

1 **Targeting fatty acid synthase suppresses tumor development in *NF2/CDKN2A*-deficient**
2 **malignant pleural mesothelioma**

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40 **Abstract**

41 Malignant pleural mesothelioma (MPM) is an uncommon yet deadly cancer linked to asbestos
42 exposure. The lack of effective early diagnosis and treatment leads to reduced life expectancy
43 among patients with MPM. This study is aimed to identify a novel molecular target inhibitor to
44 develop more effective therapeutics for MPM. Our drug screening assay showed that the fatty
45 acid synthase (FASN) inhibitor cerulenin demonstrates strong and selective anti-proliferative
46 properties against *NF2/CDKN2A(p16)*-deficient MPM cells, surpassing the effects of cisplatin
47 or pemetrexed. FASN protein is frequently detected in *NF2/p16*-deficient MPM tumor-derived
48 tissues (15/15, 100%), but rarely in *NF2/p16*-intact MPM tumors (8/25, 32%). Notably,
49 cerulenin administration successfully reduced the growth of *NF2/p16*-deficient MPM tumors in
50 xenografted mice. Cerulenin inhibits mitochondrial fission by targeting dynamin-related protein
51 1 (DRP1) in *NF2/p16*-deficient cells. Moreover, the disruption of the FASN gene leads to
52 increased ubiquitination of DRP1. These findings suggest that FASN might play a role in the
53 tumorigenesis of MPM cells through the regulation of mitochondrial dynamics. This research
54 offers a novel perspective on the potential development of precision medicine for MPM.

55

56 Keywords: CRISPR/Cas9, FASN, p-Drp1, cerulenin, MPM

57

58 **Abbreviations**

59 MPM, malignant pleural mesothelioma; IHC, immunohistochemistry; CRISPR, clustered
60 regularly interspaced short palindromic repeats; Cas9, CRISPR-associated protein 9; FASN,
61 Fatty acid synthase; DKO, double knockout; TKO, FASN-knockout DKO; MTR, MitoTracker;

62 Drp1, dynamin-related protein 1; OPA1, optic atrophy-1; PARP, Poly (ADP-ribose) polymerase;

63 ROS, reactive oxygen species; FCM, flow cytometry.

64

65

66 **Introduction**

67 Malignant pleural mesothelioma (MPM) is a highly aggressive neoplasm originating from

68 pleural mesothelial cells, commonly linked to asbestos exposure after a latency period of 30–40

69 years (1-3). Diagnosis of MPM typically occurs in advanced stages, contributing to a grim

70 prognosis for patients (4). MPM generally responds poorly to radiation and conventional

71 chemotherapy (5). Recent advancements in the management of the disease, including diagnosis,

72 staging biomarkers, and treatment strategies have provided greater understanding of MPM;

73 however, the mortality rate remains high, partially due to late diagnosis and treatment resistance

74 (6, 7).

75 Recent studies have revealed that MPMs are associated with frequent genetic

76 alterations in the neurofibromatosis 2 (*NF2*), cyclin-dependent kinase inhibitor 2A (*CDKN2A*,

77 *p16*), and BRCA1-associated protein 1 (*BAP1*) tumor suppressor genes (8-11). Previously, we

78 have observed high expression levels of *FGFR2*, *CD24*, and *CAMK2D* in *NF2*-knockout (KO),

79 *NF2/p16*-double KO (DKO), and *BAP1*-KO mesothelial cell lines, respectively (12-14),

80 suggesting representation of diagnostic and therapeutic targets for MPM. However, a library
81 containing 364 anticancer drugs was screened using *NF2/p16*-DKO human mesothelial cells to
82 precisely understand the molecular determinants of MPM, and cerulenin—a fatty acid synthase
83 (FASN) inhibitor was identified.

84 FASN—a key catalytic enzyme regulating long-chain fatty acid synthesis—is highly
85 expressed in several human cancers, including MPM (15-17). FASN activity is necessary for
86 mitochondrial priming (18) and senescence in cancer cells (19). FASN amplifies mitochondrial
87 ATP synthesis (20) and promotes mitochondrial fission, key processes in the reprogramming of
88 fatty acid metabolism in colon cancer cells (21). Mitochondria are highly pleomorphic and
89 considered primary mediators of intracellular energy production (22, 23). Alterations in
90 mitochondrial dynamics are associated with disease progression and drug resistance in various
91 cancers (24-27); however, the precise role of FASN in mitochondrial biology in MPM cells
92 remains unknown.

93 Inhibiting FASN activity prevents cancer cell proliferation, migration, invasion, cell
94 cycle, signaling pathway, and energy metabolism in breast and colon cancers, diffuse large B-
95 cell lymphomas, melanomas, retinoblastomas, prostate cancers, and mesothelioma cells (17, 28-
96 35). Cerulenin, which is available as a naturally occurring or pharmacologically synthesized
97 antibiotic, is an effective apoptosis inducer, with mitochondria being the key player in

98 cerulenin-mediated apoptosis (36). This occurs by disrupting the mitochondrial membrane and
99 dysregulating the mitochondrial membrane proteins (37). The preferential activity of FASN and
100 its inhibitor cerulenin are attractive targets for anticancer therapy; however, the underlying
101 mechanism of cerulenin-mediated FASN inhibition and cancer cell suppression in MPM
102 remains unclear.

103

104 This study aimed to determine promising diagnostic and therapeutic targets for
105 *NF2/p16*-deficient MPM by FASN and its effect on mitochondrial dynamics. Additionally,
106 considering that cerulenin exhibits a selective antiproliferative effect on MPM cells, its role as
107 potential molecular-targeted anticancer drug was determined.

108

109 **Results**

110 ***The FASN inhibitor cerulenin selectively suppresses the proliferation of NF2-p16-deficient***
111 ***MPM cells***

112 Given that we have previously shown differential gene expression between control and
113 *NF2/p16*-double knockout (DKO) isogenic human mesothelial cell clones, we attempted to
114 identify an inhibitor that specifically suppresses the proliferation of *NF2/p16*-DKO cells, which
115 are frequently observed in MPM (38). To this end, we performed the MTT assay with the

116 Screening Committee of the Anticancer Drugs library, which comprises 364 chemical
117 compounds (Supplementary Table S1). Our drug screening assay showed that the FASN
118 inhibitor cerulenin, but not C75, exhibited the most potent antiproliferative activity against
119 *NF2/p16*-DKO cells (Fig. 1a). To further evaluate cerulenin's impact on MPM cell proliferation,
120 we conducted MTT assays using various mesothelial and MPM cell lines. These included
121 *NF2*^{+/+}/*p16*^{+/+} (parental MeT-5A and HOMC-B1 cells), *NF2*^{-/-}/*p16*^{+/+} (Y-MESO-14),
122 *NF2*^{+/+}/*p16*^{-/-} (NCI-2452, ACC-MESO-4, Y-MESO-9, and MSTO-211H), and *NF2*^{-/-}/*p16*^{-/-}
123 cells (MeT-5A-DKO, HOMC-B1-DKO, NCI-H290, NCI-H2052, and MSTO-211H-DKO).
124 Interestingly, the average IC₅₀ values in the *NF2/p16*-DKO mesothelial cell lines and MSTO-
125 211H cells ranged from 5 μM to 7.5 μM, which is lower than that in the *NF2*^{+/+}/*p16*^{+/+} cell lines
126 (approximately 20 μM) (Fig. 1b). This outcome indicates that cerulenin selectively inhibits the
127 proliferation of *NF2/p16*-DKO cells. Compared with the antiproliferative effect of cerulenin,
128 treatment with pemetrexed or cisplatin was relatively ineffective even at higher doses against
129 both the *NF2*^{+/+}/*p16*^{+/+} and *NF2*^{-/-}/*p16*^{-/-} MPM cell lines (Fig. 1c). Furthermore, cerulenin
130 treatment suppressed the proliferation of MPM cell lines NCI-H290 and NCI-H2052, both of
131 which spontaneously lacked *NF2/p16* expression (Fig. 1d). These findings indicate that
132 cerulenin holds potential as a valuable molecular target for anticancer therapy in MPM.
133

134 ***FASN expression is inversely associated with NF2/p16 expression in human MPM tissues***

135 Immunohistochemistry was performed using 45 MPM tissue samples and 2 mesothelium

136 samples to assess FASN expression in mesothelioma (Supplementary Table S2). Microscopic

137 analyses revealed 3 strong (3⁺), 14 moderate (2⁺), and 11 weak (1⁺) FASN-positive signals in the

138 MPM tissues (Fig. 2a; Supplementary Table S2), whereas no positive signal was observed in the

139 normal mesothelium (Fig. 2b). Additionally, the FASN positivity rate was higher in the

140 NF2/p16-negative cases (15/15 tissues, 100%) than in the NF2/p16-positive cases (8/25 tissues,

141 32%; Fig. 2b). Notably, MPM patients demonstrating elevated FASN expression levels

142 experienced notably shorter overall survival compared to those with lower FASN expression

143 levels (Fig. 2c). These findings strongly imply a close correlation between FASN expression and

144 poorer prognosis in MPM patients.

145

146 ***Cerulenin suppresses tumor growth of NF/p16-deficient MSTO-211H cells in vivo***

147 The effect on the growth of NF2/p16-deficient MSTO-211H (NF/p16-DKO MSTO-211H)

148 tumors was examined in SCID mice to assess the potential of cerulenin for clinical application.

149 The intraperitoneal injection of cerulenin notably inhibited tumor growth in comparison to the

150 vehicle control (see Figs. 3a and 3b). Furthermore, cerulenin treatment did not result in any loss

151 of body weight (see Fig. 3c). Subsequent assessment of cerulenin's safety in normal BALB/c

152 mice revealed no noteworthy alterations in body weight (data not presented) or
153 histopathological changes in the heart, liver, or kidneys on Day 14 post-treatment (see Fig. 3d).
154 Furthermore, biochemical analyses showed no alterations in blood parameters, including levels
155 of aspartate aminotransferase (AST) and alanine aminotransferase (ALT), following cerulenin
156 administration (see Fig. 3e; Supplementary Table S3). These findings indicate the potential
157 safety and efficacy of cerulenin as a molecular-targeted anticancer medication for managing
158 patients with NF2/p16-deficient MPM.

159

160 ***Cerulenin accelerates the mitochondrial fusion processes and induces apoptosis in NF2/p16-***
161 ***DKO cells***

162 Growing evidence indicates FASN's role in tumor proliferation through the regulation of energy
163 metabolism. Consequently, we investigated cerulenin's impact on mitochondrial morphology in
164 *NF2/p16*-DKO cells. We assessed the morphology of the mitochondria using MitoTracker,
165 which permanently binds to mitochondria regardless of cell survival. Confocal microscopy
166 showed that mitochondrial hyperfusion (long tubular network structure) increased after
167 cerulenin treatment in *NF2/p16*-DKO MeT-5A and HOMC-B1 mesothelial cells, whereas no
168 obvious change was observed in the parental cells (Fig. 4a). Similarly, mitochondrial
169 hyperfusion increased after cerulenin treatment in parental MSTO-211H (spontaneously

170 *CDKN2A/p16*-deficient) cells, which was further enhanced in the *NF2/p16*-DKO MSTO-211H
171 cells (Fig. 4a). We examined whether cerulenin influences the phosphorylation status of
172 dynamin-related protein 1 (DRP1), a critical regulator of mitochondrial fission, in *NF2/p16*-
173 DKO cells. Immunofluorescence analysis revealed elevated levels of DRP1 phosphorylation in
174 all *NF2/p16*-DKO cells compared to parental cells; however, cerulenin treatment mitigated
175 these increases in DRP1 phosphorylation levels (see Fig. 4a). Additionally, Western blot
176 analysis demonstrated that cerulenin reduced the phosphorylation levels of both Akt and DRP1
177 (see Fig. 4b). We unexpectedly found that the FASN protein levels were relatively higher in all
178 the *NF2/p16*-DKO cells than in the parental cells tested (Fig. 4b). In fact, *NF2/p16* loss exhibited
179 high FASN expression levels in the MPM cells tested, as compared with NF2- and/or p16-
180 positive MPM cells (Supplementary Fig. S1). To assess cerulenin's impact on the apoptosis of
181 *NF2/p16*-double knockout (DKO) cells, we conducted flow cytometry analysis using Annexin V
182 (AxV) and propidium iodide (PI) double staining. The results revealed a significant increase in
183 the Ax⁺/PI⁺ population in *NF2/p16*-DKO mesothelial and MPM cells following cerulenin
184 treatment compared to parental cells (see Fig. 4c), suggesting a potential heightened sensitivity
185 of *NF2/p16*-DKO cells to cerulenin. Similarly, western blot analysis revealed that the cleaved
186 poly (ADP-ribose) polymerase (PARP) levels markedly increased, whereas both FASN protein
187 and phosphor-AKT levels decreased after cerulenin treatment in *NF2/p16*-DKO cells (Fig. 4b).

188 Furthermore, our gene set enrichment analysis (GSEA) demonstrated a significant suppression
189 of PI3K-AKT-MTOR signaling in NF2/p16-DKO cells treated with cerulenin (see Fig. 4d).
190 These findings strongly indicate that cerulenin selectively triggers apoptosis by targeting
191 oncogenic AKT signaling in NF2/p16-DKO cells.

192

193 ***FASN knockout (TKO) suppresses cell proliferation and DRP1 activity in NF2/p16-DKO***

194 ***MeT-5A cells***

195 *FASN knockout* (hereafter called triple knockout “TKO”) cell clones #1 and #2 were generated
196 using previously established *NF2/p16*-DKO MeT-5A cell clones by targeting exon 3 of the
197 *FASN* gene to further clarify the role of *FASN* in *NF2/p16*-DKO mesothelial cells
198 (Supplementary Figs. 2a and b). As expected, the *FASN* protein expression levels were
199 undetected in the TKO #1 and #2 cells (Supplementary Fig. S2c). Further, effect on the
200 proliferation rate by the disruption of the *FASN* gene was tested. The MTT assay indicated a
201 notably reduced proliferation rate in the TKO cell clones (#1 and #2) compared to the DKO cell
202 clones (see Fig. 5a). Furthermore, Western blot analysis revealed a decrease in Akt and DRP1
203 phosphorylation levels, alongside elevated levels of cleaved PARP and caspase-3 in the TKO
204 cell clones relative to the DKO cell clones (see Fig. 5b). Additionally, OPA1, MFN1, and MFN2,
205 which positively regulate mitochondrial fusion, increased at the protein levels in the TKO cell

206 clones (Fig. 5b). Therefore, the effect of *FASN* knockout on the mitochondrial morphology
207 under *NF2/p16*-deficient conditions was examined. Confocal microscopy showed a significant
208 increase in the elongated and tubular structure of the mitochondria in the TKO cells as
209 compared with the DKO cells (Fig. 5c). The protein expression of DRP1 decreased consistently,
210 whereas that of OPA1 increased in the TKO cell clones (Fig. 5d). The effect of proteasome
211 inhibition on DRP1 protein expression was examined to uncover the molecular mechanism
212 underlying the DRP1 expression change in the TKO cells. Western blot analysis demonstrated
213 that both the phosphorylation and protein expression levels of DRP1 were reinstated following
214 treatment with the proteasome inhibitor MG-132 in the TKO cells (see Fig. 5e). Furthermore,
215 the immunoprecipitation assay revealed a higher DRP1 ubiquitin level in the TKO cells than in
216 the DKO cells (Fig. 5f). The intracellular reactive oxygen species (ROS), largely generated in
217 the mitochondria, increased in the TKO cells (Fig. 5g), suggesting involvement of *FASN*
218 expression in the mitochondrial activity, which might affect mitochondrial ROS generation in
219 *NF2/p16*-DKO cells.
220

221 **Discussion**

222 This study investigated the involvement of *FASN* in MPM cell proliferation and revealed the
223 selective antiproliferative effect of the *FASN* inhibitor cerulenin in *NF2/p16*-deficient human

224 MPM cells. The results revealed that loss of *NF2/p16* sensitizes the human mesothelial and
225 MPM cells to cerulenin as compared with the *NF2*- and/or *p16*-intact cells.
226 Immunohistochemical analysis also showed detection of FASN expression in *NF2/p16*-negative
227 human mesothelioma tissues. The analysis of public data revealed that the overall survival was
228 significantly shorter for patients with MPM exhibiting a higher *FASN* expression level.
229 Moreover, cerulenin suppressed the tumor growth of *NF2/p16*-deficient human MPM cells *in*
230 *vivo* without any relevant toxicities.
231 FASN has been reported as a therapeutic target and an oncogene in several cancers,
232 including breast, pancreatic, and colorectal cancers (38-43). FASN overexpression is correlated
233 with poor prognosis in HER2-positive ovarian and gastric cancers (44, 45). FASN
234 overexpression was observed in a subset of MPM cell lines and human MPM tissues in this
235 study. Additionally, FASN expression was preferentially detected in the *NF2/p16*-deficient
236 mesothelial and MPM cell lines as compared with the *NF2/p16*-intact mesothelial and MPM cell
237 lines. MPM patients with higher FASN expression exhibited significantly shorter overall
238 survival than those with lower FASN expression. Our previous study reported that the loss of
239 *NF2* and *p16* genes significantly enhances anchorage-independent growth and epithelial–
240 mesenchymal transition phenotype in mesothelial cells, in which the cancer-stem cell marker
241 CD24 is upregulated in *NF2/p16*-deficient mesothelial and MPM cell lines (46, 13). Although

242 the association between *NF2/p16* deficiency and FASN overexpression remains unclear, the
243 retardation of the proliferation of *NF2/p16*-deficient cells by FASN disruption indicates the
244 possibility of important role of FASN in *NF2/p16*-deficient MPM cell survival.

245

246 In our investigation, we observed that the upregulation of FASN expression coincides with
247 the concurrent phosphorylation of oncogenic Akt in *NF2/p16*-DKO cells. Wagner et al.
248 previously reported that the FASN inhibitor impairs receptor-PI3K-mTORC1 signaling in
249 ovarian cancer cells (47). They also showed that FASN inhibitors impair the
250 phosphatidylinositol 3,4,5-trisphosphate (PIP3) levels, diacylglycerol (DAG), and subsequent
251 PI3K-AKT suppression (47). Another report showed that sensitivity to FASN inhibitors is
252 driven by DAG level reductions and DAG-PKC signaling pathway impairments (48). We found
253 that cerulenin significantly inactivates the PI3K–Akt–mTOR signaling pathway in *NF2/p16*-
254 DKO mesothelial cells. Although it is still unclear whether the membrane lipid composition
255 changes in the *NF2/p16*-DKO cells, the loss of *NF2/p16* may enhance the proliferation of MPM
256 cells via FASN-mediated PI3K-Akt signaling. Indeed, *FASN* knockout in *NF2/p16*-DKO cells
257 delays proliferation and decreases Akt phosphorylation as compared with the *FASN*-intact DKO
258 cells. Therefore, it is possible that cerulenin suppresses PI3K-Akt signaling by modulating lipid
259 membrane biogenesis.

260 The mitochondria are the main organs involved in the generation of cellular fuel ATP
261 through several metabolic processes, including lipid catabolism. Besides, the equilibrium
262 between mitochondrial fission and fusion occurs for the maintenance of mitochondrial functions,
263 which also plays an important role in both normal cells and cancer progression (49-56). In this
264 current study, the absence of FASN in the DKO (TKO) MeT-5A cells led to an increase in
265 elongated and tubular structured mitochondria, indicative of mitochondrial fusion. Similarly,
266 cerulenin treatment induces mitochondrial fusion formation in *NF2/p16*-deficient (DKO) MPM
267 cells. The absence of FASN, as well as FASN inhibition, notably elevated the count of apoptotic
268 cells in DKO MPM cells. These findings imply that FASN plays a crucial role in sustaining the
269 survival of MPM cells through its involvement in mitochondrial activity. At a molecular level,
270 both the expression and phosphorylation of DRP1 decreased in TKO MeT-5A cells and
271 cerulenin-treated DKO cells. Furthermore, the depletion of FASN in TKO cells resulted in
272 increased protein expression of OPA1, MFN1, and MFN2, all of which promote mitochondrial
273 fusion. Interestingly, proteasome inhibitor MG-132 restored DRP1 protein expression in the
274 TKO cells. Since the ubiquitinated DRP1 level is elevated in TKO cells compared to DKO cells,
275 it suggests a potential role of FASN in stabilizing DRP1 at the protein level. Given that FASN-
276 mediated fatty acids are converted to phospholipids (e.g., cardiolipin) that comprise the
277 mitochondrial membrane, it would be of interest to examine whether cerulenin affects the lipid

278 composition of mitochondrial membranes in MPM.

279 The present study has some limitations. First, the FASN protein expression was shown
280 more frequent in NF2- and/or p16-negative MPM tissues as compared in both positive tissues;
281 however, the somatic mutations of *NF2* and/or *p16* genes in the corresponding tissues was not
282 investigated. Therefore, further studies are required to clarify the association between FASN
283 expression and functional status of *NF2* and/or *p16* in MPM. Second, the antitumor effect of
284 cerulenin was demonstrated using only one human MPM cell line, MSTO-211H. Hence,
285 additional tests using primary human MPM-derived tumors are required for evaluating the
286 efficacy of targeting FASN therapy in MPM.

287 In conclusion, our genome-edited human mesothelial cell model identified that the FASN
288 inhibitor cerulenin preferentially exhibits an antiproliferative effect on MPM cells lacking NF2
289 and/or p16. These results underscore the potential clinical utility of cerulenin in crafting
290 molecular-targeted therapies for combating MPM.

291

292 **Materials and methods**

293 ***Cell culture***

294 Two immortalized normal human mesothelial cell lines, MeT-5A (pleural mesothelial) and
295 HOMC-B1 (omental mesothelial; epithelioid type), along with eight human mesothelioma cell

296 lines—ACC-MESO-4, Y-MESO-12, Y-MESO-14, Y-MESO-9, NCI-H2452, MSTO-211H, NCI-
297 H290, and NCI-2052—were generously provided by Dr. Y. Sekido from the Division of
298 Molecular Oncology, Aichi Cancer Center Research Institute (Nagoya, Japan). The HOMC-B1
299 cell lines were maintained 18s according to previously described protocols. The ACC-MESO-4,
300 Y-MESO-12, Y-MESO-14, Y-MESO-9, NCI-H2452, MSTO-211H, NCI-H290, and NCI-2052
301 cell lines were cultured in RPMI-1640 medium (Wako Pure Chemical Industries, Ltd., Osaka,
302 Japan) supplemented with 10% fetal bovine serum (Sigma) and 1% penicillin-streptomycin
303 (Wako Pure Chemical Industries, Ltd.) at 37°C in a 5% CO₂ humidified atmosphere.

304

305 ***Gene KO using the CRISPR/Cas9 system***

306 We employed the CRISPR/Cas9 system to disrupt NF2 expression in the MSTO-211H cell line
307 and FASN expression in the DKO cells, following established procedures (12). The
308 pSpCas9(BB)-2A-GFP (PX458) plasmid was generously provided by Feng Zhang (plasmid
309 #48138; Addgene, Watertown, MA, USA) (58). Briefly, sgRNA sequences were chosen using an
310 optimized CRISPR design tool (<http://crispr.mit.edu/>). The selected sgRNA sequences for NF2
311 and FASN were 5' -AACACATCTCGTACAGTGACA-3' in exon 8 and 5' -
312 CCTTCAGCTTGCCGGACCGC-3' in exon 3, respectively. Plasmids expressing hCas9 and
313 sgRNA were generated by ligating oligonucleotides into the BbsI site of PX458 (NF2/PX458 or

314 FASN/PX458). To establish a knockout (KO) clone, 1 μ g of NF2/PX458 plasmid was
315 transfected into the MSTO-211H cell line, and 1 μ g of FASN/PX458 plasmid was transfected
316 into MeT-5A-DKO#1 cells (1×10^6 cells) using a 4D-Nucleofector instrument (Lonza Japan,
317 Tokyo, Japan). After 3 days, GFP-expressing cells were sorted using fluorescence-activated cell
318 sorting (BD FACSAriaTM III Cell Sorter; BD Biosciences, San Jose, CA, USA). A single clone
319 was selected, expanded, and utilized for the biological assays.

320

321 ***Cell growth assay***

322 The growth rate of the cells was assessed using the MTT assay (59). Briefly, cells (1×10^3
323 cells/well) were seeded into 96-well plates and cultured for the specified durations.
324 Subsequently, 10 μ L of MTT solution (5 mg/mL; Sigma-Aldrich) was added to each well, and
325 the cells were incubated for 4 hours. Following this, cell lysis buffer was added to dissolve the
326 resultant colored formazan crystals. The absorbance at 595 nm was measured using a
327 SpectraMAX M5 spectrophotometer (Molecular Devices, Sunnyvale, CA, USA).

328

329 ***Western blot analysis***

330 Western blotting was conducted following established protocols (13, 60). The antibodies utilized
331 are detailed in Table S4. Immune complexes were visualized using ImmunoStar LD (Wako Pure

332 Chemical Industries, Ltd.) and imaged with a LAS-4000 image analyzer (GE Healthcare, Tokyo,

333 Japan).

334

335 ***Annexin V assay***

336 Cells were plated into six-well culture plates (5×10^5 cells/well) and treated with cerulenin

337 (7.5 $\mu\text{g/mL}$) for 48 hours. Subsequently, the cells were exposed to annexin V (Ax)-FITC and

338 propidium iodide (PI) (10 $\mu\text{g/mL}$) at 25°C for 15 minutes. The fluorescence intensities were

339 quantified using fluorescence-activated cell sorting (FACS) analysis (LSRFortessa X-20 Flow

340 Cytometer, BD Biosciences,, Franklin Lakes, NJ, USA).

341

342 ***Immunofluorescence***

343 The cells were cultured on glass coverslips and fixed with a 4% paraformaldehyde solution for

344 20 minutes at 25°C. Subsequently, the cells were permeabilized with phosphate-buffered saline

345 (PBS) containing 0.1% Triton X-100, blocked in PBS containing 7% serum for 30 minutes, and

346 then incubated with primary antibodies followed by Alexa Fluor-conjugated secondary

347 antibodies (Invitrogen). Cell staining was conducted using MitoTracker (stock solution 1 mM;

348 diluted at 1:10,000) for 1 hour at 37°C to visualize the mitochondria. Following staining, the

349 cells were washed with PBS and fixed with cold paraformaldehyde (3.2% in PBS) for 20

350 minutes at room temperature. After additional washing steps, the samples were mounted using
351 PermaFluor, and images were captured using the FLUOVIEW FV3000 Series of Confocal Laser
352 Scanning Microscopes.

353

354 ***Immunohistochemistry***

355 Immunohistochemical analysis was conducted following previously established protocols (61).
356 A human mesothelioma tissue array (MS-1001a; US Biomax, Rockville, MD, USA) was
357 utilized. The tissue sections were incubated with primary antibodies (NF2, p16, and FASN
358 antibody, 2 µg/mL). Negative controls were established using normal rabbit immunoglobulin G
359 or by omitting the primary antibody. Immunoreactivity was independently assessed by two
360 investigators (S.K. and H.M.), and staining intensity was graded as strong (+3), moderate (+2),
361 weak (+1), or negative (0).

362

363 ***Xenograft experiments***

364 All animal experiments were conducted in accordance with the protocols (approval number-
365 2022-20) approved by the ethical committee of Aichi Medical University and followed
366 established guidelines. Female Fox Chase SCID mice (CB17/Icr-Prkdcscid/IcrIcoCrl) were
367 procured from CLEA Japan Inc. (Tokyo, Japan) and housed at the Institute of Animal

368 Experiments in Aichi Medical University under pathogen-free conditions. The mice, aged 6–8
369 weeks and weighing 17–18 g each, were utilized for the study. Subcutaneous injection of
370 MSTO-211H-DKO cells (6×10^6 cells) was performed in the SCID mice. Upon reaching a
371 tumor volume of approximately $\sim 100 \text{ mm}^3$, the mice were randomly assigned to two groups
372 (treatment and control). Cerulenin (20 mg/kg, administered three times per week) was
373 intraperitoneally administered to mice in the treatment group, while PBS served as the vehicle
374 control in the control group. Tumor measurements were taken every 3–4 days, and tumor
375 volume was calculated using the modified ellipsoid formula ($1/2 \times \text{length} \times \text{width}^2$).

376

377 ***DCFH-DA-based DCF assay***

378 Cells were plated into six-well culture plates at a density of 3×10^5 cells per well. The following
379 day, they were exposed to 10 μM DCFH-DA (2',7'-dichlorodihydrofluorescein diacetate) in the
380 medium for 45 minutes in the absence of light. Subsequently, the DCFH-DA solution was
381 removed, and the cells were washed with PBS. After detachment using trypsin, the cells were
382 suspended in media, centrifuged at 1,000 rpm for 5 minutes, and resuspended in PBS in 0.5-mL
383 tubes, which were then placed on ice. The samples were analyzed via FACS using LSRII Fortessa
384 X-20 Flow Cytometer (BD Biosciences).

385

386 **Statistical analysis**

387 The data are presented as mean \pm SE values. Statistical significance among groups was assessed
388 using one-way analysis of variance followed by Dunnett's comparison. All statistical analyses
389 were conducted using SPSS 23.0 software (SPSS Inc., Chicago, Illinois, USA).

390

391 **Competing interest**

392 This publication has no conflicts of interest, and there has been no substantial financial support
393 for this work that could have influenced its outcome. The manuscript has been reviewed and
394 endorsed by all named authors, and there are no other individuals who meet the criteria for
395 authorship but are not listed. All authors have consented to the order in which they are listed in
396 the manuscript.

397

398 **Availability of data and materials**

399 All accessible data are provided either in the main manuscript or in the supplementary materials.
400 Complete and unaltered western blot data utilized in the study are available in the
401 supplementary information. Furthermore, specific inquiries, data, and materials can be obtained
402 upon reasonable request to the corresponding author.

403

404 **Ethical standards**

405 The research conducted adhered to the ethical standards outlined by the Japanese Ministry of
406 Health, Labour, and Welfare.

407

408 **Author contributions**

409 The study concept was devised by S.K. and A.O. oversaw project acquisition and supervision.
410 S.K., M.L.R., M.N.H., and M.W. formulated the statistical analysis plan and performed the
411 statistical analyses. Data and resources were acquired by S.K., H.M., L.Q.V., I.H., M.T.A.S.,
412 N.J., A.I., M.R., H.I., Y.K., Y.L., T.H., H.K., S.T., and Y.H. Acquisition of funding was
413 facilitated by S.K. and A.O. contributed to manuscript composition. All authors critically
414 reviewed and edited the manuscript draft and consented to the final version for publication.

415

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605

606

607 **Figure legends**

608 **Fig. 1. Identification of FASN inhibitor cerulenin against the DKO-deficient cells and its**
609 **effect on MPM cell line survival**

610 (a) Evaluation of compounds from the Screening Committee of Anticancer Drugs library. MeT-
611 5A cells (DKO and NF2/p16-WT [parental] cells) were exposed to 364 chemical compounds
612 (10 μ M each). Cell survival rates were normalized to the mean optical densities of untreated
613 cells (set as 100%). Results are depicted as differences in cell survival percentages between
614 NF2/p16-WT and DKO cells. Red spots indicate >40% reduction in cell viability in DKO cells
615 compared to NF2/p16-WT cells. (b) Assessment of the impact of the FASN inhibitor cerulenin
616 on MPM cell line survival. Cell survival rates were calculated as described previously. (c)
617 Examination of cisplatin, pemetrexed, and cerulenin effects on the survival of various MPM cell
618 lines (MeT-5A, HOMC-B1, Y-MESO-14, Y-MESO-12, NCI-H2452, ACC-MESO-4, Y-MESO-
619 9, MSTO-211H, MeT-5A-DKO, HOMC-B1-DKO, NCI-H290, NCI-H2052, and MSTO-211H-
620 DKO) after incubation with indicated drug concentrations (20, 15, 10, 7.5, 5, 2.5, 1.25, 0.625,
621 and 0 μ M) for 72 hours. MTT assays were conducted following the manufacturer's guidelines,
622 and absorbance was measured at 595 nm using a spectrophotometer. Cell survival percentages
623 were calculated accordingly. Cisplatin, pemetrexed, and cerulenin are denoted by red, black, and
624 green lines, respectively. Data are presented as mean \pm standard error (n = 3).

625

626 **Fig. 2. Immunohistochemical (IHC) analysis of FASN expression**

627 (a) Illustrative IHC findings demonstrating FASN expression in two MPM tissues (top panels,
628 cases 5 and 10) and two normal mesothelial tissues (bottom panel, cases 46 and 47). (b)
629 Summary of IHC outcomes in MPM tissues. The bar graph indicates the percentage of total
630 MPM cases exhibiting FASN expression by NF2/p16-positive or NF2/p16-negative cases. (c)
631 Kaplan-Meier analysis was performed to assess the impact of FASN expression on overall
632 survival in TCGA mesothelioma patients, sourced from the UCSC Xena database. Fluorescence
633 values above and below the median were defined as high (red) and low (blue) FASN expression,
634 respectively. Scale bar = 60 μ m.

635

636 **Fig. 3. Effect of cerulenin on the growth of MSTO-211H-DKO tumor cells in vivo**

637 The MSTO-211H-DKO cells (5×10^6 cells/mouse) were injected subcutaneously into SCID
638 mice. Following tumor establishment (day 0), cerulenin (20 mg/kg body weight) or PBS vehicle
639 was administered intraperitoneally on days 0, 2, 5, 8, 10, 12, 15, and 17. (a) A representative
640 image depicting tumor-bearing xenografted mice in each group. (b-c) Line graphs illustrating
641 (b) relative tumor volume and (c) mouse body weight (in grams) during cerulenin treatment.
642 Tumor volume is presented relative to the size at day 0 (defined as 100%). Data are presented as
643 mean \pm SE (n = 5). *p < 0.05. (d) Representative histochemical images of the heart, liver, and
644 kidney from BALB/c mice after the 14-day observation period (Hematoxylin and Eosin staining,
645 magnification, $\times 100$; scale bar = 100 μ m). (e) Alterations in blood chemistry (AST, ALT, TG,
646 and LDH) following cerulenin administration compared with control data.

647

648 **Fig. 4. Role of cerulenin in mitochondrial morphology and apoptosis in DKO-deficient**
649 **MPM cells**

650 (a) The mitochondria were visualized using the MitoTracker probe (red). Nuclei were stained
651 with Hoechst (blue), in MeT-5A (upper left panel), HOMC-B1 (upper center panel), and MSTO-
652 211H (upper right panel). Cells were immunostained with Drp1 antibody (red) and Hoechst

653 (blue) in the mitochondria, as depicted in the lower left panels (MeT-5A), lower center panels
654 (HOMC-B1), and lower right panels (MSTO-211H). Two representative magnified view has
655 been illustrated in bottom part. **(b)** Protein expression of FASN, p-AKT, AKT, p-Drp1, Drp1, c-
656 PARP, and GAPDH analyzed by Western blotting in MeT-5A, MeT-5A-DKO, HOMC-B1,
657 HOMC-B1-DKO, MSTO-211H, and MSTO-211H-DKO cells treated with cerulenin (7.5 μ M)
658 for 48 h. **(c)** Flow cytometry analysis. Representative results of AxV-PI-based staining are
659 presented on the left. The graphs on the right display the percentage of AxV⁺/PI⁺ apoptotic cells
660 following cerulenin treatment (7.5 μ M) for 48 h measured using LSRFortessa X-20 Flow
661 Cytometer (BD Biosciences). Data are mean \pm SE (n = 3). Asterisks denote significant
662 differences between the DKO-deficient cells (MeT-5A and HOMC-B1 cells and MSTO-211H
663 cells) (*p < 0.05). **(d)** Gene set enrichment analysis (GSEA). MeT-5A, MeT-5A-DKO, HOMC-
664 B1, and HOMC-B1-DKO cell clones were treated with cerulenin (7.5 μ M) or control solvent for
665 48 h. A cDNA microarray analysis (Agilent) was conducted to obtain a comprehensive gene
666 expression profile. Raw data were analyzed using GSEA with HALLMARK gene sets (C1).

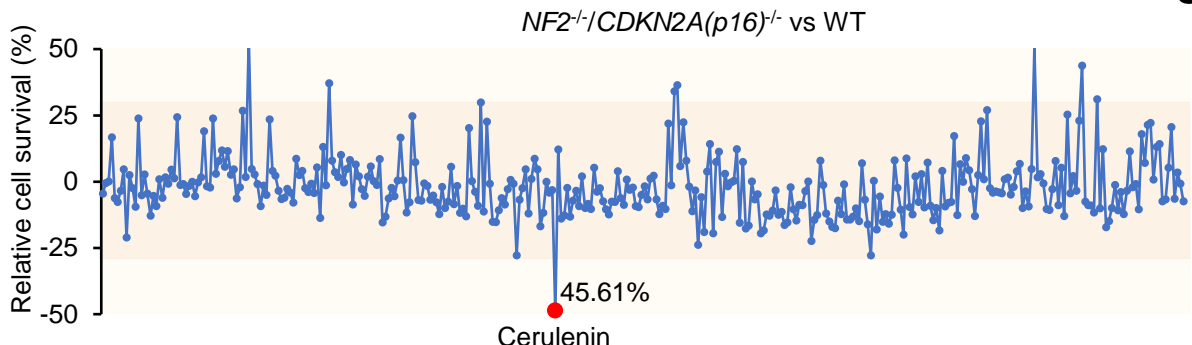
667

668 **Fig. 5. Disruption of FASN has adverse effects on mitochondrial morphology and**
669 **dynamics in MPM cells**

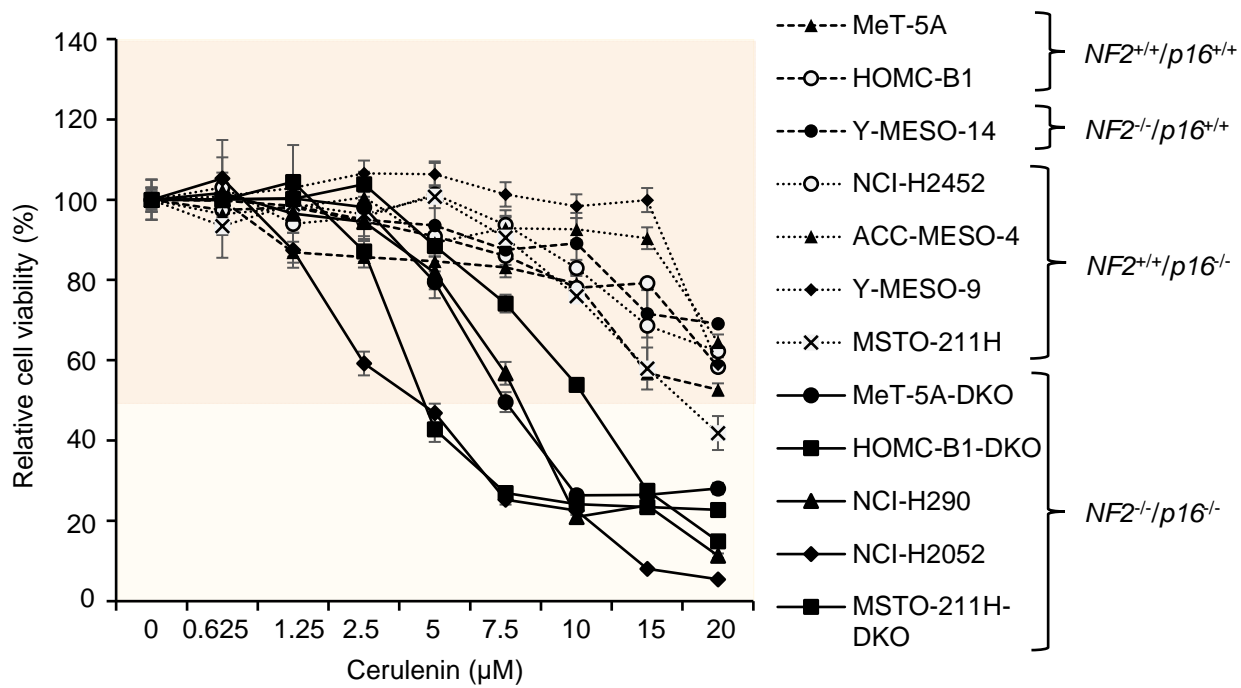
670 Cellular phenotype of MeT-5A-DKO/fatty acid synthase (DKO/FASN) triple knockout (TKO)
671 in MeT-5A cells. **(a)** MTT assay of DKO (#1 and #2) and TKO (#1 and #2) cell clones in MeT-
672 5A cells. The optical density (OD; 595 nm) at each time point (days 0, 1, 3, and 5) is presented
673 as the mean \pm SEM (n = 6). Growth ratios are expressed relative to the optical densities detected
674 at day 0, arbitrarily defined as 1. **(b)** Western blot analysis of FASN, p-AKT, AKT, p-Drp1,
675 Drp1, Opa1, MFN1, MFN2, c-PARP, and c-caspase3 expression in the DKO and TKO cells. **(c)**
676 Mitochondria labeled with the MitoTracker probe (red; lower panel). Nuclei labeled with
677 Hoechst (blue; upper panel). DKO and TKO shown in the left and right panels, respectively. **(d)**
678 Cells immunostained with Opa1 antibody (green) and Hoechst (blue) in the mitochondria as

679 shown in the left (DKO) and right (TKO) panels. (e) Effects of the proteasome inhibitor MG-
680 132 on FASN, pDrp1, and Drp1 proteins observed in TKO and DKO cells. (f) DKO and TKO
681 cells transfected with ubiquitin expression plasmid followed by MG-132 (10 μ M) treatment for
682 6 hours. Subsequently, each cell lysate immunoprecipitated with Drp1 and subjected to
683 immunoblot with anti-ubiquitin antibody (P4D1). (g) Effect of DCFH-DH (10 μ M, 45 min) on
684 reactive oxygen species production in TKO cells (Red) compared to DKO cells (Blue) in clone
685 #1 (left panel) and #2 (right panel).

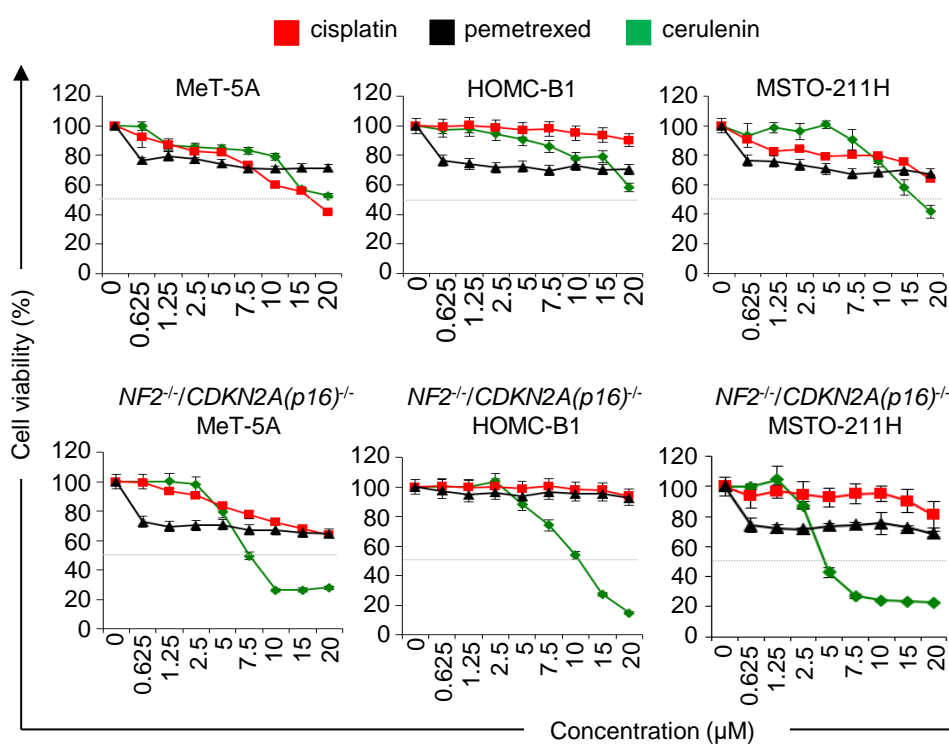
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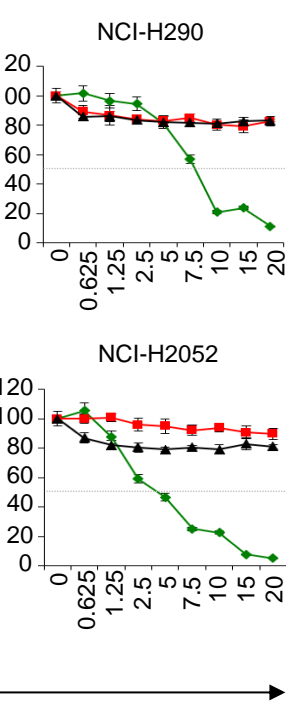
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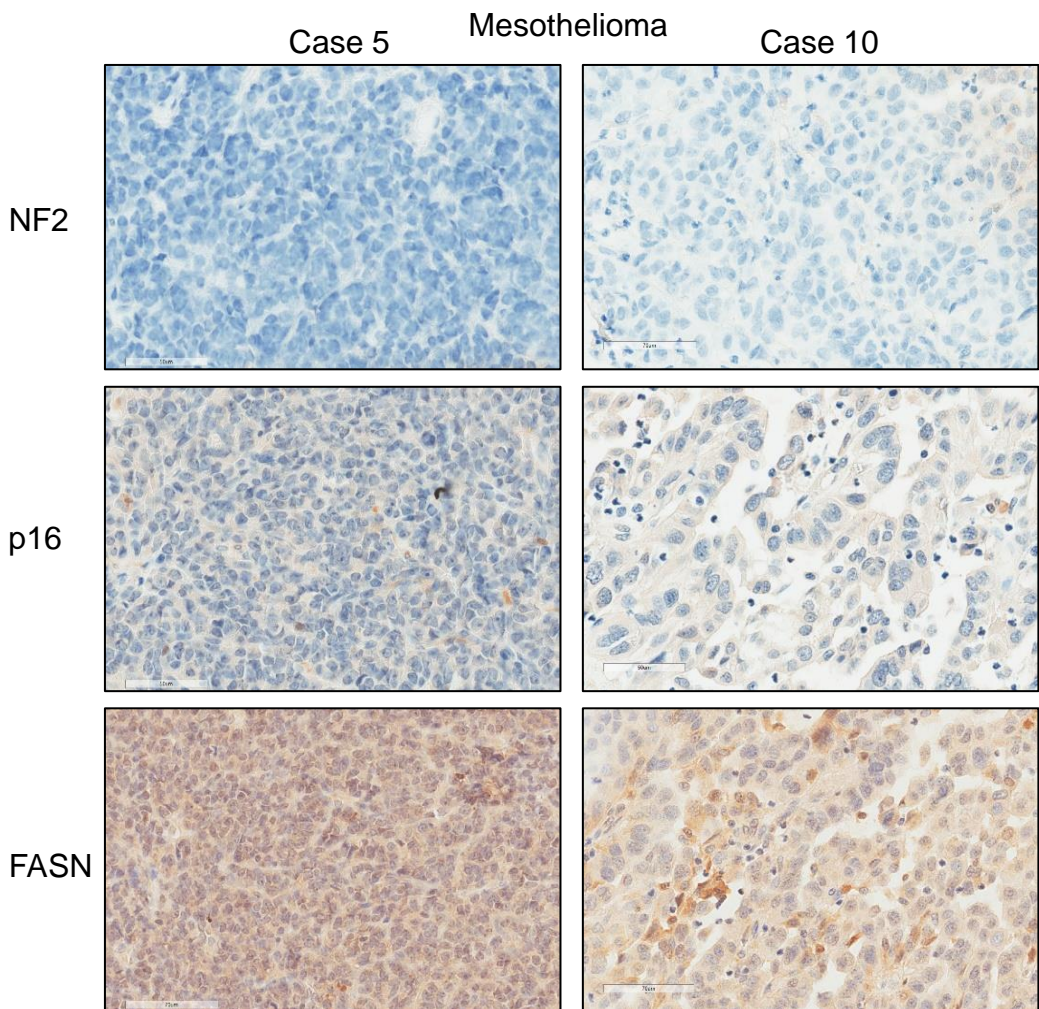
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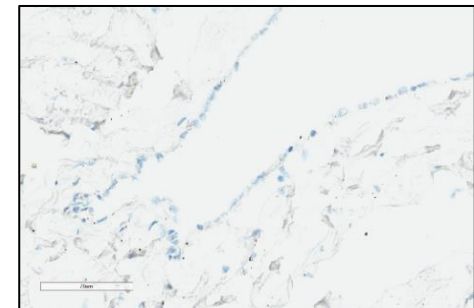


Normal Mesothelium

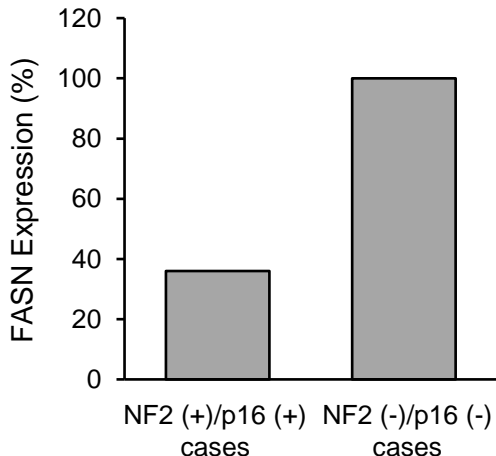
Case 46

Case 47

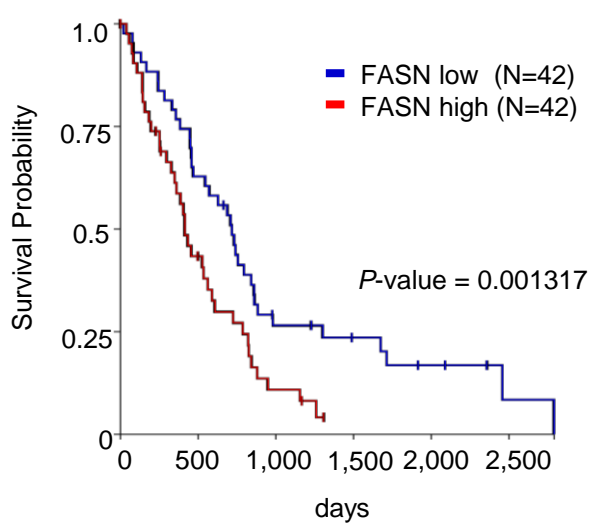
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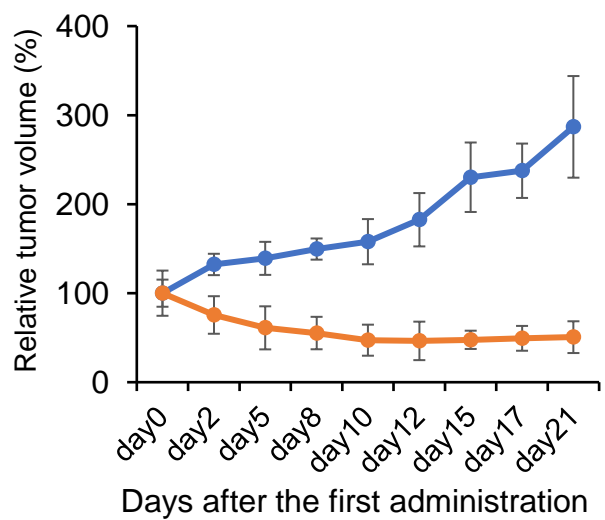
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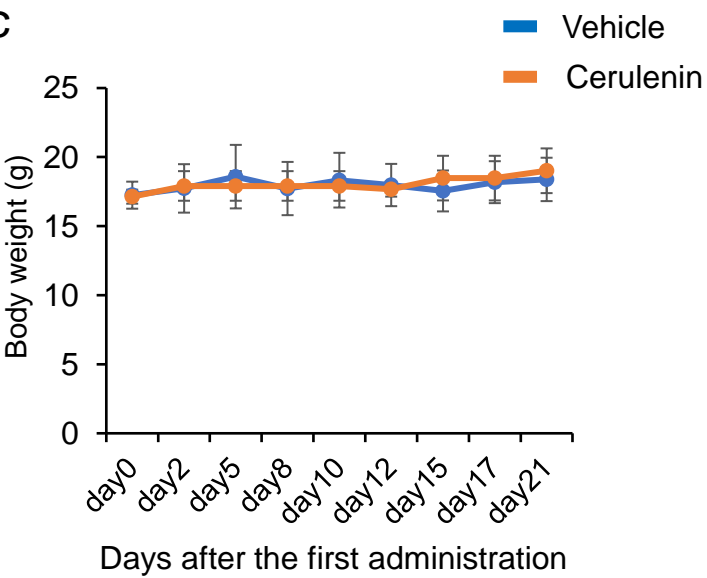
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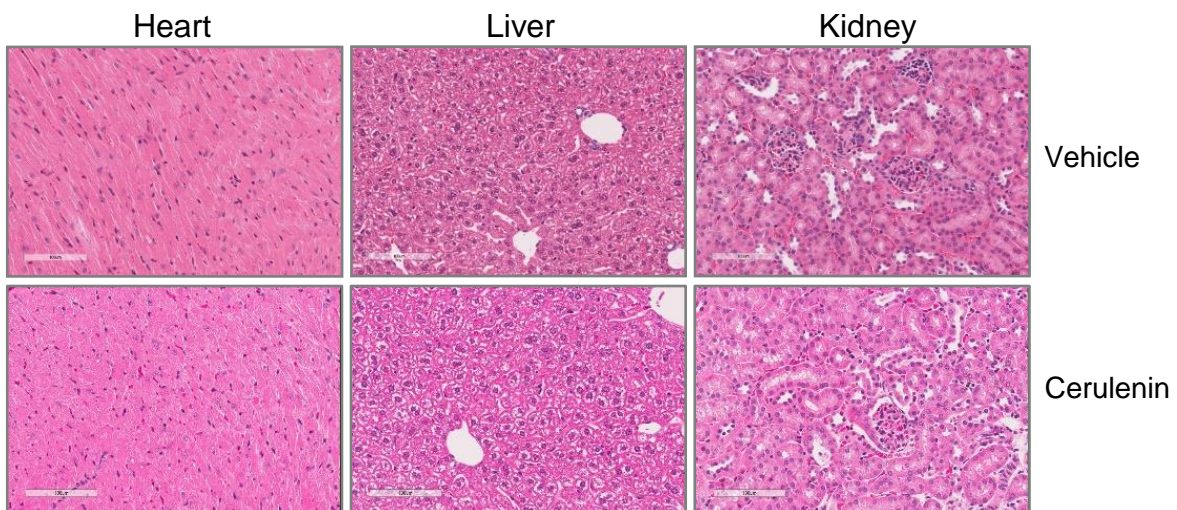
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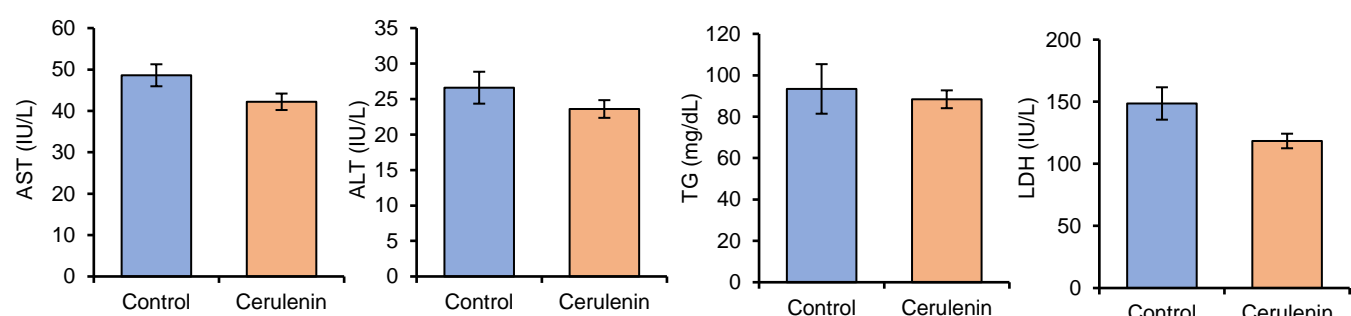
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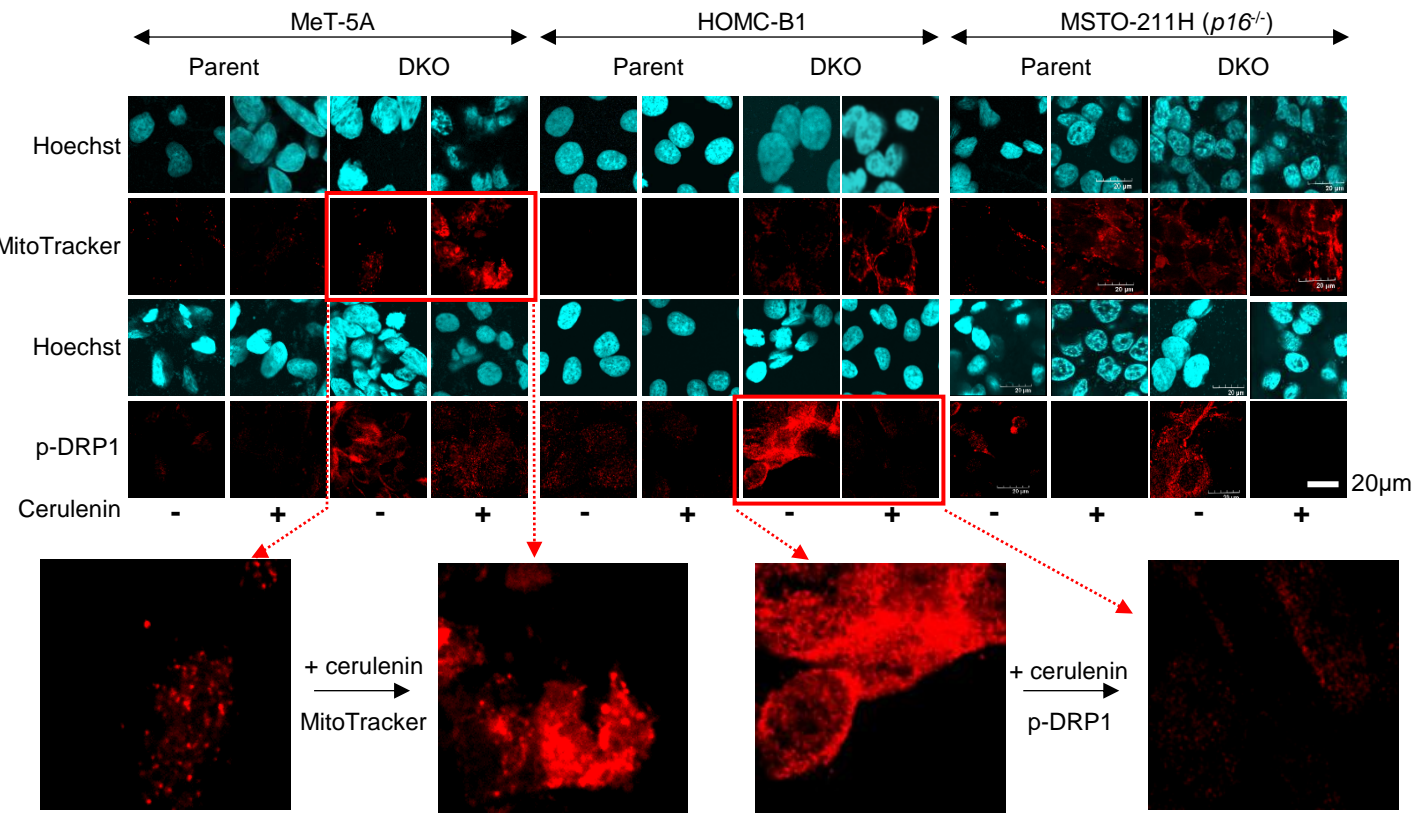
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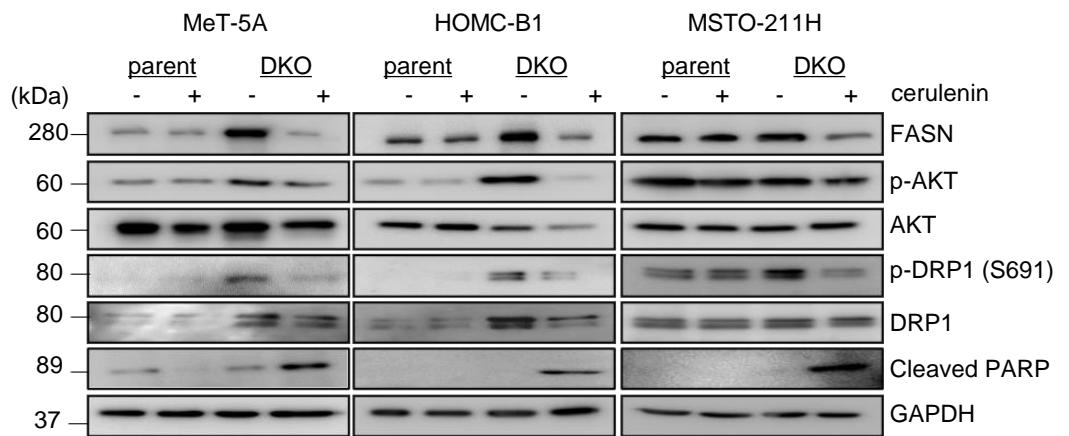
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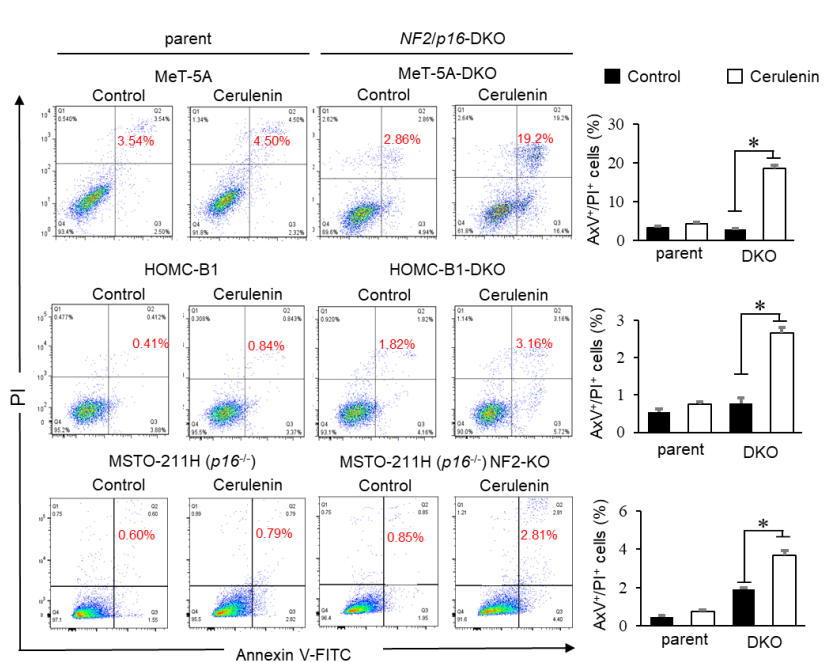
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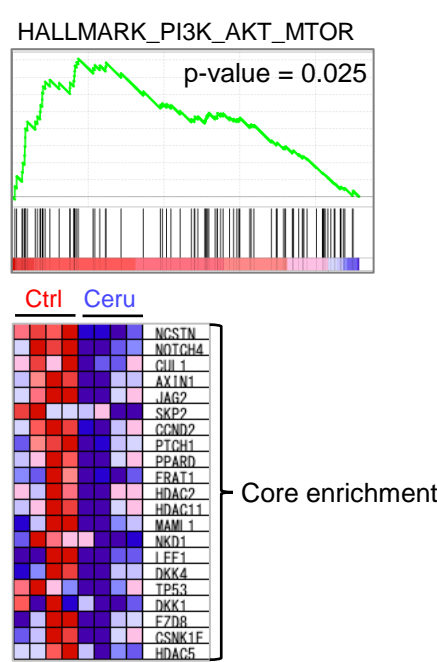
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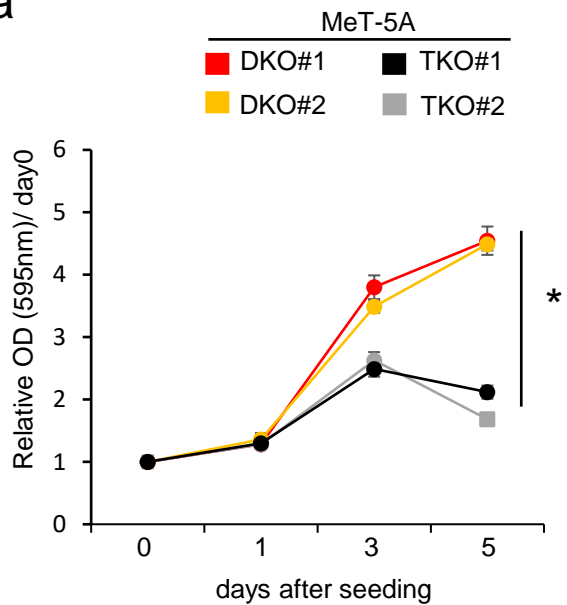
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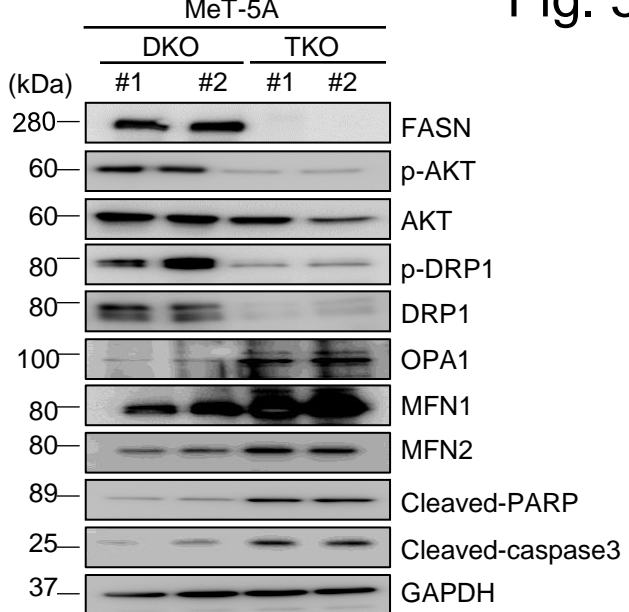
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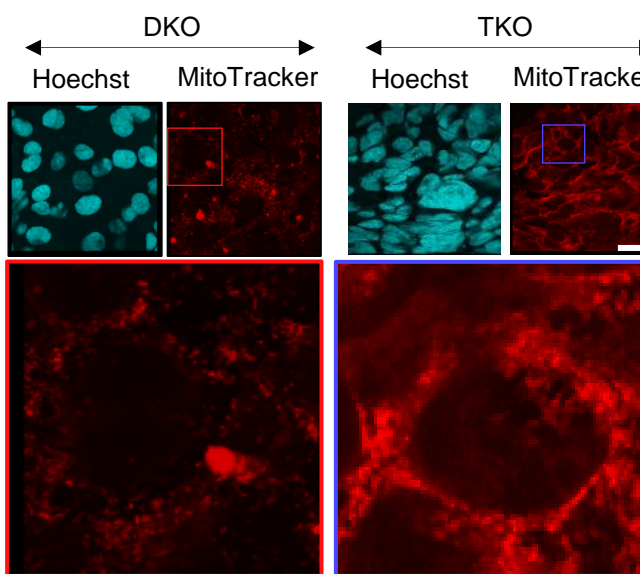
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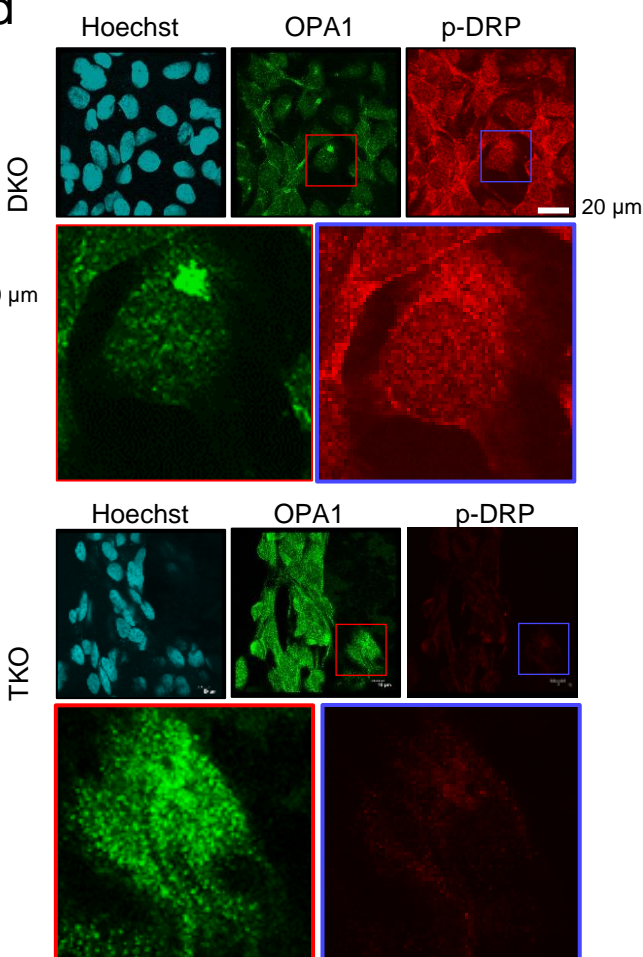
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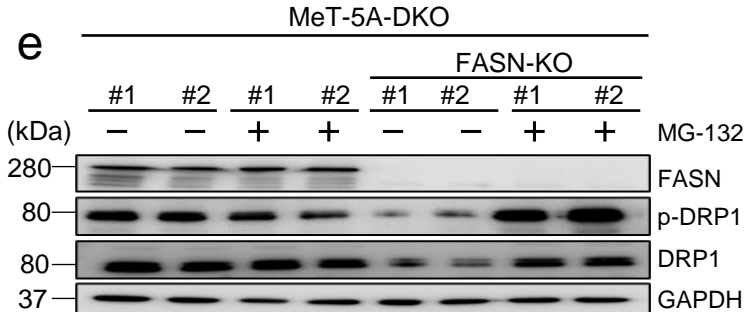
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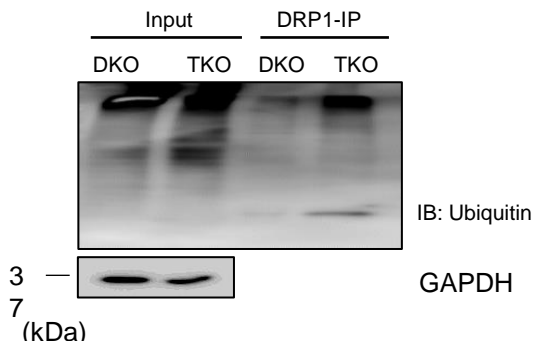
d



e



f



g

