REPORT

Conformation-specific anti-Mad2 monoclonal antibodies for the dissection of checkpoint signaling

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ABSTRACT
The spindle assembly checkpoint (SAC) ensures accurate chromosome segregation during mitosis by delaying the activation of the anaphase-promoting complex/cyclosome (APC/C) in response to unattached kinetochores. The Mad2 protein is essential for a functional checkpoint because it binds directly to Cdc20, the mitotic co-activator of the APC/C, thereby inhibiting progression into anaphase. Mad2 exists in at least 2 different conformations, open-Mad2 (O-Mad2) and closed-Mad2 (C-Mad2), with the latter representing the active form that is able to bind Cdc20. Our ability to dissect Mad2 biology in vivo is limited by the absence of monoclonal antibodies (mAbs) useful for recognizing the different conformations of Mad2. Here, we describe and extensively characterize mAbs specific for either O-Mad2 or C-Mad2, as well as a pan-Mad2 antibody, and use these to investigate the different Mad2 complexes present in mitotic cells. Our antibodies validate current Mad2 models but also suggest that O-Mad2 can associate with checkpoint complexes, most likely through dimerization with C-Mad2. Furthermore, we investigate the makeup of checkpoint complexes bound to the APC/C, which indicate the presence of both Cdc20-BubR1-Bub3 and Mad2-Cdc20-BubR1-Bub3 complexes, with Cdc20 being ubiquitinated in both. Thus, our defined mAbs provide insight into checkpoint signaling and provide useful tools for future research on Mad2 function and regulation.

KEYWORDS
APC/C; Cdc20; conformational specific; kinetochore; Mad2; SAC

Introduction
Proper partitioning of sister chromatids to the 2 new daughter cells is one of the most important aspects of mitosis. This outcome is ensured by the correct attachment of kinetochores to microtubules of the mitotic spindle resulting in biorientation of the sister chromatids and their subsequent partitioning at anaphase.1,2 Unattached kinetochores will result in the activation of the spindle assembly checkpoint (SAC) that inhibits anaphase entry until proper biorientation of all sisters have been established.3,5 SAC activation results in the generation of the mitotic checkpoint complex (MCC) that inhibits Cdc20, the mitotic co-activator of the anaphase-promoting complex/cyclosome (APC/C). The MCC is composed of the checkpoint proteins Mad2 and BubR1-Bub3 that bind directly to Cdc20.6-11 The MCC binds stably to the APC/C, and recent work shows that the MCC can bind and inhibit a second molecule of Cdc20 already bound to the APC/C, which is required for checkpoint signaling.12 The rate-limiting step in the formation of the MCC is the kinetochore-catalyzed binding of Mad2 to Cdc20 because Mad2 has to undergo a large structural change for this to occur.13 Once the Mad2-Cdc20 complex has formed, BubR1-Bub3 can bind to Cdc20, and subsequently Mad2 might dissociate to form a Cdc20-BubR1-Bub3 complex that has been referred to as the BBC complex.10,14-16 The removal of Mad2 from Cdc20 is stimulated by p31comet and the AAA-ATPase TRIP13, but the exact details of this are still to be determined.14,17-22

The Mad2 protein almost exclusively consists of a HORMA domain that can adopt 2 different conformations referred to as open-Mad2 (O-Mad2) and closed-Mad2 (C-Mad2).23-27 The conversion from O-Mad2 to C-Mad2 requires a large movement of both the N terminus and the C terminus of Mad2 (Fig. 1A). C-Mad2 is the active form of Mad2 that binds to a short sequence in the N terminus of Cdc20, and, in the Mad2-Cdc20 complex, the C terminus of Mad2 crosses Cdc20 like a seatbelt.24 Interestingly, a similar sequence is present in the Mad1 checkpoint protein that also binds C-Mad2 in a structurally similar way.26 The Mad1-Mad2 complex is a very stable complex and localizes to unattached kinetochores through Mad1 interactions with outer kinetochore proteins.28-33 Given the fact that C-Mad2 can dimerize with O-Mad2, the template model proposes that kinetochore-localized Mad1-Mad2 recruits soluble O-Mad2 to kinetochores and this converts O-Mad2 into C-Mad2.34 This model is supported by many elegant biochemical experiments but data also suggest that unknown aspects of the kinetochore environment could contribute to
Figure 1. Characterization of Mad2 monoclonal antibodies. (A) Structure of O-Mad2 and C-Mad2 with the C- and N-terminus colored in green and red, respectively. In the C-Mad2 structure, a ligand (orange) is bound mimicking the binding of Cdc20 and Mad1. O-Mad2 modified from PDB 1DUJ and C-Mad2 modified from PDB 1KLQ. (B) ITC measurements using purified Mad2 monoclonal antibodies and recombinant Mad2 L13A/R133A (C-Mad2) and Mad2 V193N/R133A (O-Mad2) to determine binding specificity and affinity of the antibodies. (C) Stable HeLa cell lines expressing Mad2-Venus, Mad2 L13A-Venus or Mad2 V193N-Venus or a control cell line (FRT) were treated with nocodazole and cells were harvested by mitotic shake-off. A mitotic extract was prepared from each cell line and Mad2 complexes were immunoprecipitated with the different Mad2 monoclonal antibodies. Samples were subsequently analyzed by SDS-PAGE and western blot and probed for the indicated proteins. (D) Immunofluorescence images of cells stained with the indicated Mad2 conformational specific antibodies. Cells were arrested in prometaphase using nocodazole or left untreated to obtain interphase cells.
To further understand Mad2 function in vivo, it is critical to have well-defined tools to probe the different Mad2-containing complexes that are formed during an active checkpoint. A mouse monoclonal C-Mad2-specific antibody has been generated by the Nigg lab, but mouse monoclonal O-Mad2 and pan-Mad2 antibodies have not been reported. Here, we describe 3 novel Mad2 mouse monoclonal antibodies (mAbs) that recognize C-Mad2, O-Mad2 and pan-Mad2, and use these to probe Mad2 complexes during mitosis.

**Results**

**Characterization of conformation-specific anti-Mad2 mAbs**

To investigate Mad2 biology in vivo, we aimed at generating Mad2-specific mAbs to address more precisely the composition of checkpoint complexes. Mice were immunized with bacterially produced His-tagged full-length Mad2, which contains a mixture of different Mad2 conformations. Following fusion with hybridoma cells, single clones were isolated and screened by ELISA for reactivity toward recombinant Mad2. Supernatants from positive clones were subsequently screened for their ability to immunopurify Mad2 from mitotic cells, which resulted in 3 positive clones (clone numbers 32, 157, 177) that were characterized further.

First, we used isothermal titration calorimetry (ITC) to measure the affinity of the antibodies for a mutant of Mad2 locked in its closed conformation (Mad2 L13A) or a mutant locked in its open conformation (Mad2 V193N) (see ref 39 for characterization of these recombinant Mad2 mutants). In addition, both Mad2 mutants harbored the R133A mutation to avoid any chance of Mad2 dimerization. All three antibodies bound with nanomolar (nM) affinity to Mad2 with a molar ratio of 2, but there was a clear difference in their recognition of the different Mad2 conformations (Fig. 1B and Fig. S1A). Clone 32 bound very specifically to O-Mad2 while clone 157 bound very specifically to C-Mad2. In contrast, clone 177 bound tightly to both O-Mad2 and C-Mad2. None of the antibodies recognized Mad2 by western blot (data not shown), clearly suggesting that they were recognizing structural rather than linear epitopes.

We further analyzed the specificity of the 3 antibodies in immunopurifications from mitotic HeLa cells stably expressing Mad2 WT-Venus, Mad2 L13A-Venus and Mad2 V193N-Venus (Fig. 1C). Cells were arrested with nocodazole, a microtubule-depolymerizing agent, to activate the checkpoint and then harvested by mitotic shake-off. From the mitotic cell extracts, we immunopurified Mad2 with the different antibodies and analyzed samples by protein gel blot. Clone 177 bound all 3 forms of exogenous Mad2 and co-purified known Mad2 interactors, which is in agreement with the ITC measurements. Clone 32 only precipitated Mad2-Venus and Mad2 V193N-Venus, while clone 157 precipitated Mad2-Venus and Mad2 L13A-Venus but not Mad2 V193N-Venus. Thus, the specificity of the antibodies determined by ITC was confirmed by these immunopurification experiments.

The Mad1-Mad2 complex localizes to the nuclear pores in interphase and to kinetochores during mitosis; we thus analyzed the ability of the different antibodies to detect Mad2 localization. Cells treated with nocodazole were stained with a centromere marker (CREST) and the different Mad2 antibodies and analyzed by immunofluorescence. Clone 157 specifically stained the kinetochore of mitotic cells and stained the nuclear envelope of interphase cells, while clone 32 did not stain any cellular structures (Fig. 1D). Clone 177 stained the kinetochores, but the signal was not as strong as for clone 157.

Based on these tests, we conclude that clone 177 is a pan-Mad2-specific antibody, clone 157 is a C-Mad2-specific antibody and clone 32 is an O-Mad2-specific antibody. We use these names rather than clone numbers in the sections below.

**Analysis of Mad2 complexes in mitotic cells**

To explore the different complexes formed by Mad2 during mitosis, we precipitated endogenous Mad2 from nocodazole-arrested cells with the 3 different antibodies and analyzed their ability to co-purify known interactors (Fig. 2A). The pan-Mad2 antibody co-purified all known Mad2 interactors efficiently suggesting that the epitope recognized by this antibody does not interfere with Mad2 interactions. In contrast, the C-Mad2 antibody strongly co-purified Cdc20 and Mad1, but very little BubR1, APC/C and p31comet. As p31comet and BubR1 binds the dimerization surface of C-Mad2, we suspect that the C-Mad2 antibody is recognizing a similar region on Mad2. This was also reported for the C-Mad2-specific antibody generated by the Nigg lab and a different C-Mad2 clone we have described, potentially reflecting the dimerization surface as a strong antigen in C-Mad2. Although the O-Mad2 antibody was specific for O-Mad2 in our previous tests, we did detect some binding to known C-Mad2 interactors, such as Cdc20 and APC/C components. This could either reflect O-Mad2 interaction with C-Mad2 in these complexes or the weak affinity of the antibody for C-Mad2 that we can detect by ITC (Fig. 1B). We note that a sheep polyclonal antibody produced by the Taylor lab might also be a O-Mad2-specific antibody, and this antibody also co-purifies small amounts of Cdc20 and Mad1. Since the O-Mad2 antibody did not purify any Mad2 L13A-Venus from cells, we believe that this antibody is specific for O-Mad2 under our standard immunoprecipitation (IP) conditions. To further analyze if O-Mad2 can associate with checkpoint complexes, most likely through dimerization with C-Mad2, we purified APC/C complexes from nocodazole-arrested cells stably expressing Mad2 L13A-Venus and Mad2 V193N-Venus well below the endogenous Mad2 levels (Fig. 2B-C). Following peptide release of APC/C complexes from an APC4 affinity column, the samples were subjected to a second purification using a Venus affinity resin. In these experiments, APC/C and MCC components were co-purified by both Mad2 L13A-Venus and Mad2 V193N-Venus. The supernatant remaining after the APC4 depletion was also subjected to purification with a Venus affinity resin and again we could detect some APC/C and MCC components in both Mad2 purifications (Figs. 2B–C). Importantly, only Mad2 L13A-Venus co-purified p31comet as we expected. Based on this, we favor that the reason why our O-Mad2 antibody can co-purify small amounts of APC/C and checkpoint proteins is due to dimerization of O-Mad2 with C-Mad2 in these complexes.
Mass spectrometry analysis of Mad2 complexes in checkpoint-arrested cells

To investigate, in an unbiased manner, the proteins interacting with the different conformations of endogenous Mad2 during an active checkpoint, we used SILAC quantitative mass spectrometry. HeLa cells were grown in the presence of different isotopes of Arg and Lys to obtain 3 different populations (light (L), medium (M), heavy (H)), the peptides of which can be distinguished in the mass spectrometer (Fig. 2D). The heavy population was arrested in mitosis with nocodazole and the medium population was in addition incubated with the Mps1 inhibitor reversine for one hour to silence the SAC. Cells treated with reversine were also treated with the proteasome inhibitor MG132 to avoid mitotic exit. The light population was arrested with nocodazole and a mouse IgG was used for control purification. We used the 3 Mad2-specific mAbs to purify Mad2 from the heavy and medium conditions to get a

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**Figure 2.** Analysis of Mad2 complexes using conformation specific antibodies. (A) HeLa cells arrested in mitosis with nocodazole were harvested by mitotic shake-off and Mad2 complexes purified with the different Mad2 monoclonal antibodies. Representative of 3 independent experiments. (B) Schematic of the sequential purification of Mad2-Venus complexes. (C) HeLa FRT/Trex cells stably expressing Mad2 L13A-Venus or Mad2 V193N-Venus were collected by mitotic shake-off after nocodazole treatment. APC/C complexes were purified with a monoclonal APC4 antibody and then eluted by the antigenic peptide. The eluate containing APC/C and the supernatant after APC/C depletion was then used for purification of Mad2-Venus complexes using GFP-Trap beads (chromotek) and analyzed by protein gel blot. Input is total cell extract prior to APC4 purification. Representative of 2 independent experiments. (D) Schematic of the triple SILAC experimental set-up. L: light condition, M: medium condition, H: heavy condition. (E) SILAC ratios determined for known Mad2 interactors in the SAC ON and SAC OFF condition. Only known interactors having a SILAC ratio higher than 2 are included except for TRIF13, which is included despite not being enriched above IgG control levels. (F) Peptide intensities for Mad2, Mad1 and Cdc20 in the indicated conditions to illustrate that the O-Mad2 antibody purifies much lower levels of Mad1 and Cdc20.
quantitative comparison of “SAC ON” and “SAC OFF” complexes and referenced this to the IgG control (Fig. 2D). In agreement with the above analysis, the pan-Mad2 antibody enriched all known Mad2 interactors, and there was a clear reduction in Cdc20, BubR1, Bub3 and APC/C components when the checkpoint was turned off (Fig. 2E). The association with Mad1 did not change much as expected and similarly p31comet association with Mad2 was not responsive to SAC status in agreement with p31comet constantly antagonizing Mad2 function.4,20 In contrast, the C-Mad2 and O-Mad2 antibodies mainly precipitated Cdc20 and Mad1, and again Cdc20 showed the expected reduction in the SAC OFF condition. It should be noted that the intensities for Cdc20 and Mad1 was highest in pan-Mad2 and C-Mad2 purifications and much lower in O-Mad2 purifications, as expected (Fig. 2F). Our pan-Mad2 purification also validated Kif20A as an endogenous Mad2 interactor45 but TRIP13 was not enriched above background in any of these purifications, likely reflecting a transient interaction of this protein with checkpoint complexes.

When combined, the mass spectrometry analysis supports current models of Mad2 checkpoint complexes and their dynamics.

**Distribution of Mad2 in different complexes**

The above analysis provided a detailed analysis of Mad2 interactors but it did not enable the separation of different checkpoint complexes containing Mad2. To do this, we combined size-exclusion chromatography and the pan-Mad2 antibody. A nocodazole extract was prepared and separated on a Superdex 200 column, and from each fraction we immunoprecipitated Mad2 using the pan-Mad2 antibody and analyzed these by western blot (Fig. 3). As expected for a pan-Mad2 antibody, the migration profile of immunoprecipitated Mad2 resembled the total Mad2 profile in the extract. Two distinct populations of Mad2 were evident, one peaking around 1000 kDa and another around 25 kDa, likely corresponding to free Mad2. The Mad2 population at 1000 kDa represented approximately 20% of the Mad2 and precipitated MCC components, Mad1 and APC/C explaining their large Stokes radius. The Mad2 migrating at 25 kDa corresponds to monomeric Mad2, and we also detected some p31comet-Mad2 complex migrating around 50 kDa. The presence of a p31comet-Mad2 complex suggests that C-Mad2 can exist unbound to its known ligands Mad1 and Cdc20.

When we purified O-Mad2 and C-Mad2 from the fractions, there was a clear difference between Mad2 migration in the input and Mad2 enrichment in the immunopurifications. The O-Mad2 antibody primarily precipitated Mad2 in low molecular weight complexes and a small fraction in high molecular weight complexes. In contrast, the C-Mad2 antibody largely precipitated Mad2 in the high molecular weight complexes and some Mad2 migrating at 25 kDa on the column. This is in agreement with C-Mad2 being the active form binding to MCC and Mad1 but it indicated that unliganded C-Mad2 might be present during an active checkpoint. We cannot, however, exclude antibody induction of O-Mad2 to C-Mad2 conversion in these experiments.

**Analysis of checkpoint complexes associated with the APC/C**

To analyze the composition of checkpoint complexes associated with the APC/C, we used a sequential purification approach (Fig. 4A). From a nocodazole extract, we first immunopurified the APC/C using a monoclonal APC4 antibody and eluted the complexes by antigen peptide competition. The purified APC/C was then split in 3 and one-third was then incubated with a BubR1 mAb recognizing residues 116–127 in the TPR domain (Fig. S1B), another one-third was incubated with the pan-Mad2 antibody and the last one-third was incubated with a mouse IgG. The complexes were then analyzed by quantitative protein gel blot and signals normalized to the BubR1 level (Fig. 4). As the BubR1 and Mad2 antibodies precipitated equal levels of APC7, it clearly indicates that very little Mad2-Cdc20 is present on the APC/C in agreement with Mad2 inhibiting and BubR1 stimulating the interaction with the APC/C.46–48 It was clear that the ratio of BubR1 to Mad2 was lower in the BubR1 purification, suggesting that indeed BBC complexes are present on the APC/C in addition to the MCC. In both the BubR1 and Mad2 purifications, Cdc20 was clearly present during an active checkpoint. We cannot, however, exclude antibody induction of O-Mad2 to C-Mad2 conversion in these experiments.
ubiquitinated, as revealed by the ubiquitination pattern recognized by the Cdc20 antibody. The level of Cdc20 ubiquitination appeared similar in the 2 purifications, and thus ubiquitination of Cdc20 does not correlate with Mad2 dissociation in endogenous complexes.

Discussion

The ability of Mad2 to convert between O-Mad2 and C-Mad2 is critical for the checkpoint and understanding how this is controlled is fundamental for understanding checkpoint signaling. Beautiful biochemical work have provided important models for how this is regulated but our ability to study Mad2 biology in vivo is still lagging behind. Here, we provide important tools in the form of defined Mad2-specific mAbs that are useful for dissecting endogenous Mad2 complexes and the conformation of Mad2 in these.

Our C-Mad2 specific antibody data is in agreement with the data from the Nigg lab showing that C-Mad2 mainly exists in complex with its known ligands Mad1 and Cdc20.40 However, in our size-exclusion chromatography experiments, we do detect a small amount of unliganded C-Mad2 and p31 comet Mad2 complexes, which could be C-Mad2 that has been removed from Cdc20. Recent work from the Cleveland lab proposes that C-Mad2 removed from the MCC could facilitate cytoplasmic amplification of the checkpoint signal by binding a new Cdc20 molecule, and potentially this is the C-Mad2 pool that we observe here.15

A puzzling observation is that while the Mad2-Cdc20 interaction is very dynamic and responds to SAC status, 2 other interactors that use a similar motif for binding Mad2, Mad1 and Kif20A, are stable (Fig. 2). Currently, we do not know the details of how C-Mad2 is removed from Cdc20 but it does not appear to depend on BubR1 because we recently showed that Mad2 is more efficiently removed from a Cdc20 mutant that does not bind BubR1.48 p31 comet binding is also not specific for the Mad2-Cdc20 complex because it also associates with Mad1-Mad2.40,49 Mad2 removal might require that it is bound to an unstructured region as in the case for Cdc20, and could explain why C-Mad2 is not removed from Mad1. However, the Mad2 binding site of Kif20A is at the extreme C terminus of Kif20A and one would assume that it should be very easy to remove C-Mad2.45 Understanding the details of specific Mad2 removal from Cdc20 is an important goal.

We find that Cdc20 is ubiquitinated in both BBC and MCC complexes and our data do not support that Cdc20 ubiquitination in itself dissociates the Mad2-Cdc20 complex,50 which is in agreement with observations from the Morgan lab.46 This does not rule out a subsequent role of ubiquitination on Cdc20 in SAC silencing. Another possibility is that Cdc20
ubiquitination is a bystander effect from the stable association of the MCC with the APC/C, and that p31comet dissociation of the MCC is required for subsequent Cdc20 degradation. The pan-Mad2 antibody described here will be useful in further investigations of Mad2 interactions since it efficiently purifies all Mad2 interactors and ubiquitinated Cdc20.

Materials and methods

Cell culture

HeLa cells were maintained in DMEM supplemented with 10% fetal bovine serum (FBS, HyClone) and 1% penicillin-streptomycin (Life Technologies). Protein expression was induced in stable cell lines by treatment with doxycycline (5 ng/ml).

Cloning and stable cell lines

Mad2 was cloned into pcDNA5/FRT/TO Venus (C-terminal tag). Stable cell lines were generated using the Flp-In system (Invitrogen) and clones were kept under selection by supplementing the growth media with 200 mg/ml hygromycin B and 5 mg/ml blasticidin S.

Antibodies

The following antibodies were used for Western blotting as indicated: APC7 1:1000 WB (rabbit, Bethyl A302-551A-1), Bub3 1:1000 WB (mouse, BD Biosciences 611731), BubR1 1:1000 WB (rabbit, raised against the TPR domain), Cdc20 1:1000 WB (mouse, Santa Cruz sc-13162), GFP 1:500 WB (mouse, Roche 11814460001), Mad1 1:1000 WB (mouse, Sigma M8069), Mad2 1:1000 WB (rabbit, Bethyl A300-301A), and p31 1:1000 WB (rabbit, raised against full length protein).

Size-exclusion chromatography

Cells were treated with nocodazole (200 ng/ml) for 18 hrs and mitotic cells were collected by shake-off. Cells were lysed for 30 mins in 20 mM Tris-HCl pH 7.8, 100 mM NaCl, 0.1% IGE-PAL, 1 mM dithiothreitol (DTT), 0.5 mM EDTA, 1x protease inhibitor cocktail (Roche) and 1x phosphatase inhibitor cocktail (Roche). Lysates were clarified at 13000 rpm for 15 mins and then for 20 mins at 55000rpm. 800 µg of protein was used for each IP in Fig 1C and 2 mgs in Fig 2A. Immunoprecipitations were performed with 10 µl of Protein-G sepharose 4B beads crosslinked to antibodies (4 µg antibody/10 µl beads) for 2 hours and then washed in lysis buffer by inversion 3 times.

For the reciprocal IPs in Figs 2C and 4B, the cell lysates were obtained as described in the above paragraph. In Fig. 2C, the APC/C was then immunoprecipitated from 6 mgs of protein, whereas in Fig. 4B the IP was preformed from 30 mg. IPs were performed for 2 hours using an APC4 Ab. IPs were then washed 3 times by inversion and the APC/C was eluted form the beads for 2 hours using 1 ml of APC4 peptide dissolved in lysis buffer (1mg/ml). After pelleting of the beads, the supernatant was divided into 300 µl aliquots and used to perform IPs with GFP-Trap beads (Chromotek) (Fig. 2C), or panMad2 or BubR1 antibodies (Fig. 4B). IPs were washed in lysis buffer by inversion 3 times.

Isothermal titration calorimetry measurements

All proteins and antibodies were dialyzed extensively in PBS prior to the ITC experiments. Protein concentrations were determined by UV spectroscopy and a molar extinction coefficient at 280 nm of 28545 M⁻¹ cm⁻¹ for both MD2L1-L13A-R133A and MD2L1-V193N-R133A. ITC experiments were performed at 25°C using an ITC200 instrument (Microcal). Three µL volumes of either MD2L1-L13A-R133A or MD2L1-V193N-R133A, at approximately 150 µM, were titrated into the ITC sample cell containing approximately 15 µM of antibody, until saturation was achieved. The heat of the reaction was obtained by integrating each peak after the injection of protein and fit to a model describing a single binding site using software provided by the ITC200 manufacturer.

Silac labeling

For quantitative proteomic experiments, cells were grown in SILAC DMEM (PAA) supplemented with 10% dialyzed FBS (PAA), penicillin-streptomycin and glutamax (Life Technologies) and isotope labeled L-arginine and L-lysine. HeLa cells were grown in light “L” SILAC medium supplemented with saline (PBS) + 0.1% Tween for 30 minutes. The cells were incubated with primary antibodies for 2 hours at room temperature (primary antibodies were used at the following concentrations: pan-Mad2 (16 µg/ml), C-Mad2 (4 µg/ml), O-Mad2 (4 µg/ml) and CREST (Antibodies Inc., 1:400), followed by 45 minutes of incubation with appropriate secondary antibodies and DAPI (1:1000). Images were acquired taking z stacks of 200 nm using a 100X/1.4NA objective on a DeltaVision Elite Microscope (GE Healthcare). Figures were generated using ImageJ.
normal arginine (R0) and lysine (K0) (Sigma). For the medium "M" condition cells were grown in SILAC DMEM supplemented with U-13C6 arginine (K6) and 4, 5, 5-D4 lysine (K4) and for the heavy "H" labeling cells were grown with U-13C6, U-15N4 arginine (R10) and U-13C6, U-15N2 lysine (K8) (Cambridge Isotope Laboratories). Cells were grown in SILAC medium for at least 5 divisions before harvest.

Mass spectrometry analysis

The eluate from the beads was reduced, alkylated and loaded onto an SDS–PAGE gel. The lane was cut into 7 pieces and subjected to in-gel tryptic digestion and subsequent sample desalting and concentration. The resulting peptide mixture was analyzed by nano-HPLC–MS/MS using an easy-nLC nanoflow system (Thermo Scientific, Odense, Denmark) connected to a Q Exactive mass spectrometer (Thermo Scientific, Bremen, Germany) through a nano-electrospray ion source. The column length was 150 mm with an inner diameter of 75 μm, and packed in-house with 1.9 μm C18 beads (Reposil-AQ Pur, Dr. Maisch). Each sample was analyzed using a 140 min gradient from 4 to 36% acetonitrile in 0.5% acetic acid. The mass spectrometer was run as described for sensitive acquisition. Data analysis was performed in the MaxQuant environment configured for triple SILAC and the Andromeda search engine. The database searched was the UniProt complete human environment concatenated with a list of commonly observed background contaminants.

Protein production and purification

Mad2 DNA and mutants thereof were cloned into pET30 and expressed in BL21(DE3) cells at 37 degrees. Cells were harvested and Mad2 affinity purified using Ni-NTA affinity resin (Qiagen) and subsequently purified on a Superdex 75 column.

Disclosure of potential conflicts of interest

No potential conflicts of interest were disclosed.

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