block and release. Temperature sensitive *cdc10* mutants were incubated for 4 h at the restricted temperature to give a synchronous G1 culture.

**The selection method of synchronization**

Another method to synchronize a population of cells uses a sugar gradient to separate cells according to their size. Since fission yeast cells are very regular in size, selection of the smallest cells in a population results in a synchronous G2 culture. The sugar of choice for gradient-based synchronization of fission yeast cells is lactose because it cannot be metabolized by fission yeast and will therefore not perturb growth. The gradient was established using a gradient maker with a 10% and a 40% lactose solution (see materials for recipe).

There were always two consecutive gradients applied to achieve the best result for separation. Cultures were grown to mid-log phase, then loaded onto the gradient at a cell concentration of 5x10^9/ml. The protocol used for lactose gradient synchronization is outlined below.

1. 500 ml cell suspension (**OD_595 = 0.5**) was centrifuged for 3 min at 1100 x g.
2. The pellet was resuspended 5 ml YE medium.
3. 5 ml cell suspension was added onto the first gradient by pipetting along the inside of the tube.
4. The gradient with the applied cell suspension was then centrifuged for 3 min at 230 x g.
5. Afterwards 6 aliquots ≈ 1 ml were collected from the top of the gradient.
5 Methods

6. To remove the remaining lactose in the collected suspension it was spun down at 1100 x g for 3 min and resuspended in 1 ml EMM.

7. 1 ml of the collected and resuspended suspension was added onto the second gradient similar to the first one by pipetting carefully along the inside of the tube.

8. The gradient with the applied cell suspension was then centrifuged for 3 min at 230 x g.

9. 500 µl aliquots were collected from the top of the second gradient; in total 6 aliquots each in separate Eppendorf tube.

10. Synchrony was checked under the microscope. For good synchrony the cell size should be uniform and there should not be any cells with septa because septated cells are not in G2 but in G1 or S phase.

11. Cells from aliquots with the best synchrony were spun down at 1100 x g for 3 min to remove excess lactose and resuspended in 30 ml EMM.

5.6. UVC irradiation

In all experiments irradiation was done in G2 phase.

A suspension of synchronous cells was poured into a Petri dish to cover the bottom with a thin layer (3 mm). The cells were stirred while they were irradiated with 254 nm UVC light. The dose given was 1100 J/m² in about 3 min measured with a radiometer (UVCP, Upland, CA, USA). Samples were taken immediately after irradiation and then every 15 min.

5.7. Flow cytometry

Flow cytometry is a technique which facilitates analysis of DNA content in yeast cells. Analysis was performed with a BD™LSR II Flow Cytometer with 488-nm excitation light. Collected samples were prepared as follows:

1. 1 ml of cell suspension was spun down at 16 000 x g for 2 min, the supernatant removed and cells fixed in 1 ml ice-cold 70% EtOH on a vortexer. (Samples can be stored in the refrigerator.)
2. 500 µl of the sample was spun down at 16 000 x g for 2 min

3. The pellet was resuspended in 1 ml 20 mM EDTA and spun down at 16 000 x g for 2 min twice.

4. The supernatant was discarded and pellet resuspended in 500 µl 20 mM EDTA containing 100 µg/ml RNase A.

5. Samples were incubated over night at 36 °C.

6. Finally 500 µl 20 mM EDTA containing 2 µM Sytox Green was added.

7. Before running the flow cytometer samples were sonicated for 5 min and transferred to 5 ml tubes.

LSR II Flow Cytometer was run as described in the core facility manual. Analysis was performed as in Knutsen et al. (2011). This new method enables separation between mononuclear and binuclear cells by measuring the width as well as the total area of the DNA signal. Fluorescently labeled DNA remains in the excitation focus for a longer period of time when there are two separated nuclei than when there is only one nucleus in the cell. Therefore, the DNA signal (DNA-W) is expected to last longer for a binuclear than for a uninuclear cell (see figure 5.6 Panel C-E versus F-K). We also showed that by employing light-scatter measurements it is possible to exclude cell doublets (figure 5.6 Panel, and L-N) Binuclear cells were gated using the flow cytometry software to estimate the relative amount of binuclear cells (the binuclear index, BI).
Figure 5.6: Analysis of exponentially growing cells. Flow cytometric and microscopic analyses of wild-type S. pombe cells in an exponentially growing culture. First column: cytograms showing DNA-W versus DNA-A; second column: microscopy images of the sorted cells; third column: cytograms showing FSC versus SSC. The two-parametric DNA-A/DNA-W cytograms show four main subpopulations of cells: 1 G2 cells with a single nucleus and a 2C DNA content, 2 cells in G1 phase or late in mitosis with two nuclei and 2x 1C DNA content, 3 cells in S phase, with a DNA content between 2C and 4C, 4 cell doublets. From Knutsen et al. (2011).
5.8. Cell staining

Cell-cycle progression was not only monitored by flow cytometry but also by two different staining techniques: Staining of nuclear DNA or septa with fluorescent dye (4',6-diamidino-2-phenylindole (DAPI) and aniline blue, respectively) and staining of tubulin with a fluorescently-labeled antibody. The two dyes staining DNA and tubulin, visualize cellular structures (nucleus and mitotic spindle, respectively) that specifically enable determination of the number of cells in mitosis. Septated cells, determined by aniline blue staining are an index of cells in G1/S phase.

DAPI staining
Samples were fixed in 70% ice-cold EtOH, rehydrated and stained with DAPI to visualise chromosomes under a fluorescence microscope. Cells in mitosis show to distinct sets of chromosomes in the same cell. A detailed procedure of how samples were prepared for DAPI staining is outlined below.

1. 1 ml of an early-mid log phase culture was spun down at 16 000 x g for 2 minutes, the supernatant removed and cells fixed in 1 ml ice-cold 70% EtOH on a vortexer.
2. 500 µl of EtOH fixed cells was spun down at 16 000 x g for 2 minutes. The supernatant was discarded and pellet resuspended in 20 µl H₂O.
3. 5 µl suspension was placed onto a microscope slide and allowed to dry. A cover slip with 2,5 µl DAPI in 50% glycerol was applied.

Samples were observed with a fluorescence microscope (Leica DM6000B, Wetzlar, Germany) with 63x magnification. The MI was determined counting those cells that had two discernable nuclei but no septum.

Staining with aniline blue
Aniline blue stains the septum and was used to monitor the frequency of septated cells (septation index). A septum appears after nuclear division and is normally located at the midpoint of the newly divided cell. The procedure for staining with aniline blue is as for DAPI staining. Samples were fixed in 70% EtOH.
1. 1 ml of cell suspension was spun down at 16 000 x g for 2 minutes, the supernatant removed and cells fixed in 1 ml ice-cold 70% EtOH on a vortexer.

2. 500 µl of EtOH fixed cells was spun down at 16 000 x g for 2 minutes. The supernatant was discarded and pellet resuspended in 20 µl H₂O.

3. 5 µl suspension was placed onto a microscope slide and allowed to dry. A cover slip with 2,5 µl aniline blue was applied.

Samples were observed with a fluorescence microscope (Leica DM6000B, Wetzlar, Germany) with 63x magnification. The septation index (SI) was determined counting cells with a septum (Figure 5.7).

![Figure 5.7](image)

_Figure 5.7:_ Determining the frequency of septated cells. From Forsburg (2003a).

**Immunostaining with anti-tubulin antibodies**

β-tubulin is the major component of microtubules. Stained with a fluorescently-labeled anti-tubulin antibody it is possible to distinguish mitotic cells from interphase cells, based on their structural differences between interphase and mitotic microtubules (figure 5.8). Immunostaining with anti-tubulin antibodies and counting cells with a mitotic spindle provides another method to determine the MI.

For tubulin staining 10⁸ cells were harvested for each sample. Samples were fixed in methanol (MeOH) prior to immunofluorescence labeling. A description of the procedure is outlined below.
1. Each sample was spun down at 1100 x g for 3 min and the supernatant was removed thoroughly.

2. Cells were fixed by adding 15 ml ice-cold MeOH.

3. To prepare samples for immunostaining with anti-tubulin antibodies, cells were spun at 1100 x g for 8 min.

4. Cells were resuspended in 1 ml PEM and transferred to Eppendorf tubes.

5. Then spun again at 6 000 x g for 1 min and resuspended in 1 ml PEM with 3 mg/ml Lysing enzyme and 1 mg/ml Zymolyase 20T.

6. Cells were incubated at 37 °C until ~80% cells were digested.

7. Afterward samples were washed with 1ml PEM, spun at 6 000 x g for 1 min and resuspended in 1 ml 1% Triton in PEM for 30 seconds.

8. Samples were spun again at 6 000 x g for 1 min and wash 3x in 1 ml PEM.

9. The volume of the final pellet was roughly assessed.
10. Cells were resuspended in 1 ml PEMBAL and a volume which will give a 20-30 µl pellet upon a subsequent spin was transferred to each of two Eppendorf tubes.

11. Samples were put on a rotating wheel for 60 min at room temperature and after that spun at 6 000 x g for 1 min.

12. Cells were resuspended in 100 µl primary antibody (1/10 TAT1 in PEMBAL - supplied by Keith Gull) and incubated 16 h at 4°C on a rotating wheel.

13. Cells were washed 3x in 1 ml PEMBAL.

14. Cells were resuspended in 50 µl CY3-linked anti-mouse secondary antibody (1:200 in PEMBAL) for 1-2 h at room temperature. Samples tubes were wrapped in foil to protect the fluorophore from light to avoid bleaching.

15. Subsequently samples were washed 3x in 1 ml PEMBAL.

16. Finally the pellet was resuspended in 100 µl PEMBAL and cells were mounted on PolyL-lysine coated cover slips.

17. They were dried on a heating block and a cover slip with 2,5 µl DAPI in 50% glycerol was applied.

Samples were observed under a fluorescence microscope with 63x magnification (Leica DM6000B, Wetzlar, Germany) immediately after preparation. The MI was determined counting those cells that showed a single microtubule structure.

5.9. Immunoblotting

The third method applied to monitor cell-cycle progression is immunoblotting. Immunoblotting can be used to monitor the presence of cell-cycle regulated proteins and/or their modified forms. Dephosphorylation of the checkpoint kinase Cdc2 triggers entry into mitosis. By comparing the presence of dephosphorylated Cdc2-Y15 in untreated and UVC-irradiated samples it is possible to distinguish G2-arrested and M-arrested cells. Immunoblotting consists of several steps. Those included snap-freezing of cells after sampling, extraction of proteins from frozen samples, separation of proteins by sodiumdodecyl sulphate polyacrylamid gel electrophoresis (SDS-PAGE), transferring proteins from gel to membrane (Western blotting) and finally immunodetection using protein-specific antibodies.
Each technique is outlined below.

**Snap-freezing of protein samples**
10⁸ cells were harvested for each sample.

1. Each sample was spun down at 1 100 x g for 3 min (4 °C) and the pellet was resuspended in 1 ml H₂O.

2. The cell suspension was transferred to a 2 ml tube which was followed by another centrifugation step at 1 100 x g for 2 min (4 °C).

3. The supernatant was removed and the cell pellet was frozen in liquid N₂ (-196 °C) to stop all cellular processes immediately.

Samples were stored at -80 °C.

**TCA protein extraction**

1. The cell pellet was thawed at room temperature and resuspended in 300 µl of 20% trichloroacetic acid (TCA)

2. 500 µl glass beads was added.

3. The tube containing cells, TCA and glass beads was placed in the Ribolyzer and cells were lysed 3x 20 sec at 6.5 m/s until broken cells were seen under a light microscope.

4. 600 µl 5% TCA was added and mixed thoroughly.

5. The cell lysate was spun into a new tube as follows: the lid of the tube with protein sample and beads was punctured with a needle and placed on top of a 1.5 ml screw cap tube upside down. Both tubes were placed in a 15 ml Falcon tube (figure 5.9)

6. Samples were spun for 1 min at 1 100 x g to remove the cell lysate from the glass beads.

7. Following another centrifugation at 16 000 x g for 10 min the supernatant was discarded and the protein pellet resuspended in 200 µl sample buffer (1x Laemmli buffer + 200 mM Tris-HCl pH 8)
8. Boiled samples (heating block; 100 °C) were spun 1 min at 16 000 x g. The supernatant was transferred to a new tube and the pellet was discarded as it mainly contains unlysed cells.

Protein samples were frozen at -80 °C.

**SDS-PAGE**

SDS-PAGE is a qualitative method to analyze protein samples where proteins are separated based on their molecular weight. Proteins are dissolved in Laemmli buffer (sample buffer, see 9. under TCA protein extraction), which contains both dithiothreitol (DTT) that reduces disulfide bonds in the tertiary structure of proteins and SDS, an anionic detergent which denatures proteins. SDS binding unfolds polypeptide chains and confers a uniform negative charge per unit length. The migration of proteins is therefore dependent on their molecular weight rather than their intrinsic charge.

In preparation to SDS-PAGE, frozen protein samples were boiled in a heating block (100°C) to ensure total denaturation of the proteins. The gel was placed in a gel chamber filled with 1x running buffer and 10 µl samples were loaded onto the gel. The Precision Plus Dual Color was used as sizing marker (Appendix). Proteins were run on a polyacrylamid gel at 160 V for about 50 min. The current applied makes the negative charged protein-SDS molecules move towards the anode. Smaller molecules migrate faster than larger ones.
Western blotting

Western blotting is a technique where proteins separated by SDS-PAGE are transferred from the gel onto a membrane before reaction with antibodies. When an electrical current is applied negatively charged proteins migrate towards the positive pole. The current is applied at 90° to the gel. Therefore proteins will move out of the gel and onto the membrane (figure 5.10). A description of the procedure follows below.

For Western blotting an immuno-P membrane was prewet in 100% methanol and afterwards immersed in H$_2$O. The immuno-P membrane and the gel from the SDS-PAGE were soaked in transfer buffer for 5 min. Additionally two Whatman filter pads were soaked in transfer buffer for 10 min. Afterwards the membrane sandwich was assembled as shown in figure 5.10 and air bubbles were removed with a pipette that was rolled over the membrane sandwich. A current of 15 V was applied for 45 min.

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{western blotting.png}
\caption{Assembly of membrane sandwich for Western blotting}
\end{figure}

Immunodetection

After completing SDS-PAGE and Western blotting proteins on the membrane can be detected using antibodies. The membrane was incubated with a primary antibody that specifically binds the protein of interest. The primary antibody was detected by reaction with a second antibody conjugated to an enzyme which catalyzes a light-emitting reaction. In this study the enzyme linked to the secondary antibody was an alkaline phosphatase which dephosphorylates the ECF™ substrate to produce a fluorescent product.

The first step in immunodetection was to wash the membrane in 5 ml TBS-Tween for about 5 min at room temperature. Blocking was carried out in 5% dry milk
in TBS-Tween for 1 h. The membrane was incubated with the primary antibody (table 5.1) over night at 4°C.

**Table 5.1.:** Overview over primary antibodies used for detection of phosphorylated Cdc2p and tubulin

<table>
<thead>
<tr>
<th>Primary antibody</th>
<th>Concentration</th>
<th>Protein size</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anti-cdc2-P antibody, rabbit</td>
<td>1:300</td>
<td>34 kDa</td>
</tr>
<tr>
<td>Anti- tubulin antibody, mouse</td>
<td>1:30 000</td>
<td>50 kDa</td>
</tr>
</tbody>
</table>

The antibody was removed and 3 consecutive washes were performed: 1x10 min and 2x5 min in 5 ml TBS-Tween. After that the membrane was incubated with the secondary antibody for 1 h at room temperature (table 5.2).

**Table 5.2.:** Overview over secondary antibodies which react with the primary antibodies

<table>
<thead>
<tr>
<th>Secondary antibody</th>
<th>Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alkaline phosphatase, anti-rabbit</td>
<td>1:5 000</td>
</tr>
<tr>
<td>Alkaline phosphatase, anti-mouse</td>
<td>1:10 000</td>
</tr>
</tbody>
</table>

The secondary antibody was removed and 3 consecutive washes were performed: 1x10 min and 2x5 min in 5 ml TBS-Tween. Then the membrane was carefully drained for excess TBS-Tween on a paper towel and laid on a glass plate. 1-1.5 ml ECF™ was poured over the membrane and spread to cover it completely. The reaction was stopped when bands became visible but not later than 5 min and was then dried in the dark between two filter papers. Fluorescent bands appeared after exposure to 540 to 560 nm-light.

The membrane can be stored at 4°C covered in foil.
Chapter 6.

Results

In order to investigate the role of Gcn2 in the regulation of G2/M, we first constructed *gcn2 rad* double mutants and then followed their progression into M after UVC irradiation in G2.

6.1. Making *gcn2 rad3* double mutants

<table>
<thead>
<tr>
<th>Strain</th>
<th>Referred to as</th>
<th>Genotype</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>1153</td>
<td><em>rad3</em>Δ</td>
<td><em>rad3::ura4+ leu1-32 ura4-D18 ade6-704 h</em>-</td>
<td>Lab collection</td>
</tr>
<tr>
<td>1137</td>
<td><em>gcn2</em>Δ</td>
<td><em>gcn2::ura4+ cdc10-M17 ura4-D18 h</em>+</td>
<td>Lab collection</td>
</tr>
<tr>
<td>1638</td>
<td><em>rad3</em>Δ<em>rad3</em>Δ</td>
<td><em>rad3::ura4+ cdc10-M17 ura4-D18 h</em>+</td>
<td>This work</td>
</tr>
</tbody>
</table>

Deletion of *gcn2* alone was not expected to show any effect on cell-cycle progression after UVC irradiation due to activation of the powerful DNA damage checkpoint. Therefore a double mutant was produced crossing a *rad3*Δ mutant with a *gcn2*Δ mutant (see table 6.1). Possible effects of Gcn2 after UVC irradiation on cell-cycle progression could now be seen more easily.

When crossing the *rad3*Δ strain with the *gcn2*Δ strain, recombination can give 32 possible genotypes of the progeny. The 32 genotypes are due to five different genes: *rad3, gcn2, cdc10, leu1-32* and *ade6-704*. Each gene has two possible versions: wild type or mutated ($2^5=32$). Among the 32 possible genotypes there is just one that we are interested in and will employ: *rad3::ura4+ gcn2::ura4+ cdc10-M17 ura4-D18* (*rad3*Δ*rad3*Δ mutant, see table 6.1). The strategy used to
distinguish between the 32 genotypes was to grow them on appropriate media. By comparing growth on the different plates it was possible to limit the number of progeny whose genotype was further analyzed by PCR. All the possible genotypes and their growth characteristics on the employed selective plates are summarized in table 6.2. The rationale behind the selection strategy is given below.

After crossing, the spores were allowed to form colonies on YE plates at 25°C. For genotype testing colonies from a YE plate were replica-plated on the following media:

- **EMM**
  to identify *ura4*+ colonies. This allows determination of progeny that has either the *gcn2*Δ or the *rad3*Δ or both.

- **YEP+HU**
  to identify the colonies that carry the *rad3*Δ. HU is an inhibitor of DNA synthesis (Enoch & Nurse, 1990; Elledge *et al*., 1992) and leads to incomplete DNA replication. *rad* mutants are deficient in the S/M checkpoint and will continue into mitosis and cell division despite a defective S phase. Exposed to HU *rad3*Δ cells show aberrant mitosis and sometimes anucleated cells. When exposed to HU cells lose viability which leads to increased cell death (Alkhodairy & Carr, 1992). This means that *rad3*+ grow in the presence of low concentrations of HU, while colonies with the mutated *rad3* are sensitive to HU and contain an increased number of dead cells. Phloxin B was used for better discrimination between dead and living cells and *rad3*Δ cells formed dark pink colonies and *rad3*+ cells were pale pink.

- **YEP, 36°C**
  to identify *cdc10-M17* colonies. The *cdc10* mutation confers temperature sensitive growth. Cells with the temperature sensitive *cdc10* gene can not survive at restrictive temperature (36°C), while wild type cells do. Also here phloxin B was used for easier discrimination of dead and living cells.

In summary, by comparing growth on the three different media, colonies that grew on EMM but not on YEP+HU or on YEP + 36°C were picked. As shown in table 6.2 this leaves just 2 out of 32 genotypes possible for the picked colonies. For further identification, the genotype of these colonies was determined by PCR.
Table 6.2.: The table shows growth of progeny on different media. There are 32 possible genotypes from this cross. Growth on EMM and YEP supplemented with hydroxyurea at 25°C and on YEP at 36°C characterizes the genotype.

<table>
<thead>
<tr>
<th>Genotype</th>
<th>EMM</th>
<th>HU</th>
<th>36°C</th>
</tr>
</thead>
<tbody>
<tr>
<td>rad3::ura4+ leu1-32 ura4-D18 ade6-704</td>
<td>x</td>
<td></td>
<td></td>
</tr>
<tr>
<td>rad3::ura4+ ura4-D18 ade6-704</td>
<td></td>
<td></td>
<td>x</td>
</tr>
<tr>
<td>rad3::ura4+ leu1-32 ura4-D18</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>rad3::ura4+ ura4-D18</td>
<td>x</td>
<td></td>
<td></td>
</tr>
<tr>
<td>rad3::ura4+ leu1-32 ura4-D18 ade6-704 cdc10M-17</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>rad3::ura4+ ura4-D18 ade6-704 cdc10M-17</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>rad3::ura4+ leu1-32 ura4-D18 cdc10M-17</td>
<td>x</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

rad3::ura4+ ura4-D18 cdc10M-17 |     |    |  |

gen2::ura4+ leu1-32 ura4-D18 ade6-704 | x   |   | x  |
| gen2::ura4+ leu1-32 ura4-D18 |     |   | x |
| gen2::ura4+ ura4-D18 ade6-704 | x |     |  |
| gen2::ura4+ ura4-D18 cdc10M-17 |     |    |  |
| gen2::ura4+ ura4-D18 ade6-704 cdc10M-17 |     |    |  |
| gen2::ura4+ ura4-D18 cdc10M-17 | x |     |  |
| gen2::ura4+ rad3::ura4+ leu1-32 ura4-D18 ade6-704 | x |   |  |
| gen2::ura4+ rad3::ura4+ ura4-D18 ade6-704 | x |     |  |
| gen2::ura4+ rad3::ura4+ leu1-32 ura4-D18 | x |     |  |
| gen2::ura4+ rad3::ura4+ ura4-D18 | x |     |  |
| gen2::ura4+ rad3::ura4+ leu1-32 ura4-D18 ade6-704 cdc10M-17 | x |   |  |
| gen2::ura4+ rad3::ura4+ ura4-D18 ade6-704 cdc10M-17 | x |     |  |
| gen2::ura4+ rad3::ura4+ leu1-32 ura4-D18 cdc10M-17 | x |     |  |
| gen2::ura4+ rad3::ura4+ ura4-D18 cdc10M-17 | x |     |  |

6.1.1. Polymerase Chain Reaction (PCR) of rad3 and gcn2

To further investigate the genotype of progeny from the gcn2Δ x rad3Δ cross, candidates with a potential rad3Δgcn2Δ genotype were picked and grown to saturation in liquid YE medium. Genomic DNA was isolated and two different PCRs were run, each with specific primers for either rad3 or gcn2 amplification. The PCR fragment of rad3Δ is expected to have a size of 1.8 kb while there is no product expected for the wild type of rad3 (see Chapter 5 - Methods: figure 5.3). DNA from a rad3Δ strain was used as a positive control. In figure 6.1 bands
with the expected size for rad3Δ of 1.8 kb are observed for both candidates. In addition, the right product size is confirmed by the positive control of a rad3Δ mutant having a band of the same size. PCR of a wild type strain does not show any bands which eliminates unspecific binding of the primers. Together with the results from replica plating on YEP+HU, this confirms that both candidates carry a rad3 deletion.

![Figure 6.1: Samples from putative rad3Δgcn2Δ double mutants, positive control (rad3Δ) and negative control (rad3+).](image)

Since there is no appropriate medium available to select for gcn2 mutants, identification of gcn2Δ was done by PCR alone. The obtained band of 1.5 kb seen on the gel for gcn2 PCR in figure 6.2 matches with the expected size of the gcn2 deletion product (see chapter 5-Methods: figure 5.1). For comparison, the size of wild type gcn2 is 4.8 kb. This suggests that candidates contain a gcn2 deletion as well as a rad3 deletion.
Both candidates show PCR products with the right size for rad3 and gcn2 deletion and generation of a rad3Δgcn2Δ double mutant is thereby confirmed. One of the two rad3Δgcn2Δ double mutant was used in further experiments.

6.2. Measuring cell-cycle progression using cdc10 block and release

We wished to compare the kinetics of entry into mitosis after UVC irradiation in rad3Δgcn2Δ to that in rad3Δ cells. For this purpose cells had to be synchronized and then irradiated in G2 phase. Synchronization was achieved using cdc10 block and release.

6.2.1. Cell-cycle analysis by flow cytometry

In order to follow cell-cycle progression, we used flow cytometry. Traditional flow cytometry analysis of the fission yeast cell cycle cannot distinguish between cells in G2, mitosis and G1 (2C versus 2x 1C). Therefore we have developed a new method (Knutsen et al., 2011) that enables us to distinguish 2C cells with one nucleus (G2 and early M) from cells with two nuclei (anaphase - G1). In order to validate the new method, we have compared mitotic indices measured by flow cytometry and by traditional staining techniques. My major contribution to the paper was these comparisons in cdc10 block-and-release experiments (figure 5 in...
Knutsen et al). The new method was used to first establish when the cells should be irradiated, then to follow entry into mitosis in wild type and mutant cells after UVC irradiation in G2.

### 6.2.2. Synchronization and timing of irradiation

Figure 6.3 below shows selected time points of *cdc10*<sup>ts</sup> cells after release. The DNA histograms show DNA content on the x-axis with 50 as the value for cells with one copy of each chromosome and 100 as the value for cells with two copies of each chromosome.

71% of wild type cells and 92% of *gcn2Δ* cells arrested at the G1 stage when incubated at 36°C (panel A). 120 min after release from the temperature block the majority of cells in both strains have two copies (2C) of each chromosome (panel B). 150 minutes after release the population of binucleate cells (BI) increased to 8% in wild type cells and 23% in *gcn2Δ* cells (panel C), showing entry into mitosis. In the cytogram showing DNA content-W (DNA content-width) versus DNA content-A (DNA content areal) it can be seen that most of the cells are in G2. Hence, an incubation time of 4 h at 36°C is sufficient to induce a good synchrony in both strains.
Figure 6.3: Flow cytograms of wt (489) and gcn2Δ (1138) strain at time points 0, 120 and 150 min after release from the 4h-temperature block.
We monitored the frequency of binuclear cells (BI) as a function of time after shift down from the temperature block. Samples were taken for 195 min. Analysis by flow cytometry shows that the frequency of binuclear cells of wild type and \( gcn2\Delta \) starts increasing at about 135 min after shift down from the temperature block (figure 6.4). This means that by 135-150 min cells enter mitosis. It can be noticed that the \( gcn2\Delta \) strain goes into mitosis faster than the wild type strain.

![Figure 6.4](image)

**Figure 6.4.** Binuclear index (BI) of wt (489) and \( gcn2\Delta \) (1138) strain monitored for 195 min after shift down from temperature block.

To confirm results measured by flow cytometry, the degree of synchrony was also analyzed using cell staining. Samples were taken as for flow cytometry but then stained with aniline blue to count septated cells. Septated cells appear concurrently with S phase and therefore later in the cell cycle than binuclear cells. Knowing that, the septation index can also be used to estimate entry into mitosis. In figure 6.5 wild type cells start septation at about 150 min after shift down. As seen for the BI in figure 6.4 \( gcn2\Delta \) cells show septa earlier than the wild type, at about 135 min after shift down from the temperature block.

Based on the graphs for the binuclear and the septation index, cells enter mitosis between 135 min and 150 min after release from the temperature block. Therefore cells were irradiated approximately 110 min after shift down from the \( cdc10 \) block to the permissive temperature to measure cell-cycle progression after UVC. At this time point both strains are still in G2 and we can expect that any G2/M
entry into mitosis after UVC irradiation

In this study cell-cycle progression of four different strains was compared (table 6.3).

Table 6.3.: Strains used in the experiments measuring cell-cycle progression.

<table>
<thead>
<tr>
<th>Strain</th>
<th>Referred to as</th>
<th>Genotype</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>489</td>
<td>wild type</td>
<td>cdc10-M17 h-</td>
<td>lab collection</td>
</tr>
<tr>
<td>1137</td>
<td>gcn2Δ</td>
<td>gcn2::ura4+ cdc10-M17 ura4-D18 h+</td>
<td>lab collection</td>
</tr>
<tr>
<td>1153</td>
<td>rad3Δ</td>
<td>rad3::ura4+ leu1-32 ura4-D18 ade6-704 h-</td>
<td>lab collection</td>
</tr>
<tr>
<td>1353</td>
<td>rad3Δ</td>
<td>rad3::ura4+ cdc10-M17 ura4-D18 h+</td>
<td>lab collection</td>
</tr>
<tr>
<td>1638</td>
<td>gcn2Δ rad3Δ</td>
<td>gcn2::ura4+ rad3::ura4+ cdc10-M17 ura4-D18</td>
<td>this study</td>
</tr>
</tbody>
</table>

Exponentially growing cdc10\textsuperscript{ts} cells were synchronized in G1 phase and exposed to UVC-light in G2 phase. After irradiation cell-cycle progression was measured by flow cytometry. Samples were taken immediately after irradiation and then every 15 min.

Analysis by flow cytometry (figure 6.6) shows that the frequency of binuclear cells increases in untreated samples after the time point of irradiation. All strains continue the cell cycle and enter mitosis as expected. By 75 min after the time of irradiation most of the untreated cells in all strains had passed mitosis having...
more than 60% binuclear cells. When irradiated with UVC the frequency of binuclear cells of wild type and \textit{gcn2}Δ cells does not increase significantly. The BI of both strains is below 15% throughout the whole experiment. We conclude that after treatment with UVC these two strains delay mitosis. The \textit{rad3}Δ strain and the double mutant strain \textit{rad3}Δ\textit{gcn2}Δ show continous increase in the BI after UVC treatment. 75 min after irradiation about 40% of the cells had passed mitosis. The increase in binuclear cells of the \textit{rad3}Δ and the \textit{rad3}Δ\textit{gcn2}Δ strain is significantly slower than in the untreated cultures. This result is surprising since the \textit{rad3}Δ mutant is checkpoint deficient (Bentley et al., 1996) and was expected to enter mitosis without any major delay.

6.2.4. Cell-cycle progression monitored by fluorescent microscopy

To further test the new flow cytometry-based method and confirm the conclusions drawn above, we monitored cell-cycle progression of two of the above strains (wild type and \textit{rad3}Δ\textit{gcn2}Δ) using a fluorescent staining technique. Samples were taken at the same time points as for flow cytometry but prepared for analysis with DAPI staining. Figure 6.7 shows the kinetics of entry into mitosis as measured by DAPI staining for UVC-treated and untreated samples of wild type and \textit{rad3}Δ\textit{gcn2}Δ strain. The frequency of mitotic cells determined using DAPI staining (mitotic index, MI) is compared with BI obtained using flow cytometry.

The graphs in figure 6.7 show that both methods to estimate number of mitotic cells give the same result. The wild type strain delays mitosis at least
Figure 6.7: Comparison of checkpoint response after UVC irradiation in wild type (wt) and rad3Δ gcn2Δ double mutant. The cells were synchronized using cdc10 block and release and irradiated with UVC in G2. For graphs with open symbols binuclear cells were determined by DAPI staining, for closed symbols flow cytometry was used.

75 min after irradiation with UVC. The slow, gradual entry into mitosis of the rad3Δ gcn2Δ double mutant is observed both when analyzing samples by flow cytometry and fluorescent microscopy.

We also employed immunofluorescent staining of β-tubulin as a third method to measure progression into mitosis. Here we compare entry into mitosis of wild type and rad3Δ.

In the beginning of mitosis the microtubules are reorganized to form the spindle apparatus which facilitates the distribution of the chromosomes to the daughter cells. We have stained β-tubulin using indirect immunofluorescence to visualize the microtubules. The mitotic spindle can be clearly distinguished from the interphase microtubule array (see methods). In the figure shown below the mitotic index was determined counting mitotic cells stained with a fluorescently-labeled tubulin antibody.

We found that already 35-40% of cells from rad3Δ-strain were in mitosis directly after irradiation. In contrast, there were only 0-5% mitotic cells observed in the wild type strain. The maximum of mitotic cells was reached 15 min after irradiation for the rad3Δ-strain, while wild type cells had a maximum of mitotic cell 45 min after irradiation. This indicates that cells with a rad3 deletion complete S phase considerably earlier than wild type cells and were therefore irradiated when they had already passed any G2/M checkpoint. While the current
6 RESULTS

Figure 6.8.: Comparison of checkpoint response after UVC irradiation in wild type (wt) and rad3Δ mutant. The cells were synchronized using cdc10 block and release and irradiated with UVC in G2. The MI was determined by counting cells with a mitotic spindle.

experiments were in progress, other results in the lab indicated that rad3Δ cells enter S phase earlier than the wild type cells after release from a cdc10 block (Cathrine Arnason Bøe, personal communication). Therefore we decided to employ a selection synchronization method in further experiments.

6.3. Measuring cell-cycle progression using a selection method for synchronization

To further investigate cell-cycle progression of the rad3Δ strains we chose lactose gradient synchronization, a method which does not disrupt the cell cycle to the same extent as the induction method. This method is based on isolating the smallest cells from the population. Due to the short G1 (see Theoretical background 2.3), this results in a population of early-G2 cells. Therefore irradiation was performed immediately after synchronization.

6.3.1. Cell-cycle progression monitored by flow cytometry

As for cells synchronized with block and release, cell-cycle progression was monitored by flow cytometry to measure the frequency of binuclear cells. The binuclear index shows that all strains of the untreated samples have entered mitosis 30-45 min after the irradiation time point. Irradiated cells from all strains show a delay and do not enter mitosis until 75 min after irradiation with UVC. Consistent with the previous experiments using cdc10 block and release, these
results indicate that Gcn2 does not have any effect on G2/M transition after UVC irradiation. Surprisingly, synchronized with the lactose gradient the rad3Δ strains showed a clear delay into mitosis, with the same kinetics as the wild type strain.

As Rad3 is an important component in the signaling pathway of the DNA damage checkpoint, a deletion of rad3 was thought to result in a checkpoint-deficient mutant. Thus, the cell cycle delay of the rad3Δ strain upon UVC irradiation was unexpected in light of the general view in the field. We decided to pursue this observation and the main focus of this work shifted from exploring the role of Gcn2 to exploring the delay induced in the absence of Rad3.

6.4. The role of Rad3 after UVC irradiation

6.4.1. Confirming the rad3Δ strain

Having made the surprising observation that the rad3Δ strain delays mitosis after UVC irradiation, we considered the possibility that the strain has picked up a suppressor and/or there was a mistake in the main lab collection. First, we verified that the strain carries the rad3Δ by PCR. The strain clearly carries a deletion of rad3, since we could amplify the band specific for the deletion (figure 6.10).
Then we checked other, well-studied phenotypes of the rad3Δ mutant, namely entry into mitosis in the presence of HU and survival rates after UVC treatment. Consistent with the results of Bentley et al. (1996) we achieve a low survival of the rad3Δ strain when irradiating with a dose of 1 100 J/m² of UVC in G2. This dose gives a survival rate of 100% in the wild type strain, but only 7% in the rad3Δ strain (Gro Elise Rødland, personal communication).

After HU treatment, wild type cells arrest in early S phase and elongate. In contrast, the rad3Δ cells are known to enter mitosis with unreplicated chromosomes, which results in the so-called cut phenotype. As the cell tears apart unreplicated chromosomes, uneven chromosome segregation is typical for such mitoses. We compared wild type and rad3Δ cells after treatment with HU (figure 6.11). rad3Δ cells show newly divided cells with septa and a clearly visible cut phenotype. In contrast the wild type strain has elongated cells, arrested with only one nucleus after the HU treatment. Septa could not be observed in the wild type strain. We conclude that the rad3Δ strain is the right strain as judged by PCR and that it behaves as described in the literature after treatment with HU (Enoch et al., 1992).
6.4.2. Cell-cycle progression of rad3Δ monitored by immunoblotting

The binuclear index measures the number of cells in anaphase, G1 and S. It is therefore unclear whether the rad3Δ cells delay in G2 prior to entry into mitosis or in early mitosis, such as metaphase. In order to investigate more closely where in the cell cycle the rad3Δ cells delay, we followed the presence of the Y15 phosphorylated form of Cdc2 by immunoblotting, using a phosphospecific antibody. Dephosphorylation of Cdc2-Y15 is a hallmark event for entry into mitosis. Protein samples were taken at time points 0, 80 and 100 min after irradiation. Since lactose gradient synchronization only provides a limited amount of synchronous cells, it was not possible to take samples more often.

Cells of both strains, rad3Δ and wild type, show phosphorylated Cdc2-Y15 at time point 0 consistent with cells being in G2. We expected a decrease in the intensity of the phosphorylated Cdc2 band at 80 minutes, but this is difficult to see in this experiment. At 100 min after irradiation control samples show again phosphorylated Cdc2-Y15.
For samples irradiated with UVC, both rad3Δ and wild type show an increase in phosphorylated Cdc2-Y15 at 80 min compared to control samples suggesting that the cells arrest in G2 phase. At 100 min after UVC treatment the wild type strain shows Cdc2-Y15 phosphorylated suggesting a continued delay. Due to uneven loading it is difficult to conclude as to the length of the delay in the rad3Δ strain, but it appears somewhat less prominent than that in wild type cells.

From these rather preliminary results we conclude that the rad3Δ cells delay in G2 after UVC irradiation in early G2 in the same way as wild-type cells do, but the length of the delay might be somewhat shorter in the absence of Rad3.

Figure 6.12.: Immunoblots probed with antibodies against phosphorylated Cdc2 and against tubulin as loading control showing samples from UVC irradiated wild type (wt) and rad3Δ mutant.
Chapter 7.

Discussion

Gcn2 and its role in the G2/M transition

In 2007 Tvegard et al. reported that UVC activated a Gcn2-dependent checkpoint in G1. Since activation of Gcn2 has also been observed in G2 phase (Krohn et al., 2008) we investigated whether Gcn2 has any effect on cell-cycle progression from G2 phase into mitosis. In order to see a possible minor effect of Gcn2 on the G2/M transition, we constructed a rad3\Delta gcn2\Delta double mutant and monitored cell-cycle progression after UVC irradiation by flow cytometry to measure the frequency of binuclear cells. We have shown that the kinetics of entry into mitosis after UVC irradiation of rad3\Delta and rad3\Delta gcn2\Delta cells were the same. This result is independent of the synchronization method we used (figure 6.9). Based on the literature and the general view in the field, we expected the rad3\Delta strain to enter mitosis directly after irradiation with UVC in G2 but could not observe this when synchronizing with cdc10 block and release. We suspected that the induction method we used might have affected the cell-cycle progression of the strains. Lactose gradient synchronization is a more gentle synchronization method and does not perturb cell-cycle progression to the same extent as synchronization with cdc10. Moreover this selection method was also used by Bentley et al. (1996) in their experiments characterizing Rad3. Therefore we decided to use lactose gradient synchronization in further experiments. However, the kinetics of entry into mitosis was similar between the rad3\Delta and rad3\Delta gcn2\Delta strains also when lactose gradient centrifugation was used to synchronize the cells. While Gcn2 is clearly required for the G1/S checkpoint, we show here that it is not required for the regulation of G2/M after UVC irradiation.

The role of Gcn2 in the G1 phase is not totally elucidated yet. We know that it is required for translation downregulation and for the G1/S checkpoint response.
However, it is not clear if and how these two functions of Gcn2, translation downregulation and checkpoint activation, are connected and how they lead to the inhibition of S phase entry. One possibility is that Gcn2 acts directly on a cell cycle protein. In this scenario, Gcn2 might phosphorylate a G1-specific cell cycle protein. Alternatively, the effect on the cell cycle at G1/S might derive from Gcn2’s effect on translation downregulation. In this scenario the difference between the roles of Gcn2 in G1 versus G2 might stem from different requirements for regulating translation of specific mRNAs that are necessary in G1 but not in G2.

Another observation was that the $gcn2\Delta$ mutant cells enter mitosis about 15 min earlier than wild type cells, using $cdc10$ block-and-release (Fig 6.4-6.6). However, the early entry into mitosis is lost in irradiated $gcn2\Delta$cells (figure 6.6). It is difficult to find a plausible explanation for why the $gcn2\Delta$ enters mitosis earlier than the wild type strain, but it fits with our observation that under some growth conditions the $gcn2\Delta$ strain seems to have a slightly shorter generation time than the wild type strain (Beata Grallert, unpublished observation). As this enhancement of cell-cycle progression is clearly seen first when measuring the entry into mitosis but not in earlier cell cycle stages, one possibility is that the $gcn2\Delta$ mutant might be slightly faster in several or every cell cycle phase. We cannot exclude that Gcn2 has a subtle role also in the regulation of G2/M, but we can conclude that it has no major role in the regulation of G2/M after UVC irradiation in G2.

**A Rad3-independent delay after UVC irradiation in G2**

In the course of this work we observed that $rad3\Delta$ cells delayed mitosis in the same way as wild type cells do upon UVC-irradiation in G2 phase. We confirmed that this observation is made on a bona fide $rad3\Delta$ strain, which does have the deletion as judged by PCR. Our $rad3\Delta$ strain otherwise behaves as previously shown (Enoch et al., 1992): it is sensitive to HU showing small septated cells with unevenly distributed chromosomes (cut phenotype) and has a low survival after UVC treatment. We therefore conclude that the $rad3\Delta$ strain is correct and that it did not pick up a suppressor.

This finding was most surprising as it contradicts the general view in the field that assumes that $rad3\Delta$ cells are checkpoint deficient after UVC irradiation.
This view is based on the findings of Bentley et al. (1996) who showed that \textit{rad3}\textDelta mutants are checkpoint deficient after IR treatment and have low survival rates after UVC irradiation. To our knowledge checkpoint deficiency has not been shown after UVC irradiation, only generally assumed based on the profound UVC sensitivity and the established checkpoint deficiency after IR. As we show that \textit{rad3}\textDelta cells do delay mitosis (either entry into or in early M, see below), we have to conclude that (i), an alternative pathway is activated that prevents mitosis in \textit{rad3}\textDelta cells and (ii), Rad3 has an important, checkpoint-independent function(s) required for survival after UVC irradiation.

Cell-cycle progression measured by DAPI-staining and flow cytometry determines the frequency of cells that have entered anaphase since mitotic cells are first detected when chromosomes separate and two distinct nuclei appear. From those results we can therefore not conclude whether \textit{rad3}\textDelta cells stand in G2 phase before entering mitosis or in metaphase in early mitosis. To investigate where cells arrest we employed immunoblotting to look at the phosphorylation state of Cdc2 after UVC irradiation in G2 since dephosphorylation of Cdc2 is a prerequisite for entry. We observed that at 80 minutes after irradiation Cdc2 accumulates in its phosphorylated form in both strains as compared to unirradiated control samples. At this time point \textit{rad3}\textDelta cells show a clear arrest when monitoring cell-cycle progression with DAPI-staining and flow cytometry. This results indicate that cells are arrested in G2 rather than in metaphase. However, this experiment is hampered by technical difficulties in that we cannot obtain enough synchronous cells from lactose gradients to follow the kinetics of Cdc2 dephosphorylation more closely. Therefore further studies are needed to confirm this result. Another approach would be to identify mitotic cells by tubulin staining. If \textit{rad3}\textDelta cells do not show the mitotic spindle after UVC irradiation, this would be another evidence for an arrest in G2 phase.

If the cell-cycle delay following UVC-irradiation is independent of Rad3, this raises the question which other proteins might be responsible for the delay.

In mammals two functionally distinct pathways depending on the type of DNA damage exist. Either of two protein kinases, ATM or ATR, initiates the signaling pathway (Alberts \textit{et al.}, 2008). ATR, the human homolog of Rad3, is activated by UVC-induced DNA damage, while ATM, the human homolog of Tel1, plays an
important role after DNA double strand break. However, until now no checkpoint role of Tel1, the ATM homologue in *S. pombe*, has been described. So far the only known role of Tel1 is in telomere maintenance in fission yeast. To investigate the possible requirement for Tel1 in the G2 delay of *rad3*Δ cells, we are currently constructing a *rad3*Δ*tel1*Δ double mutant. Cell-cycle progression of *rad3*Δ*tel1*Δ after UVC irradiation in G2 will give information about whether Tel1 is involved in a checkpoint.

Previously a study in budding yeast demonstrated that the role of Tel1 in checkpoint signaling is not limited to mammals (Clerici *et al.*, 2004). They showed that when *MEC1*Δ cells are irradiated in G1, they delay entry into mitosis. The delay depends on both Tel1 and components of the spindle assembly checkpoint (SAC). The activation of Tel1 in the absence of Mec1 led to the phosphorylation of the downstream effector kinase Chk1. Since activation of Tel1/ATM after DNA damage is observed in both human and budding yeast, Tel1 might have a role in checkpoint activation in *S. pombe* as well.

As for the requirements for the SAC after UVC irradiation in G1, it is important to note that budding yeast has the unique feature of assembling a short mitotic spindle already at the beginning of S phase. Thus irradiation in G1 might affect multiple components leading to activation of the SAC, possibly due to defects in the spindle pole body (SPB) and its duplication, kinetochores, and/or centromeres.

We irradiated our cells in G2 when duplication of the SPB is already completed, but the cells have not yet formed the mitotic spindle. As described above our preliminary results suggest that UVC-irradiated *rad3*Δ cells delay in G2 and not in mitosis. Therefore we think it unlikely that the SAC is responsible for the cell-cycle delay observed in *rad3*Δ cells.

An alternative candidate responsible for the mitotic delay of *rad3*Δ cells is Sty1. The Sty1 kinase is a MAP(mitogen-activated protein) kinase activated by various environmental stresses such as heat, oxidative stress and heavy metals but also by DNA damaging agents such as UVC or IR treatment. Rad3 and Sty1 share some of the same targets which include genes in the stress responses mentioned above (Alao & Sunnerhagen, 2008). Interestingly, Srk1 kinase, one of the proteins regulated by Sty1, is activated upon osmotic stress and causes phosphorylation
of Cdc25 at the same site as where Chk1 phosphorylates Cdc25, leading to cell cycle arrest (Lopez-Aviles et al., 2005). Since Sty1 is activated by UVC light, it is tempting to think that a subsequent activation of the Sty1 target, Srk1, might be a mechanism that could keep rad3Δ cells arrested in G2 after UVC. In order to test the role of Srk1 and/or Sty1 in the cell cycle delay of rad3Δ cells, entry into mitosis could be monitored in rad3Δsty1Δ and rad3Δsrk1Δ mutants.

Checkpoint arrest is important for cell survival after DNA damage because it allows time for the induction of repair mechanisms that eliminate the damage. rad3Δ cells have a low survival after UVC treatment in spite of arresting at G2/M, leading us to speculate that Rad3 might have a role in DNA repair. It has been shown that Rad3 homologs in humans and budding yeast (ATR and Mec1, respectively) co-localize with replication protein A (RPA) at sites of DNA damage (Wu et al., 2005; Nakada et al., 2005). RPA is important in many DNA repair pathways including double strand brake repair by homologous recombination (HR). Following DNA damage RPA is phosphorylated by Phosphoinositide 3-kinase-related kinases including ATR and interacts with DNA repair proteins Rad51 and Rad52 (Binz et al., 2004). Rad51 and Rad52 are important for homologous recombination which is the predominant repair pathway in G2. As DNA repair and checkpoint pathways are tightly coupled through RPA interacting both with Rad51/52 and ATR, a deletion of ATR might have an influence on DNA repair. Furthermore, ATR has been shown to phosphorylate XPA, an important component of the nucleotide excision repair (NER) pathway. Phosphorylation of XPA results in its increased nuclear localization and increased repair activity (Wu et al., 2007; Shell et al., 2009). Inefficient repair by HR and NER might well explain the profound UVC sensitivity of the rad3Δ mutant.
Appendix A.

Molecular weight standards
Bibliography


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Christiane Rothe

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