RESEARCH ARTICLE



SIRT2 deacetylates GRASP55 to facilitate post-mitotic Golgi assembly

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ABSTRACT

Sirtuin 2 (SIRT2) is an NAD-dependent sirtuin deacetylase that regulates microtubule and chromatin dynamics, gene expression and cell cycle progression, as well as nuclear envelope reassembly. Recent proteomic analyses have identified Golgi proteins as SIRT2 interactors, indicating that SIRT2 may also play a role in Golgi structure formation. Here, we show that SIRT2 depletion causes Golgi fragmentation and impairs Golgi reassembly at the end of mitosis. SIRT2 interacts with the Golgi reassembly stacking protein GRASP55 (also known as GORASP2) in mitosis when GRASP55 is highly acetylated on K50. Expression of wild-type and the K50R acetylation-deficient mutant of GRASP55, but not the K50Q acetylation-mimetic mutant, in GRASP55 and GRASP65 (also known as GORASP1) double-knockout cells, rescued the Golgi structure and post-mitotic Golgi reassembly. Acetylation-deficient GRASP55 exhibited a higher self-interaction efficiency, a property required for Golgi structure formation. These results demonstrate that SIRT2 regulates Golgi structure by modulating GRASP55 acetylation levels.

KEY WORDS: SIRT2, Golgi, Acetylation, GRASP55, Golgi assembly, Cell cycle

INTRODUCTION

Sirtuin 2 (SIRT2) is a canonical NAD-dependent sirtuin deacetylase located in the endoplasmic reticulum (ER)–Golgi intermediate compartment (ERGIC) (Budayeva and Cristea, 2016). SIRT2 traffics between the ER and Golgi, shuttles between the cytoplasm and the nucleus, and associates with the centrosomes and the mitotic spindle during metaphase (Budayeva and Cristea, 2016; Inoue et al., 2007; North and Verdin, 2007). Accordingly, known SIRT2 substrates range from microtubule, membrane trafficking and metabolic proteins to chromatin-bound proteins and mitotic regulators.

SIRT2 deacetylates α -tubulin and impairs cell motility and neurite outgrowth (Janke, 2014; North et al., 2003; Pandithage et al., 2008). SIRT2 has thus been implicated in neurodegenerative diseases and its inhibition has neuroprotective effects in Parkinson's and Alzheimer's disease models (Chen et al., 2015; Godena et al., 2014; Harting and Knöll, 2010; Outeiro et al., 2007; Silva et al., 2016). Furthermore, SIRT2 regulates cell survival via deacetylation of transcription factors FOXO1, nuclear factor κB

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(NF- κ B) and p53 (Jin et al., 2008, 2007; Rothgiesser et al., 2010), which may contribute to its tumor suppressive effect, as found in mammary tumors, hepatocellular carcinoma and skin squamous cell carcinoma (Bosch-Presegue and Vaquero, 2011; Kim et al., 2011; Serrano et al., 2013). SIRT2 may also exert its tumor-suppressive function by ensuring normal mitotic progression via activation of the anaphase-promoting complex (Kim et al., 2011) and by destabilizing ATP-citrate lyase (Lin et al., 2013) and phosphoglycerate mutase (Tsusaka et al., 2014). Finally, SIRT2 counteracts replication stress and preserves genomic integrity by deacetylating histone H4K16 at the G2/M transition (Vaquero et al., 2006), stimulating H4K20 monomethylation (Serrano et al., 2013) as well as CDK9 (Zhang et al., 2013) and ATRIP activity (Zhang et al., 2016).

SIRT2 interactomes from MRC5 and HEK293 cells pointed to a new connection between SIRT2 and the ER-Golgi trafficking pathway (Budayeva and Cristea, 2016). Consistent with this, our previous proteomic study revealed ER and Golgi proteins as SIRT2interacting partners (Kaufmann et al., 2016), indicating a role for SIRT2 in ER and Golgi structure formation. To date, the regulation of Golgi structural proteins by acetylation has not been reported. The Golgi is a membranous organelle that mediates protein and lipid trafficking. It has been estimated that proteins encoded by over onethird of the genes in the human genome travel through the Golgi to their final destinations. Most of these proteins are also modified by Golgi-localized enzymes. In animals, the Golgi has a unique structure, comprising a stack of flattened cisternal membranes. Formation of the Golgi stacked structure requires two Golgi reassembly stacking proteins, GRASP55 and GRASP65 (also known as GORASP2 and GORASP1, respectively), which form trans-oligomers to hold Golgi membranes into stacks (Shorter et al., 1999; Zhang and Wang, 2015, 2016). Both GRASP55 and GRASP65 are peripheral membrane proteins located in the medial/ trans- and cis-Golgi, respectively. GRASP55 and GRASP65 selfinteract to form trans-oligomers (Wang et al., 2003; Xiang and Wang, 2010); this property is believed to be essential for their roles in Golgi cisternal stacking (Shorter et al., 1999; Tang et al., 2010b; Xiang and Wang, 2010; Zhang and Wang, 2015) and ribbon linking (Feinstein and Linstedt, 2008). GRASP55 and GRASP65 play complementary roles in Golgi structure formation, as single knockdown of GRASP55 or GRASP65, by means of siRNA, or knockout by CRISPR/Cas9, cause a mild effect on the Golgi structure, while double knockdown or knockout of both GRASP proteins causes the disassembly of both Golgi stack and ribbon structures (Bekier et al., 2017; Xiang and Wang, 2010; Xiang et al., 2013).

The Golgi structure is fragmented at the onset of mitosis and reassembled in telophase (Tang and Wang, 2013; Zhang and Wang, 2015). The process of Golgi cisternal unstacking at the onset of mitosis and restacking after its completion is known to be regulated by GRASP55 and GRASP65 phosphorylation (Duran et al., 2008; Feinstein and Linstedt, 2007; Xiang and Wang, 2010). In this study,

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we made an unexpected finding that SIRT2 interacts with and deacetylates GRASP55. Our results revealed a novel role for SIRT2 in Golgi structure formation through the modulation of GRASP55 acetylation levels, which affects GRASP55 self-interaction efficiency and post-mitotic Golgi reassembly.

RESULTS

SIRT2 interacts with and deacetylates GRASP55

Our published SIRT2 interactome revealed that SIRT2 interacts with various Golgi proteins such as GRASP55, Golgin-160 (also known as GOLGA3) and GCP60 (also known as ACBD3) (Kaufmann et al., 2016). GRASP55 ranked second in our previously published interactome analysis and was identified in a proximity biotinylation (BioID) experiment in which SIRT2 was fused to a promiscuous biotin ligase from E. coli (BirA) at both the N- and the C-terminus (Roux et al., 2012). As BioID reveals mitotic interactors due to prolonged biotin labeling (Lambert et al., 2015), we examined the interaction between SIRT2 and GRASP55 in asynchronous and nocodazole-arrested mitotic cells by performing co-immunoprecipitation experiments. To preserve the acetylation state, we included deacetylase inhibitors, trichostatin A and nicotinamide, in the lysis buffer. Our results showed that SIRT2 interacts with GRASP55 predominantly in mitosis (Fig. 1A). Another Golgi protein, Golgin-84 (GOLGA5), did not show interaction with SIRT2 (Fig. 1A), consistent with our mass

spectrometry data showing that Golgin 84 ranked relatively low on the SIRT2 interactome list (Kaufmann et al., 2016). Given that the endogenous SIRT2 protein level is very low (Fig. S1), we failed to detect an interaction between endogenous SIRT2 and GRASP55. Therefore, we performed reciprocal co-immunoprecipitation with HA–SIRT2, which exhibited strong interaction with GRASP55 during mitosis (Fig. 1B). These results confirm our mass spectrometry data and emphasize the value of BioID in revealing interactions that occur within specific stages of the cell cycle in an asynchronous cell population.

Since SIRT2 interacts with its substrate only when the substrate is acetylated, our results suggest that GRASP55 may be a SIRT2 target on the Golgi membranes. To determine whether SIRT2 directly deacetylates GRASP55, we performed *in vitro* acetylation and deacetylation assays using purified proteins. The acetylation level of GRASP55 was detected by an anti-acetylated lysine (AcK) antibody. GRASP55 was acetylated by p300 acetyltransferase (EP300) and was deacetylated by wild-type (WT) SIRT2 (Fig. 1C, lanes 2 and 3) but not its catalytically inactive H150Y mutant (Fig. 1C, lanes 4 and 5) (Pandithage et al., 2008).

To determine the acetylation state of GRASP55 *in vivo*, we immunoprecipitated GRASP55–GFP from asynchronous or mitotic HEK293T cells and analyzed its acetylation by mass spectrometry. GRASP55 was found to be highly acetylated on K50 in mitosis (Table S1). We then used parallel reaction monitoring (PRM) to

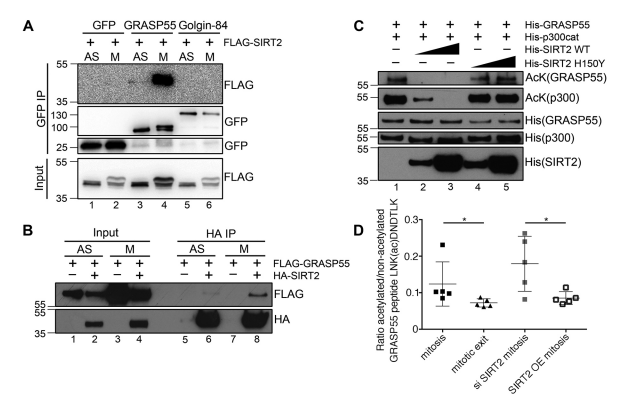


Fig. 1. SIRT2 interacts with GRASP55 in mitosis and deacetylates GRASP55 at mitotic exit. (A) SIRT2 co-immunoprecipitates with GRASP55 but not Golgin-84 during mitosis. Asynchronous (AS) and nocodazole-arrested mitotic HeLa cells co-overexpressing GFP-tagged GRASP55 or Golgin-84 and FLAG–SIRT2 were lysed, immunoprecipitated with an anti-GFP antibody, and blotted for GFP and FLAG. The upper bands of FLAG–SIRT2 and GRASP55–GFP in mitotic samples correspond to their phosphorylated forms. (B) Western blots of the HA immunoprecipitation from HEK293T cells expressing HA–SIRT2 and FLAG–GRASP55. (C) *In vitro* acetylation of GRASP55 with p300 acetyltransferase and deacetylation of GRASP55 by SIRT2. 1.5 µg of GRASP55, 0.15 µg of p300 acetyltransferase and 0.15 or 1.5 µg (as indicated by triangle) of SIRT2 or SIRT2 H150Y were present in the loaded samples. The acetylation level of GRASP55 and p300 was detected by an anti-acetylated lysine (AcK) antibody. (D) Targeted mass spectrometry analysis of GRASP55 K50 acetylation in HEK293T cells transfected with GRASP55–GFP. Mitotic samples were collected after 16 h synchronization with nocodazole. Mitotic exit samples were collected 2 h after release from nocodazole. SIRT2 siRNA was transfected for 72 h. HA–SIRT2 was overexpressed for 24 h. K50 acetylation level was determined as a ratio between modified and unmodified LNK(ac)DNDTLK peptide (*n*=5). Quantification results are presented as mean±s.e.m. **P*<0.05 (two-tailed Student's *t*-test).

analyze changes in GRASP55 acetylation at mitotic exit, as well as after SIRT2 depletion or overexpression in mitotic cells. GRASP55 K50 acetylation was significantly reduced at mitotic exit and after SIRT2 overexpression, whereas SIRT2 depletion further increased the K50 acetylation level in mitosis (Fig. 1D; Table S2). These results demonstrate that GRASP55 acetylation is regulated by SIRT2.

SIRT2 depletion causes Golgi fragmentation

The identification of GRASP55 as a novel SIRT2 substrate raised a possibility that SIRT2 may regulate Golgi morphology through GRASP55. To test the role of SIRT2 in Golgi structure formation, we depleted SIRT2 in HeLa and U2OS cells through siRNA and determined the effect on the Golgi morphology by immunostaining for GRASP55 and GM130 (also known as GOLGA2). SIRT2 silencing caused Golgi fragmentation in both cell lines (Fig. 2A–D), whereas overexpression of SIRT2 in WT cells had no significant effects on the Golgi morphology, although the Golgi appeared slightly more compact than that in control cells (Fig. 2, compare E with B). In addition, the majority of cellular SIRT2 localized in the cytosol including the Golgi region (Fig. 2E), consistent with its role in Golgi structure formation. To further understand the effect of SIRT2 depletion on the Golgi morphology, we analyzed the Golgi ultrastructure by electron microscopy (EM). The number of cisternae per Golgi stack and the length of cisternae were both significantly reduced in SIRT2-depleted cells (Fig. 2F-H; Fig. S2), indicating that SIRT2 is required for proper maintenance of the Golgi structure.

Based on the fixed cell samples, we were unable to discern the cell cycle stage at which Golgi fragmentation occurred in SIRT2-depleted cells. Therefore, we generated a U2OS cell line stably expressing GRASP55–GFP and performed live-cell imaging to monitor the Golgi morphology in control and SIRT2-depleted cells during the cell cycle. Remarkably, SIRT2 silencing impaired post-mitotic Golgi reassembly (Fig. 2I,J). While the Golgi was completely assembled 60 min after metaphase in WT cells, it remained as scattered fragments at 120 min in SIRT2-depleted cells (Fig. 2I; Movies 1 and 2). Golgi fragmentation was 2-fold more pronounced in SIRT2-silenced cells than in control cells (Fig. 2J), which correlates well with the Golgi fragmentation observed upon immunostaining of endogenous GRASP55 in SIRT2-depleted cells (Fig. 2A–D).

The deacetylase activity of SIRT2 is required for maintaining the Golgi structure

To validate the specific effect of SIRT2 silencing on the Golgi morphology, we transfected SIRT2-depleted cells with siRNA-resistant SIRT2 constructs. As shown in Fig. 3A–C, expression of WT SIRT2, but not its inactive H150Y mutant, largely rescued Golgi fragmentation in SIRT2-depleted cells. These results indicate that the deacetylase activity of SIRT2 is required for maintaining the Golgi structure.

It has been previously shown that SIRT2 depletion may increase or decrease the protein levels of its substrates (Budayeva and Cristea, 2016). We therefore tested the effect of SIRT2 depletion or overexpression on the level of Golgi structural proteins by western blotting. Knockdown or overexpression of SIRT2 did not affect the level of various Golgi proteins tested in this experiment, including GRASP55, GRASP65, GM130, Golgin-160 and Golgin-84 (Fig. 3D). This suggests that aberrant acetylation of Golgi proteins upon SIRT2 depletion causes Golgi fragmentation by perturbing protein–protein interactions rather than protein stability.

GRASP55 deacetylation is required for post-mitotic Golgi structure formation

Previous work has shown that double knockdown or knockout of GRASP55 and GRASP65 causes the disassembly of both Golgi stack and ribbon structures (Bekier et al., 2017; Xiang and Wang, 2010; Xiang et al., 2013), whereas reintroduction of a single GRASP protein in GRASP55 and GRASP65 (hereafter GRASP55/65) double-knockout cells can largely restore the Golgi structure (Bekier et al., 2017). To test whether aberrant acetylation of GRASP55 could be responsible for Golgi fragmentation in SIRT2-depleted cells, we generated K50R and K50Q mutants of GRASP55 to mimic hypoand hyper-acetylation, respectively. Expression of these constructs did not affect GRASP55 phosphorylation in mitotic cells (Fig. S3). Consistent with previous reports, WT GRASP55-GFP, but not GFP alone, effectively rescued the Golgi structure in GRASP55/65 double-knockout HeLa cells (Fig. 4). The acetylation-deficient K50R mutant restored the Golgi structure with similar efficiency to WT GRASP55, whereas the acetylation-mimetic K50Q mutant had a much weaker effect (Fig. 4A-C).

Furthermore, we specifically examined the Golgi morphology in the later stages of mitosis in GRASP55/65 double-knockout cells expressing GRASP55 WT, K50R and K50Q (Fig. 5). During mitosis, the Golgi is disassembled into mitotic clusters (observed as dots under the microscope) and vesicles (observed as a haze because the size of the vesicles is below the resolution of conventional microscopy). As cell cycle progresses, the Golgi structure reassembles in telophase and cytokinesis; the Golgi 'haze' gradually disappears and the Golgi dots accumulate and merge into a single membrane complex. Expression of WT GRASP55, as well as the acetylation-deficient K50R mutant, accelerated Golgi reassembly in telophase and cytokinesis (Fig. 5). Conversely, the acetylation-mimetic K50Q mutant was less effective in facilitating post-mitotic Golgi reassembly (Fig. 5). This result is consistent with impaired post-mitotic Golgi reassembly caused by SIRT2 depletion, as shown in Fig. 2I. Taken together, GRASP55 deacetylation is required for optimal Golgi structure formation, in particular at mitotic exit.

The K50R acetylation-deficient mutant of GRASP55 partially rescues Golgi fragmentation caused by SIRT2 depletion

Given that (1) GRASP55 is a substrate of the SIRT2 deacetylase (Fig. 1), (2) SIRT2 depletion induces Golgi fragmentation (Figs 2 and 3), and (3) expression of the K50R acetylation-deficient GRASP55 mutant rescues Golgi fragmentation in GRASP55/65 double-knockout cells (Figs 4 and 5), we further asked whether GRASP55 acetylation mutants could rescue the SIRT2-depletion phenotype. Therefore, we expressed GFP-tagged GRASP55 WT, K50R and K50Q in SIRT2-depleted cells. As shown in Fig. 6, the acetylation-deficient K50R mutant exhibited a stronger effect than WT GRASP55 in rescuing the Golgi structure in SIRT2-depleted cells, whereas the acetylation-mimetic K50O mutant had no effect. These results suggest that GRASP55 deacetylation by SIRT2 contributes to Golgi structure maintenance. The incomplete rescue of the Golgi structure by the acetylation-deficient K50R mutant indicates that SIRT2 may have additional substrates on the Golgi, or a cycle of acetylation and deacetylation may be required for Golgi structure formation.

GRASP55 deacetylation facilitates self-interaction

To address how GRASP55 acetylation may affect its function in Golgi structure formation, we probed the self-interaction efficiency of WT GRASP55 and its acetylation mutants (Fig. 7). GFP-tagged WT, K50R and K50Q GRASP55 and GRASP55–FLAG were

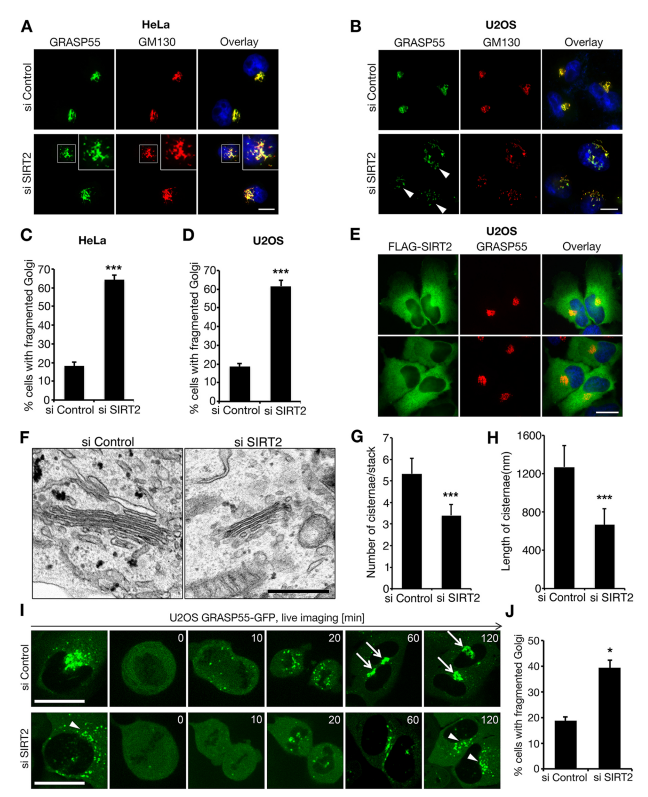


Fig. 2. SIRT2 depletion causes Golgi fragmentation and impairs Golgi structure formation. (A–D) Representative immunofluorescence images of HeLa (A) and U2OS (B) cells transfected with control or SIRT2 siRNA and co-stained for GRASP55 (green) and GM130 (red). Representative fragmented Golgi are enlarged in the insets in A. Arrowheads in B indicate cells exhibiting fragmented Golgi. Scale bars: 20 µm. (C,D) Quantification of Golgi fragmentation in A and B from three independent experiments (*n*=210). (E) SIRT2 overexpression does not significantly perturb the Golgi morphology in U2OS cells. Cells were transfected with FLAG–SIRT2 for 24 h and co-stained for FLAG (green) and GRASP55 (red). Two sets of cells are shown. Scale bar: 20 µm. (F) HeLa cells were transfected with control or SIRT2 siRNA and analyzed by EM. Representative EM images are shown. Scale bar: 500 nm. (G) Quantification of the number of cisternae per stack (*n*=20). (H) Quantification of the length of Golgi cisternae (*n*=20). (I) Live-cell images of U2OS cells stably expressing GRASP55–GFP 72 h after SIRT2 siRNA transfection. Note that the Golgi is assembled in control cells (arrows) at 60 min but remains fragmented in SIRT2-depleted cells after 120 min (arrowheads). Scale bar: 20 µm. (J) Quantification of Golgi fragmentation from I in control and SIRT2-knockdown cells using GRASP55–GFP as a marker (*n*=100 for siControl; *n*=95 for siSIRT2). All quantification results in this figure are presented as mean±s.e.m. **P*<0.05; ****P*<0.001 (two-tailed Student's *t*-test).

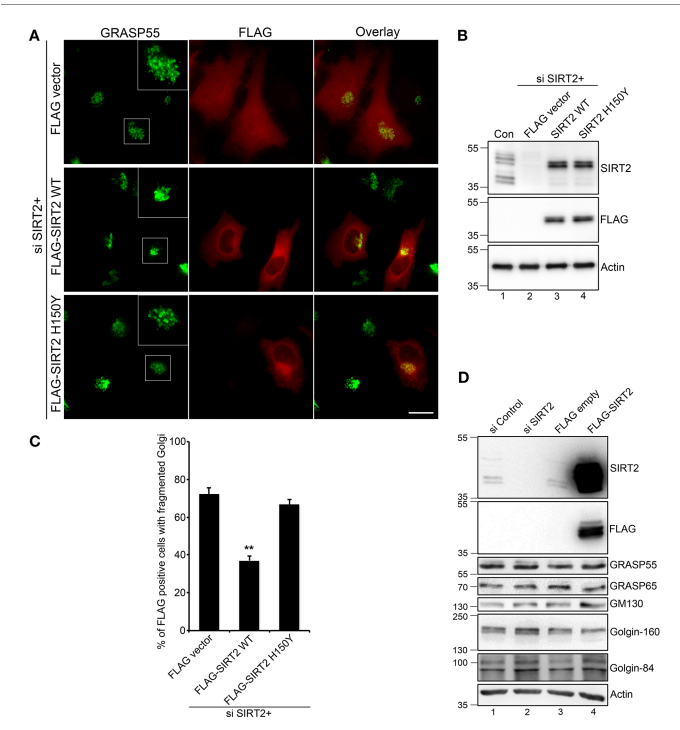


Fig. 3. The deacetylase activity of SIRT2 is required for maintaining an intact Golgi structure. (A) HeLa cells were transfected first with SIRT2 siRNA for 48 h, then with a FLAG vector (as control) or siRNA-resistant FLAG–SIRT2 WT or H150Y mutant for another 24 h, and co-stained for GRASP55 (green) and FLAG (red). Scale bar: 20 μm. (B) Cells in A were analyzed by western blotting to show the knockdown efficiency of endogenous SIRT2 and the expression level of FLAG-tagged SIRT2. (C) Quantification of Golgi fragmentation in A from three independent experiments (*n*=200). Results are presented as mean±s.e.m. ***P*<0.01 (two-tailed Student's *t*-test). (D) HeLa cells were transfected with SIRT2 siRNA for 72 h or with FLAG–SIRT2 for 24 h, lysed, and blotted for the indicated Golgi proteins. Control siRNA or an empty FLAG vector were used as controls.

co-expressed, and synchronized mitotic cells were analyzed by GFP immunoprecipitation and anti-FLAG western blotting. GRASP55 K50R showed higher self-interaction efficiency compared to WT or K50Q (Fig. 7A,B, lane 7 versus 6 and 8). Co-expression of GRASP55 K50R and K50Q exhibited an intermediate self-interaction efficiency, being lower compared to K50R–K50R but higher than K50Q–K50Q (Fig. 7C,D, lanes 9 and 11 versus 8 or 12).

Reduced acetylation of GRASP55 therefore facilitates its selfinteraction and contributes to post-mitotic Golgi structure formation.

DISCUSSION

Recent proteomic analyses have revealed proteins in the ER and Golgi membrane networks as novel SIRT2-interacting partners

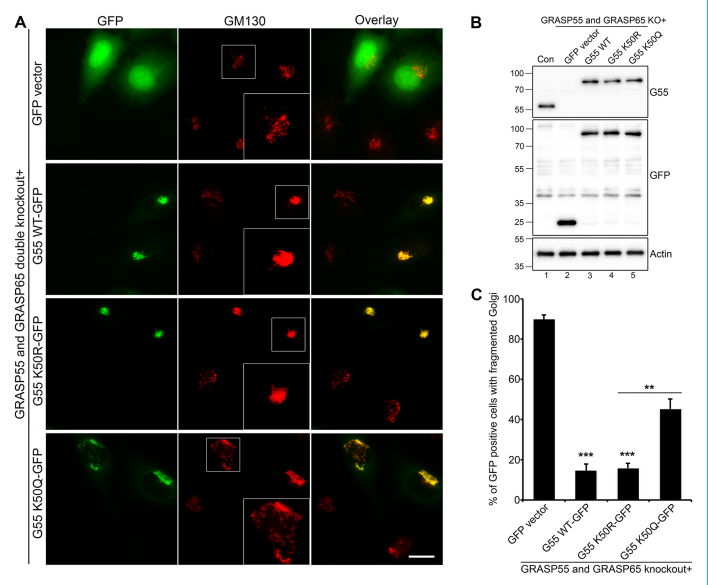


Fig. 4. GRASP55 deacetylation is required for Golgi structure formation. (A) GRASP55/65 double-knockout cells were transfected with a GFP vector (as control), or GFP-tagged WT, acetylation-deficient mutant K50R or acetylation-mimetic mutant K50Q GRASP55 (labeled G55) for 24 h, and stained for GM130. Note that expression of WT and K50R efficiently rescued the Golgi structure. Scale bar: $20 \,\mu$ m. (B) Cells in A were analyzed by western blotting to show the knockout efficiency of endogenous GRASP55 and the expression level of GFP-tagged GRASP55 WT and mutants. (C) Quantification of Golgi fragmentation in A from three independent experiments (*n*=210). Quantification results are presented as mean±s.e.m. ***P*<0.01; ****P*<0.001 (two-tailed Student's *t*-test).

(Budaveva and Cristea, 2016). In this study, we confirmed the interaction between SIRT2 and the Golgi reassembly stacking protein GRASP55, which occurs specifically in mitosis. We found that SIRT2 depletion not only increases GRASP55 acetylation level but also induces Golgi fragmentation and impairs post-mitotic Golgi reassembly. Furthermore, we discovered that GRASP55 is highly acetylated in mitosis on K50 and deacetylated at mitotic exit, indicating that SIRT2 regulates Golgi structure formation by deacetylating GRASP55 at the end of mitosis. Importantly, expression of WT or the acetylation-deficient K50R mutant of GRASP55, but not the acetylation-mimetic K50Q mutant, rescued Golgi fragmentation and restored post-mitotic Golgi reassembly in GRASP55/65 double-knockout cells, indicating that GRASP55 deacetylation is required for post-mitotic Golgi structure formation. Acetylation of GRASP55 in mitosis presumably facilitates Golgi disassembly, whereas its deacetylation is clearly required for proper Golgi reassembly at the mitotic exit. Indeed, acetylation-deficient

GRASP55 has a higher self-interaction efficiency, which facilitates Golgi reassembly. Thus, our study revealed acetylation as a novel mechanism that regulates Golgi structure formation.

There are a growing number of proteins similar to GRASP55 that undergo cell cycle-dependent changes in acetylation. For example, histone H4 is highly acetylated on K16 during S phase and deacetylated by SIRT2 at the G2/M transition (Vaquero et al., 2006); cyclin A acetylation in mitosis targets it for ubiquitylation and subsequent degradation (Mateo et al., 2009); ANKLE2 is highly acetylated in mitosis and the regulation of its acetylation state by SIRT2 is necessary for proper nuclear envelope reassembly at the end of mitosis (Kaufmann et al., 2016). Acetylation is highly abundant during mitosis (Chuang et al., 2010) and will doubtlessly continue to emerge as a post-translational protein modification that regulates cell cycle progression. Cell cycle regulation of protein acetylation necessitates tight regulation of deacetylases; phosphorylation during mitosis stabilizes SIRT2, whereas dephosphorylation at the

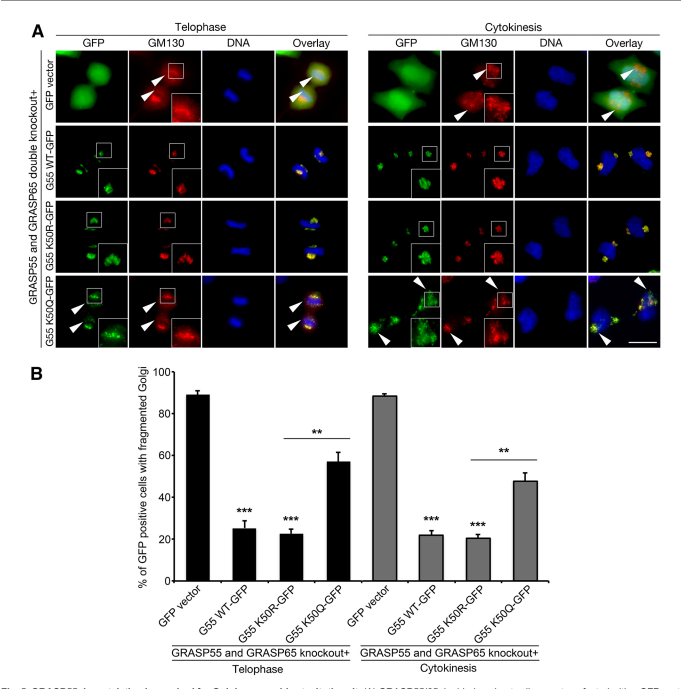


Fig. 5. GRASP55 deacetylation is required for Golgi reassembly at mitotic exit. (A) GRASP55/65 double-knockout cells were transfected with a GFP vector (as control), or GFP-tagged WT, K50R or K50Q GRASP55 (labeled G55), and stained for GM130. Representative fluorescence images of telophase and cytokinesis cells are shown. Arrowheads indicate cells with fragmented Golgi. Indicated areas, showing the representative Golgi morphology, are enlarged in the insets. Scale bar: 20 µm. (B) Quantification of Golgi fragmentation in A from three independent experiments (*n*=200). Quantification results are presented as mean ±s.e.m. ***P*<0.01; ****P*<0.001 (two-tailed Student's *t*-test).

mitotic exit leads to proteasome-mediated degradation (Dryden et al., 2003; North and Verdin, 2007).

It has been reported that tubulin acetylation on kinetochore microtubules in the mitotic spindle is regulated by SIRT2 (Nagai et al., 2013). Given that the microtubule cytoskeleton is required for an intact Golgi, this raises a question as to whether SIRT2 regulates the Golgi structure through microtubules. However, expression of the GRASP55 K50R mutant largely rescues the Golgi structure in SIRT2-depleted cells, suggesting that SIRT2 directly regulates Golgi structure through GRASP55.

Identifying SIRT2 deacetylase substrates within the ER-Golgi network raises the question of where and how these substrates are initially acetylated. Lysine acetylation can occur in the lumen of the ER (Pehar and Puglielli, 2013), while the Golgi lumen is known to possess deacetylase activity (Costantini et al., 2007). Transient acetylation of ER membrane proteins seems to facilitate protein folding by neutralizing the positive charge in kinetically unfavorable areas of the nascent protein and may protect correctly folded proteins from disposal (Pehar and Puglielli, 2013). For example, transient acetylation of the nascent ER form of the β -secretase BACE1 is

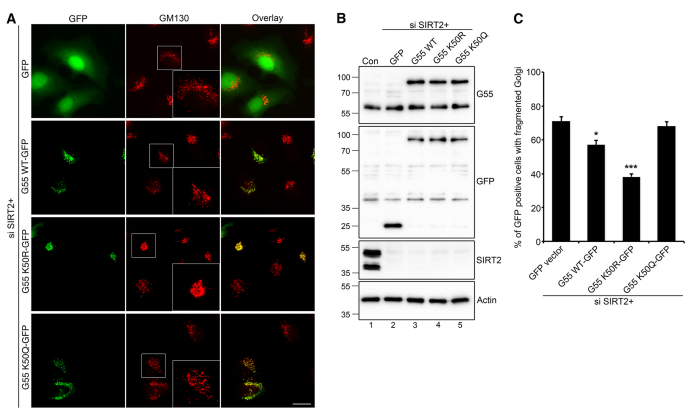


Fig. 6. Acetylation-deficient GRASP55 partially rescues Golgi fragmentation caused by SIRT2 depletion. (A) HeLa cells were transfected first with SIRT2 siRNA for 48 h, then with a GFP vector (as control) or GFP-tagged WT, K50R or K50Q GRASP55 (labeled G55) for another 24 h, and stained for GM130. Indicated areas, showing the representative Golgi morphology, are enlarged in the insets. Scale bar: 20 μ m. (B) Cells in A were analyzed by western blotting to show the knockdown efficiency of endogenous SIRT2 and the expression level of GFP-tagged GRASP55. (C) Quantification of Golgi fragmentation in A from three independent experiments (*n*=221). Quantification results are presented as mean±s.e.m. **P*<0.05; ****P*<0.001 (two-tailed Student's *t*-test).

required for its maturation within the Golgi where it is deacetylated (Costantini et al., 2007). However, unlike secretory and membrane proteins that are synthesized by the ER and transported to the Golgi, GRASP55 is a peripheral membrane protein on the cytoplasmic surface of the Golgi. While we showed that p300 acetyltransferase can acetylate GRASP55 *in vitro*, the acetylation mechanism *in vivo* will be a subject for future investigation.

Golgi fragmentation accelerates protein trafficking by enhancing vesicle budding from the Golgi membranes in interphase cells (Wang et al., 2008; Zhang and Wang, 2015). Fragmentation of the Golgi is associated with neurodegenerative diseases such as Alzheimer's and Parkinson's (Fujita et al., 2006; Gonatas et al., 2006; Joshi and Wang, 2015). In Alzheimer's disease, Golgi fragmentation was linked to GRASP65 phosphorylation by the Cdk5 kinase, which results in increased trafficking of the amyloid precursor protein (APP) and increased amyloid β (A β) production (Joshi et al., 2014; Joshi and Wang, 2015). In Parkinson's disease, Golgi fragmentation was linked with altered levels of Rab proteins and the SNARE protein syntaxin 5, and precedes α -synuclein aggregation (Rendon et al., 2013). Interestingly, SIRT2 is also implicated in neurodegenerative diseases and SIRT2 inhibition was shown to have a neuroprotective effect in various Parkinson's disease models (Chen et al., 2015; Godena et al., 2014; Outeiro et al., 2007) and in Alzheimer's disease (Silva et al., 2016). It would thus be interesting to examine Golgi morphology in neurodegenerative disease models following SIRT2 depletion or inhibition.

While our results indicate that GRASP55 is a specific target of SIRT2 through which SIRT2 regulates Golgi morphology, future

work may still be needed to characterize other potential SIRT2 targets within the Golgi network.

MATERIALS AND METHODS Cell lines

Tissue culture media, supplements and antibiotics were purchased from Sigma-Aldrich or Invitrogen. Bovine serum was obtained from Sigma-Aldrich or Gemini. All cell lines were maintained in Dulbecco's modified Eagle's medium (DMEM 4.5 g/l glucose) supplemented with 10% fetal bovine serum, 1% L-glutamine, 1% penicillin-streptomycin under 5% CO₂ at 37°C. HEK293T (ATCC, CRL-3216) and HeLa cells (ATCC, CCL-2) were used for transient transfections. To obtain mitotic cells, HEK293T and HeLa cells were treated with 330 nM nocodazole (Sigma) for 18 h. U2OS cells (gift from Dr Xiaochun Yu, City of Hope National Medical Center, Duarte, CA) stably expressing GRASP5–GFP were generated by transfection of the respective plasmid and selection of stable clones in the presence of 400 µg/ml G418 (Sigma-Aldrich). Transfection was performed with polyethylenimine (PEI; Polysciences).

Plasmids and proteins

The cloning and expression of SIRT2 and p300 acetyltransferase were described previously (Kaufmann et al., 2016). Human GRASP55 was cloned into pDEST N3xFLAG from pDONR23 (human ORFeome 5.1 collection; kindly provided by Ivana Grbesa, Laboratory for Advanced Genomics, Ruđer Bošković Institute, Zagreb, Croatia). Rat GRASP55 was cloned into pEGFP-N1 between BgIII and HindIII for mammalian expression and into pET23 (Novagen) for bacterial expression (Zhang et al., 2019, 2018). siRNA-resistant SIRT2 constructs were generated by site-directed mutagenesis according to the FastCloning protocol (Li et al., 2011) using the following primers: 5'-ACC-AAAATGCGAGGATTGTCAGAGCCTGGTGAAGCCTGATATCGTC-3'

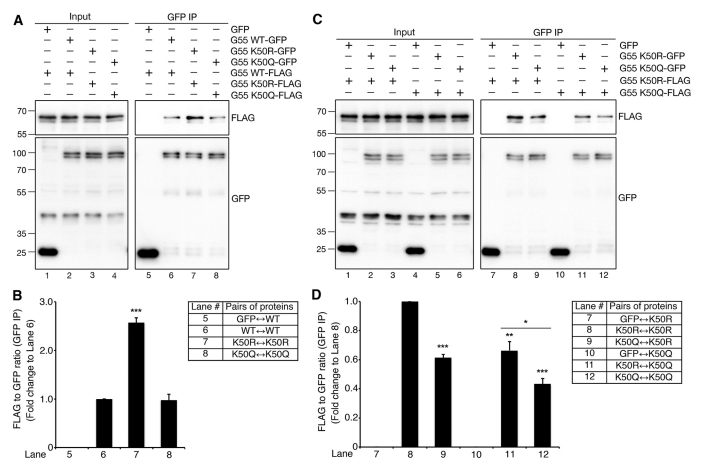


Fig. 7. GRASP55 deacetylation facilitates self-interaction. HeLa cells were co-transfected with GFP- and FLAG-tagged GRASP55 (G55) WT, K50R or K50Q constructs for 24 h and synchronized with nocodazole for another 18 h. Mitotic cells were collected, lysed, immunoprecipitated with a GFP antibody, and immunoblotted for GFP and FLAG. (A,B) Homologous GFP- and FLAG-tagged GRASP55 constructs were co-transfected, analyzed by co-immunoprecipitation, and their levels quantified. (C,D) Homologous or heterologous GFP- and FLAG-tagged GRASP55 constructs were co-transfected, analyzed, and their levels quantified. (A,C) Western blots of the co-immunoprecipitated proteins. (B,D) Quantification of the interaction efficiency found in experiments shown in A and C as determined by calculating the ratio of FLAG to GFP intensity in the GFP immunoprecipitates from three independent experiments. The homologous interaction of WT GRASP55 or the K50R mutant was normalized to 1 in B and D, respectively. Quantification results are presented as mean±s.e.m. **P*<0.05; ***P*<0.01; ****P*<0.001 (two-tailed Student's *t*-test).

and 5'-CAATCCTCGCATTTTGGTGTCACCTCAGAGAAGATCTTCTC-TTTCA-3'. All new cDNA constructs were confirmed by DNA sequencing. Proteins were expressed in *E. coli* Rosetta2 (DE3) cells (Novagen). His-tagged GRASP55 was purified on HisPur Ni-NTA Resin (Pierce, Thermo Fisher Scientific) according to a standard procedure (Zhang et al., 2018).

Antibodies

The following antibodies were used for western blotting: rabbit anti-SIRT2 (1:1000; Cell Signaling, 12650), rabbit anti-GRASP55 (1:2000; Proteintech, 10598-1-AP), mouse anti-GM130 (1:2000; BD Transduction Laboratories, 610823), rabbit anti-GRASP65 (1:2000; a kind gift from Joachim Seeman, Department of Cell Biology, UT Southwestern Medical Center, Dallas, TX), rabbit anti-Golgin 160 (1:2000; Proteintech, 21193-1-AP), rabbit anti-Golgin 84 (1:3000; a kind gift from Ayano Satoh, Graduate School of Interdisciplinary Science and Engineering in Health Systems, Okayama University, Okayama, Japan), anti-FLAG M2-peroxidase clone M2 (1:10,000; Sigma, A8592), mouse anti-HA.11 clone 16B12 (1:1000; Covance, MMS-101R), mouse antimyc clone 4A6 (1:1000; Merck Millipore, 05-724), rabbit anti-acetylated lysine (AcK, 1:1000; Cell Signaling, 9441), mouse anti-His (1:5000; GE Healthcare, 27-4710-01), mouse anti-GFP (1:1000; Roche, 11814460001), rabbit anti-actin (1:10,000; Sigma, A2066). The following antibodies were used for immunofluorescence: rabbit anti-GRASP55 (1:200; Proteintech, 10598-1-AP), mouse anti-GM130 (1:200; BD Transduction Laboratories, 610823). Secondary HRP-conjugated antibodies for western blotting (Jackson ImmunoResearch) were used at 1:10,000 dilution. Secondary FITC- or

TRITC-conjugated antibodies for immunofluorescence (Jackson ImmunoResearch) were used at 1:100 dilution.

RNA interference

siRNA transfections were performed using Lipofectamine RNAiMax (Ambion, Life Technologies) according to the manufacturer's instructions. SIRT2 siRNAs used for immunofluorescence analysis and live-cell imaging were from Invitrogen (Silencer[®] Select; 5'-GCCCAAGTGTGAAGACTGT-3') and Dharmacon (SMARTpool ON-TARGETplus), respectively. Cells were transfected with siRNAs at a final concentration of 50 nM and assayed 72 h after transfection.

Mass spectrometry analysis of GRASP55 acetylation

GRASP55–GFP was transfected into HEK293T cells and immunoprecipitated using GFP-Trap magnetic beads (Chromotek). Bands were excised from silver-stained or Coommassie-stained gels and submitted to mass spectrometry analysis as described in Kaufmann et al., 2017, except that 20 mM iodoacetamide was used as alkylating reagent before digesting the proteins with trypsin (Trypsin Gold, Promega). Samples analyzed by parallel reaction monitoring (PRM) were spiked with 100 fmol Pierce Retention Time Calibration standard (PRTC, Thermo Fisher Scientific) per injection for quality control and retention time monitoring. PRM assay generation was performed using Skyline (MacLean et al., 2010), resulting in a scheduled assay with 12-min windows and not more than eight concurrent precursors per window. For data acquisition, we operated an Ultimate 3000 RSLC nano-flow chromatography system (Thermo Fisher Scientific), using a pre-column for sample loading (PepMapAcclaim C18, 2 cm×0.1 mm, 5 μ m, Thermo Fisher Scientific), and a C18 analytical column (PepMapAcclaim C18, 50 cm×0.75 mm, 2 μ m, Thermo Fisher Scientific), applying a linear gradient from 2% to 30% solvent B (80% acetonitrile, 0.1% formic acid; solvent A, 0.1% formic acid) at a flow rate of 230 nl/min over 60 min. Eluted peptides were analyzed on a Q Exactive HF Orbitrap mass spectrometer (Thermo Fisher Scientific), equipped with a Proxeon nano-spray-source (Thermo Fisher Scientific). PRM mass spectrometry parameters were: survey scan with 60k resolution, AGC 1E6, 50 ms IT, over a range of 380 to 1100 *m/z*; PRM scan with 60k resolution, AGC 2E5, 250 ms IT, isolation window of 0.7 *m/z* with 0.2 *m/z* offset, and NCE of 27%.

The spectral library for PRM data analysis in Skyline was generated as follows: a spectral library was prepared in Proteome Discoverer (Thermo Fisher Scientific, version 2.3) using MS Amanda (Dorfer et al., 2014) and Percolator (Kall et al., 2007) with the following search parameters: database compiled of the *Rattus norvegicus* reference Proteome (UniProt, April 2019) and common contaminants, 10 ppm MS1 and 20 ppm MS2 mass tolerance, carbamidomethylation of cysteine as fixed and lysine acetylation and methionine oxidation as variable modifications, three missed cleavages allowed, 1% false discovery rate (FDR) on PSM level. Results were imported into Skyline using a score cut-off of 0.95 and filtering for GRASP55 peptides.

Data analysis, including manual validation of all peptides and their transitions (based on MS2 spectra, retention time, relative ion intensities and mass accuracy), and relative quantification was performed in Skyline. Six of the most-intense non-interfering transitions of the target peptides were selected, and their peak areas were summed for peptide quantification (total peak area). As a quality control for instrument performance, PRTC peptides were monitored using MS1 full-scan filtering. Peak areas were exported from Skyline. To allow a relative quantitative comparison of K50 acetylation independently of GRASP55 abundance, we normalized the intensity of the peptide LNK(ac)DNDTLK to the intensity of its non-modified counterpeptide LNKDNDTLK. Although such an approach is usually problematic due to the missed tryptic cleavage on acetylated lysine residues, it is reasonable in this case because the aspartate residue in the +1 position suppresses cleavage to an extent large enough to provide a peptide signal one order of magnitude higher than in its acetylated form.

In vitro acetylation and deacetylation assays

In vitro acetylation assay was performed by incubating 5 µg human His_{6} -tagged GRASP55 with 0.5 µg of p300 acetyltransferase (Kaufmann et al., 2016) in 50 µl of 10 µM acetyl-CoA (Sigma), 50 mM Tris-HCl pH 8.0, 10% glycerol, 0.1 mM EDTA, 1 mM DTT and 1 mM PMSF for 2 h at 30°C with rotation. The acetylation reaction was stopped with 20 µM anacardic acid. Subsequent *in vitro* deacetylation assay was performed by adding 0.5 or 5 µg His-tagged SIRT2 WT or its catalytically inactive H150Y mutant (Kaufmann et al., 2016) and incubating for 90 min at 37°C. The acetylation levels of GRASP55 and p300 were detected by an anti-acetylated lysine antibody (AcK, Cell Signaling).

Immunoprecipitation

To determine the interaction between FLAG–SIRT2 and GRASP55–GFP, transfected HeLa cells were lysed at 4°C for 20 min in 20 mM Tris-HCl pH 8.0, 150 mM NaCl, 1% Triton X-100, 20 mM β -phosphoglycerol, 10 μ M trichostatin A (TSA; Sigma), 20 mM nicotinamide (NAM; Sigma) and protease inhibitors. Cell lysates were rotated with anti-GFP antibodies coupled to magnetic beads (GFP-Trap_M, Chromotek) for 2 h at 4°C. Beads were washed five times with the lysis buffer and eluted by boiling. Samples were then subjected to western blotting.

To determine the interaction between HA–SIRT2 and FLAG–GRASP55, cells were harvested by scraping and cell pellets were lysed in the lysis buffer containing 50 mM Tris-HCl pH 8.0, 150 mM NaCl, 1% Triton X-100, 50 U/ml benzonase (Novagen), protease inhibitors (Complete Mini Protease Inhibitor Cocktail Tablets, EDTA-free; Roche), 5 μ M TSA (Sigma) and 20 mM NAM (Sigma) for 1 h with rotation at 4°C. Anti-HA magnetic beads (Pierce) were equilibrated by washing the beads twice with Tris-buffered saline (TBS). Lysates were incubated with the beads for 2 h at 4°C with

rotation. Beads were washed three times with the lysis buffer and eluted with glycine pH 2.0, samples were then subjected to western blotting.

To determine the self-interaction efficiency of GRASP55 WT or its acetylation mutants K50R and K50Q, HeLa cells were transfected with the indicated plasmids, synchronized with nocodazole for 18 h, and lysed in 20 mM Tris-HCl, pH 8.0, 150 mM NaCl, 1% Triton X-100 and protease inhibitors. Lysates were cleared by centrifugation (16,873 *g* for 20 min), incubated with anti-GFP antibody overnight at 4°C, and subsequently incubated with protein A beads for another 2 h, re-isolated and analyzed by western blotting.

Immunofluorescence

Cells were grown on sterile glass coverslips, rinsed with PBS, fixed in 4% paraformaldehyde for 10 min and permeabilized with 0.3% Triton X-100 in PBS for 5 min. Cells were subsequently blocked in 0.2% gelatin in PBS for 15 min, incubated with primary antibodies for 45 min, and washed and probed with the appropriate secondary antibodies conjugated to FITC or TRITC for 45 min. DNA was stained with Hoechst 33342 for 5 min, rinsed with PBS, mounted with Mowiol onto slides and sealed with nail polish. Images of the samples were taken with a $63 \times$ oil objective on a Zeiss Observer Z1 epifluorescence microscope. Axiovert software was used for image acquisition and analysis.

Live-cell imaging

Cells were seeded in multi-well dishes or 6-channel slides suitable for high resolution microscopy (Ibidi) the day before imaging. Live-cell imaging was performed on an Axio Observer Z1 (Zeiss) equipped with an EM-CCD camera (Evolve EM-512) and environmental control.

Electron microscopy

EM was performed as previously described (Tang et al., 2010a). Briefly, HeLa cells were transfected with control and SIRT2 siRNA for 72 h, and then plated in six-well dishes. After another 24 h culture, cells were processed for Epon embedding. Sections of 60 nm were mounted onto Formvar-coated nickel grids and stained with 2% uranyl acetate for 5 min and 3% lead citrate for another 5 min. Grids were imaged using a JOEL transmission electron microscope. Images from >25 cells were captured at 10,000× magnification and were taken from the perinuclear region of the cell where Golgi membranes were normally concentrated. A cisterna was defined as a membrane-bound structure in the Golgi cluster whose length is at least four times its width, and the latter does not exceed 60 nm; a stack is the set of flattened, disk-shaped cisternae piling up together. The longest cisternae in a Golgi stack were measured as the length of cisternae using ImageJ, and the number of cisternae in the Golgi stack with the most cisternae layers was counted as the number of cisternae per stack (Wang et al., 2005). In our assays, defects in Golgi stack formation are reflected as (1) a reduced number of cisternae per stack, and (2) improper alignment of the cisternae within each stack accompanied with increased number of vesicles.

Quantification and statistics

Error bars represent standard error of mean (s.e.m.) estimated from two to four independent experiments. Statistical significance was calculated using a two-tailed *t*-test. *P*-values smaller than 0.05 were considered statistically significant and are indicated with asterisks (*P<0.05; **P<0.01; ***P<0.001). Fragmented Golgi was defined as disconnected or scattered dots. To quantify the percentage of fixed cells with fragmented Golgi, we counted more than 200 cells in each treatment. To quantify the percentage of live cells with fragmented Golgi, we counted more than 90 cells in each treatment. To determine the Golgi phenotype with EM, we counted the number of cisternae per stack or the length of Golgi cisternae from 20 different control RNAi cells or SIRT2 siRNA cells.

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Competing interests

The authors declare no competing or financial interests.

Author contributions

Conceptualization: X.Z., D.S., Y.W.; Methodology: X.Z., A.B., E.K., D.S.; Validation: X.Z., D.S., Y.W.; Formal analysis: X.Z., D.S., Y.W.; Investigation: X.Z., A.B., E.K., D.S., Y.W.; Data curation: X.Z., A.B., E.K., D.S.; Writing - original draft: X.Z., D.S., Y.W.; Writing - review & editing: X.Z., D.S., Y.W.; Visualization: X.Z., D.S.; Supervision: D.S., Y.W.; Project administration: Y.W.; Funding acquisition: D.S., Y.W.

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Supplementary information

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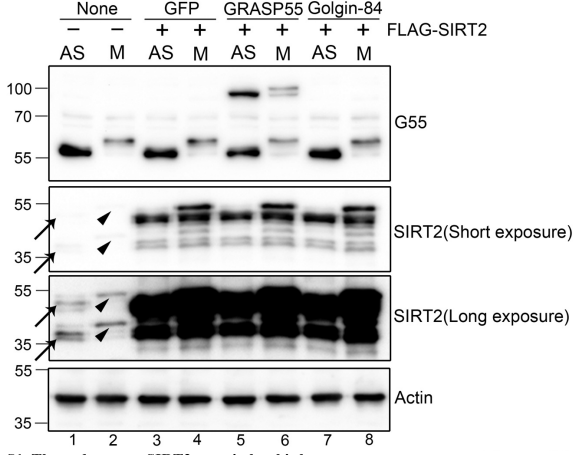


Figure S1. The endogenous SIRT2 protein level is low.

Asynchronous (AS) and mitotic HeLa cells or HeLa cells co-overexpressing GFP-tagged GRASP55 or Golgin-84 and FLAG-SIRT2 were lysed, blotted for GRASP55 (G55 in the figure) and SIRT2. Note that the endogenous SIRT2 level (lane 1 and 2) is very low compared to the overexpressed FLAG-SIRT2 (lane 3-8). Arrows indicate asynchronous SIRT2 bands, while arrowheads indicate mitotic SIRT2 bands.

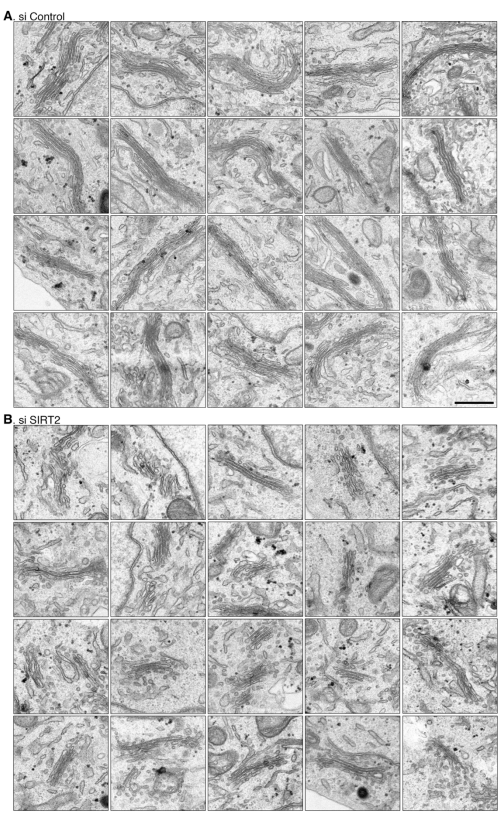
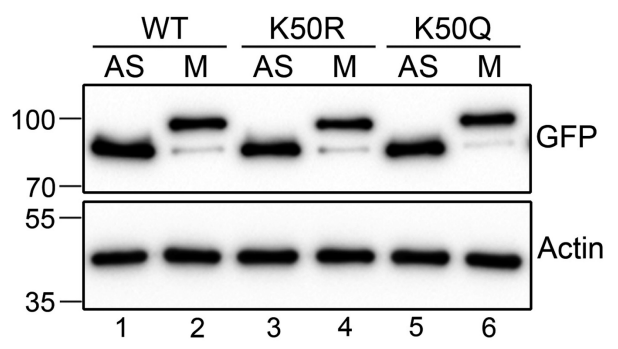


Figure S2. SIRT2 is required for proper Golgi structure maintenance.

HeLa cells were transfected with control (**A**) or SIRT2 (**B**) siRNA and analyzed by EM. Shown are galleries of EM images. The reduced average number or length of Golgi cisternae were frequently observed after SIRT2 depletion as displayed in **B**. Scale bar: 500 nm.



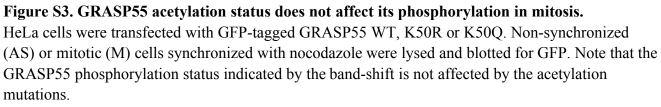
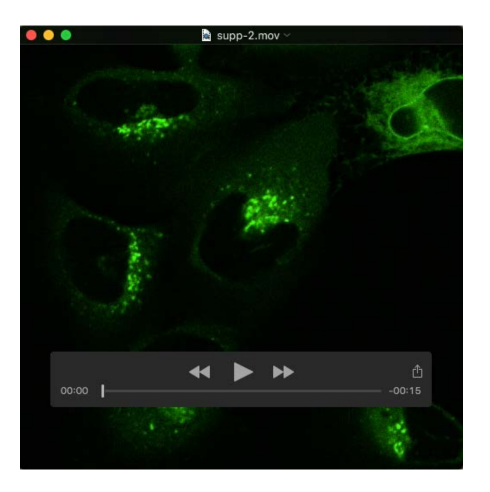


Table S1. Mass spectrometry analysis of GRASP55 post-translational modifications(acetylation, phosphorylation and ubiquitination) in asynchronous and mitotic HEK293T cellsexpressing GRASP55-GFP.

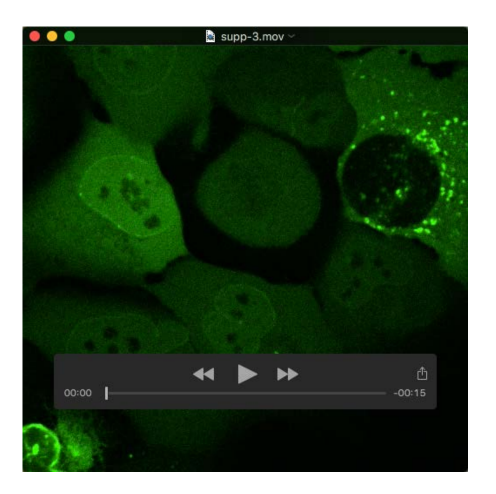
Click here to Download Table S1

Table S2. PRM (parallel reaction monitoring) mass spectrometry analysis of GRASP55-GFP K50 acetylation in mitotic HEK293T cells after silencing or overexpressing SIRT2.

Click here to Download Table S2



Movie 1. Live-cell imaging of GRASP55-GFP U2OS cells 72 h after transfection with si Control.



Movie 2. Live-cell imaging of GRASP55-GFP U2OS cells 72 h after transfection with si SIRT2.