1 Cytosolic EZH2-IMPDH2 complex regulates melanoma progression and 2 metastasis via GTP regulation

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4 Highlights

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- EZH2 has non-canonical methyltransferase-independent and GTP-dependent
 tumorigenic and metastatic functions in melanoma.
- The N-terminal EED-binding domain of EZH2 interacts with the CBS domain of
 IMPDH2 in a polycomb repressive complex 2- (PRC2-) and methylation independent manner.
- EZH2 accumulates with IMPDH2 in the cytoplasm and increases IMPDH2's
 tetramerization-mediated activity independently of EZH2 methyltransferase.
- EZH2 upregulates GTP synthesis by IMPDH2 activation and thereby activates
 ribosome biogenesis via rRNA synthesis and actomyosin contractility via RhoA
 GTPase.
- Sappanone A (SA) inhibits IMPDH2-EZH2 interactions and is anti-proliferative
 across a range of cancers including melanoma, but not in normal cells.

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19 Graphical Abstract





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23 ABSTRACT

The enhancer of zeste homolog 2 (EZH2) oncoprotein is a histone methyltransferase 24 that functions canonically as a catalytic subunit of the polycomb repressive complex 2 25 (PRC2) to tri-methylate histone H3 at Lys 27 (H3K27me3). Although targeting EZH2 26 promising therapeutic 27 methyltransferase is а strategy against cancer. methyltransferase-independent oncogenic functions of EZH2 are described. 28 29 Moreover, pharmacological EZH2 methyltransferase inhibition was only variably effective in pre-clinical and clinical studies, suggesting that targeting EZH2 30 methyltransferase alone may be insufficient. Here, we demonstrate a non-canonical 31 mechanism of EZH2's oncogenic activity characterized by interactions with inosine 32 monophosphate dehydrogenase 2 (IMPDH2) and downstream promotion of 33 guanosine-5'-triphosphate (GTP) production. EZH2-IMPDH2 interactions identified by 34 35 Liquid Chromatography-Mass Spectrometry (LC-MS) of EZH2 immunoprecipitates

from melanoma cells were verified to occur between the N-terminal EED-binding 36 domain of cytosolic EZH2 and the CBS domain of IMPDH2 in a methyltransfersase-37 independent manner. EZH2 silencing reduced cellular GTP, ribosome biogenesis, 38 RhoA-mediated actomyosin contractility and melanoma cell proliferation and invasion 39 by impeding the activity of IMPDH2. Guanosine, which replenishes GTP, reversed 40 these effects and thereby promoted invasive and clonogenic cell states even in EZH2 41 silenced cells. IMPDH2 silencing antagonized the proliferative and invasive effects of 42 EZH2, also in a guanosine-reversible manner. In human melanomas, high cytosolic 43 44 EZH2 and IMPDH2 expression were associated with nucleolar enlargement, a marker of ribosome biogenesis. EZH2-IMPDH2 complexes were also observed in a range of 45 cancers in which Sappanone A (SA), which inhibits EZH2-IMPDH2 interactions, was 46 anti-tumorigenic, although notably non-toxic in normal cells. These findings illuminate 47 a previously unrecognized, non-canonical, methyltransferase-independent, and GTP-48 dependent mechanism by which EZH2 regulates tumorigenicity in melanoma and 49 other cancers, opening new avenues for development of anti-EZH2 therapeutics. 50

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52 **Keywords:** EZH2, IMPDH2, melanoma, Sappanone A, ribosome biogenesis, 53 actomyosin contractility, GTP metabolism

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55 INTRODUCTION

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Neoplastic cells, including melanoma, are highly dependent on *de novo* biosynthesis of purine nucleotides¹. For example, the activity of Rho-GTPases in melanoma cells, and thereby formation of the actomyosin cytoskeleton which promotes cell migration and invasion, is regulated by intracellular GTP^{2,3,4}. Consistent with this, cellular GTP levels, critical for purine nucleotide synthesis, are significantly higher in melanoma cells compared to their normal cell counterparts, melanocytes².

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Increased rRNA synthesis⁵ and nucleolar hypertrophy⁶ have long been recognized
as features of malignant transformation. The requirement of GTP for Pol I transcription
and nucleolar hypertrophy has been shown recently in glioblastoma⁷, and nucleolar
hypertrophy has been associated with thicker and more mitotically active melanomas⁸.
Selective inhibition of rRNA synthesis using the RNA polymerase I inhibitor CX-5461
decreased melanoma tumorigenicity *in vitro* and *in vivo*⁹.

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Inosine monophosphate dehydrogenase 2 (IMPDH2), an oncoprotein in various 71 72 cancers¹⁰, is a key rate-limiting enzyme in nucleotide synthesis. It maintains GTP levels needed for nucleic acid synthesis, protein production via ribosome biogenesis, 73 and molecular signaling through guanine nucleotide-binding proteins (G-proteins) that 74 regulate cell functions such as cytoskeletal rearrangement, membrane trafficking, and 75 signal transduction¹¹. IMPDH2 is regulated transcriptionally, post-translationally, and 76 allosterically¹², and tetramerization is essential for its activity^{13,14}. It contains two major 77 domains: a catalytic domain for substrate interactions and the Bateman domain (CBS), 78 which is not required for catalytic activity but exerts allosteric autoregulatory 79 effects^{13,15,16}. A naturally occurring compound, Sappanone A (SA), demonstrated 80 inhibitory effects on neuroinflammation by directly targeting the conserved cysteine 81 residue 140 (Cys140) in the CBS domain of IMPDH2. Interestingly, SA selectively 82 targets and inactivates IMPDH2 but not the IMPDH1 isoform, potentially minimizing 83 lymphotoxic effects of non-specific IMPDH family targeting¹⁶. IMPDH2 is 84 overexpressed in melanoma cell lines compared to melanocytes^{17,18}, and depletion of 85 GTP via IMPDH2 inhibition with MPA induced melanocytic differentiation in melanoma 86 cells¹⁹. 87

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Enhancer of zeste homolog 2 (EZH2), a component of Polycomb Repressor 89 Complex 2 (PRC2), catalyzes tri-methylation of histone H3 at lysine 27 (H3K27me3) 90 to regulate gene expression²⁰⁻²². It has critical roles in the progression of numerous 91 malignancies²³, including melanoma²⁴⁻²⁸, where EZH2 activation represses tumor 92 suppressor genes associated with cell differentiation, cell cycle inhibition, repression 93 of metastasis, and antigen processing and presentation pathways²⁷⁻³⁰. EZH2 94 methyltransferase inhibitors have anti-cancer activity preclinically^{31,32} and in patients 95 ^{31,33}, albeit with notable toxicity. 96

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Additional to EZH2's methyltransferase activity, it also regulates gene transcription in a PRC2- and methylation-independent manner. This limits the therapeutic potential of specific EZH2 methyltransferase targeting³⁴; compounds that degrade total EZH2 protein or that target methyltransferase-independent mechanisms of EZH2 might be required to avail context-dependent therapeutic potentials of EZH2 targeting.

We recently demonstrated that EZH2 is a negative regulator of melanocytic differentiation (pigmentation), whose suppression by knockdown or degraders decreased melanoma cell clonogenicity and invasion, and induced melanocytic differentiation³⁵. In contrast, conventional EZH2 methyltransferase inhibitors displayed only minimal anti-melanoma efficacy *in vitro*³⁵. These data further suggested methyltransferase-independent, non-catalytic functions of EZH2 in melanoma tumorigenesis and invasion, prompting us to look for novel EZH2 interactions.

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Although EZH2 is mainly intranuclear, some studies have shown its cytosolic localization in fibroblasts, T lymphocytes, breast cancer, and prostate cancer cells³⁶⁻ ³⁹. Cytoplasmic functions of EZH2 are largely unknown, as most studies have focused on its nuclear functions. EZH2 is known to promote cancer progression by facilitating glucose, lipid, and amino acid metabolism^{40,41}, but other mechanisms of action are likely.

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Here, we identify a previously unrecognized methyltransferase-independent role of 119 EZH2 in melanoma tumorigenesis and invasion. We found that cytosolic EZH2 120 121 contributes to rRNA metabolism and Rho GTPase activity by regulating cytosolic IMPDH2 tetramerization-mediated activity and, in turn, promoting GTP production in 122 melanoma cells. Sappanone A (SA) inhibited interactions between EZH2 and the 123 IMPDH2 CBS domain and was anti-clonogenic in melanoma and a range of other 124 cancer types, but not in normal cells. These findings suggest novel avenues for 125 improved anti-EZH2 therapeutics. 126

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128 MATERIALS AND METHODS

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130 **Mice**

All animal experiments were performed in accordance with the Alfred Research Alliance Animal Ethics Committee protocols #E/1792/2018/M. All mice used in this study were supplied by and housed in AMREP Animal Services. Eight-week-old female NOD SCID. IL2R-/- Mice (NSG) mice were used for subcutaneous injection of pLV empty vector, shEZH2-3'UTR, shEZH2+EZH2-WT or shEZH2+ EZH2-H689A containing A375 melanoma cells (1x10⁴ cells mixed with 50 ul GFR-Matrigel, n=8 mice per group). Tumours were measured with callipers weekly, and all mice weresacrificed once the first tumour reached 20mm in diameter.

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140 Human Patient Samples

All human tissue related experiments, including human melanoma tissue FFPE samples, isolated primary human melanocytes and bone marrow samples were performed in accordance with the Alfred Human Research Ethics Committee protocols #155/18 and #29/05.

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62 human melanoma tumor tissue sections ranging from grade I to IV were obtained
from Melanoma Research Victoria (MRV) under the guidelines approved by the
Victorian Government through the Victorian Cancer Agency Translational Research
Program. MRV obtained the informed consent from all participants.

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151 Cell lines and primary cells

The HEK293, C32, SK-MEL28, IGR39, A375, B16-F10 and IGR37 cell lines were 152 obtained from ATCC and cultured under conditions specified by the manufacturer. 153 154 C006-M1 cell line was from QIMR Berghofer Medical Research Institute. LM-MEL28, LM-MEL33, LM-MEL43, LM-MEL45 were from Ludwig Institute for Cancer 155 Research⁴². LM-MEL28: B4:F3 is the monoclonal line derived from LM-MEL28 cells in 156 our lab previously. MCF7 and MDA-MB-231 cell lines were provided by Prof Jane 157 Visvader (WEHI), OVCAR3, OVCAR8 cell lines were kindly provided by Prof David 158 Bowtell (Peter MacCallum Cancer Centre), PC3, LNCaP, C4-2 cell lines by A Prof 159 Renea A. Taylor (Monash Biomedicine Discovery Institute), OMM1 was kindly 160 provided by Prof Bruce R. Ksander (Harvard Medical School) and 92.1 cell line by Prof 161 Martine Jager (Leiden University Medical Centre). Mycoplasma tests were routinely 162 performed in our laboratory and short tandem repeat (STR) profiling was conducted 163 by the Australian Genome Research Facility (AGRF) to authenticate the human cell 164 lines. 165

166

167 Chemicals

168 The chemicals used for treating cells were GSK126 (Selleckchem, S7061), EPZ6438 169 (Selleckchem, S7128), Sappanone A (Cayman Chemicals, 23205), MPA 170 (Selleckchem, S2487), Ribavirin (Selleckchem, S2504), DZNep (Sigma, S804983)

and MS1943 (MedChemExpress, HY-133129); all are listed in Table S1.

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173 Plasmids, Cloning, Overexpression, and siRNA

pCMVHA hEZH2 (#24230) and pLV-EF1a-V5-LIC (#120247) plasmids were 174 purchased from Addgene and MYC/FLAG-hIMPDH2 (#RC202977) plasmid from 175 Origene. EZH2 (1-170), EZH2 (1-340), EZH2 (1-503), EZH2 (1-605), EZH2 (171-751) 176 deletion mutants, full length EZH2 (1-751) and IMPDH2 (1-187) deletion mutant was 177 cloned into pLV-EF1a-V5-LIC vector backbone's Srfl/Notl RE using the cloning 178 primers listed in Table S1. pCMVHA hEZH2 and V5-EZH2 vector was used to 179 generate EZH2-H689A mutant vector using the mutagenesis primers listed in Table 180 S1 with QuikChange II site-directed mutagenesis kit (Agilent) following the 181 manufacturer's instructions. Custom designed siRNA oligonucleotides listed in Table 182 S1 were purchased from Bioneer Pacific. For transient transfection, 25x10⁴ cells were 183 transfected with 2.5 µg of DNA using Lipofectamine 3000 transfection reagent 184 (Invitrogen). For siRNA experiments, 25x10⁴ cells were transfected with 10 nM of the 185 indicated oligonucleotides in Table S1 using the Lipofectamine RNAiMAX transfection 186 187 reagent (Invitrogen). 72 hours after siRNA transfection, cells were used for functional assays or collected for western blot analysis. 188

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Virus-containing supernatant was collected 48 hours after co-transfection of pCMV-190 191 VSV-G, psPAX2, pMD2.G and the EZH2 vectors into HEK293 cells, and then added to the target cells. Stable knockdown and rescue of EZH2 was achieved by lentiviral 192 transduction of EZH2 with V5-EZH2-WT or V5-EZH2-H689A. After transduction, cells 193 were selected for antibiotic resistance with 2 µg/mL puromycin (Sigma Aldrich, 194 #P8833), followed by knockdown using stable short hairpin interfering RNA (MISSION 195 shRNA, Sigma Aldrich) targeting the 3'UTR of human EZH2 (TRCN0000286227), as 196 previously reported³⁸. 197

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199 GST pull-down Assay

GST pull-down assay was performed as previously described⁴³ with minor modifications. The plasmid GST-EZH2 (1-170), -EZH2 (1-340), -EZH2 (1-503), or – EZH2 (1-605) or GST only was expressed in BL-21 bacteria in the presence of 0.5mM IPTG for 2.5 h at 37°C. Bacterially expressed GST only (control) or each GST–EZH2

mutant peptide were solubilized in NETN buffer (1% NP-40, 20mM Tris-HCl, pH 8.0, 204 100mM NaCl, 1mM EDTA) and sonicated in 30 second bursts followed by 30 seconds 205 rest for 15 cycles. Then they were purified by affinity chromatography on Glutathione 206 Magnetic Agarose Beads (Pierce, Thermo Fischer)) and stored in PBS at 4°C until 207 use. For GST-pull-down assays, purified GST control or GST-EZH2 mutant peptides 208 were mixed with total lysates isolated from HEK293 cells, overexpressing V5-IMPDH2-209 CBS, grown in serum-fed condition and then incubated for 2 h at 4°C with constant 210 rotation. The lysates from HEK293 cells were used as a source of IMPDH2-CBS 211 212 domain. After extensive washing of unbound proteins, bound protein was eluted and analyzed by sodium dodecyl-PAGE (SDS-PAGE). 213

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215 Colmmunoprecipitation and HA/ FLAG pulldown assays

Pellets of 1x10⁷ cells were lysed with 1mL Co-IP Lysis Buffer (300mM NaCl, 50mM 216 Tris HCL pH7.4, 0.5% NP40, 0.1% Sodium deoxycholate, 2% SDS) with PhosSTOP 217 (Roche) and cOmplete (Roche) rolled at 4°C for one hour. DynaBeads™ Protein G 218 (Thermofisher) were washed three times with Co-IP lysis buffer and chilled in 219 preparation. Lysates were centrifuged at 15,000 RPM for 15 minutes at 4°C and the 220 221 supernatant was collected, pre-cleared with 20µL of prewashed DynaBeads and incubated on a roller for 1 hour at 4°C. Lysates had pre-cleared beads removed and 222 were split with 500µL for IgG control and 500µL for EZH2 sample, topped to 1 mL with 223 Co-IP lysis Buffer. These were incubated overnight at 4°C with 1:1250 of Rabbit (DAIE) 224 225 mAB IgG Isotype control or 1:300 of anti-EZH2 (D2C9) XP Rabbit antibody, respectively. After 16 hours of incubation, 35µL of pre-washed DynaBeads were 226 added to IgG control or EZH2 sample and returned to the roller for 2-4 hours incubation 227 at 4°C. Beads were washed with Co-IP buffer once, and then buffers of increasing salt 228 concentrations (Buffer 1, 50mM Tris, pH8.0, 150mM NaCl; Buffer 2, 50mM Tris, pH8.0, 229 450mM NaCl; and buffer 3, 1M Tris, pH8.0). For Mass Spectrometry, proteins were 230 eluted by resuspending in 150µL of 0.2M Glycine, pH2.5, for 5 minutes on ice and 231 collecting supernatant, which was repeated twice. To the 450 µL of sample, 100 µL of 232 1M Tris-HCI (pH8.0) was added and the samples were kept at -80°C until LC-MS 233 analysis. For CoIP-WB analysis beads were washed three times with Co-IP Lysis 234 Buffer. 235

For HA pulldown assays the cells were lysed with 500 μ L of IP Lysis Buffer containing cOmplete (Roche) protease inhibitor cocktail and incubated at 4°C for 35 min on a rotator followed by centrifugation at 15000 rpm for 15 min at 4°C. 50 μ L of the lysates were kept for inputs. 25 μ L of Pierce anti-HA magnetic beads (Thermo Fischer Scientific) were added onto the lysate and incubated at RT for 30 minutes on a rotator. Beads were washed with 300 μ L of TBST three times and the beads were boiled in 2x SDS-Laemmli Sample Buffer for 10 minutes.

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245 Western blot

Total proteins were extracted from cell lines and tumor xenografts in ice-cold lysis 246 buffer (10mM Tris HCL pH8.0 1mM EDTA 1% TritonX100 0.1% Sodium Deoxycholate 247 2% SDS 140mM NaCl, protease inhibitors and phosphatase inhibitors). Lysates were 248 prepared after incubation on ice for 1h and centrifugation for 15 minutes cold at 15,000 249 rpm. Supernatants were boiled in 6x SDS-Laemmli Sample Buffer for 10 minutes. 250 Proteins were run on 4-20% Mini-PROTEAN TGX Stain-Free Protein Gels (BioRad, 251 4568096) and then transferred to PVDF membrane by wet transfer system. 252 Membranes were blocked with PBS containing 0.1% Tween-20 and 5% (w/v) skim 253 254 milk, followed by incubation with the antibodies listed in Table S1. Signals were detected using Clarity ECL Western blotting Substrate (BioRad). Where applicable, 255 signal intensities were quantified by ImageJ densitometry analysis software. 256

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258 Non-reducing SDS-PAGE

Samples were lysed in 2x Laemmli Sample Buffer without SDS and DTT and run withSDS free running buffer.

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262 DSS Crosslinking

 1×10^{6} cells were precipitated and washed once with 1xPBS. The cell pellet was resuspended in 500 µL of 1xPBS. Cell suspensions were treated with either DMSO (control) or 1mM DSS (A39267, Thermo Fisher) and incubated for 30min at RT. Then, the cells were quenched with 50mM Tris-HCl pH:8.0 by incubating for 15min at RT. Finally, the cells were centrifuged, and the cell pellets were boiled in 50 µL of 2x SDS-Laemmli Sample Buffer for 10 min at 95°C.

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270 Cytoplasmic and nuclear fractionation

Cytoplasmic and nuclear extracts were isolated using a nuclear extraction kit according to the manufacturer's protocol (Affymetrix; Santa Clara, CA) with modifications⁴³. Co-IP was performed with anti-EZH2 or anti-IMPDH2 antibody at 4°C as described in the Co-IP method section. The immune complexes were collected with Protein G-Dynabeads (Thermo Fischer) and washed in lysis buffer. Bound proteins were analyzed by SDS–PAGE and WBs.

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278 RNA isolation and quantitative PCR

279 Total RNA from cells was extracted using Purelink RNA mini isolation kit according to the manufacturer's instructions (Thermo Fischer Scientific) with the additional Purelink 280 On-Column DNA purification (ThermoFisher Scientific) step. Complementary DNA 281 (cDNA) was synthesized using total RNA (1 µg per reaction) with SuperScript Vilo 282 cDNA synthesis kit (Thermo Fisher Scientific) as per manufacturer's protocol. 283 Quantitative PCR (qPCR) was carried out using Fast SYBR Green Master Mix 284 (Invitrogen) and LightCycler 480 Instrument II (Roche). RNA expression changes were 285 determined using a $\Delta\Delta$ Ct method⁴⁴. RPLP0 mRNA was used as an internal control in 286 all qPCR reactions. Table S1 shows the qPCR primers used for IMPDH2, pre-rRNA, 287 288 pre-tRNA, pre-GAPDH, p53, CDKN1A, CDKN2A, MDM2, PUMA and RPLP0 mRNA amplifications. 289

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291 Cell Proliferation, Clonogenicity and Sphere Formation Assay

To measure cell proliferation rates, we plated equal numbers of cells in 6-well plates. Cells were trypsinized and counted on the indicated days by haemocytometer after trypan blue staining.

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For clonogenicity assay, cells were fixed with ice-cold absolute methanol for 20 min and air-dried for 15 minutes. Cells were stained with 0.5% Crystal Violet for 20 min at room temperature and then rinsed with tap water to remove excess dye. Five random fields of stained cells were imaged using bright field microscopy at 40 × magnification and average cell numbers per field were plotted as a function of time.

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Sphere formation assays were performed as described⁴³. Briefly, cells (500 cells per well) suspended in 100 mL ice-cold Matrigel in RPMI medium (1:1 ratio) were overlaid onto the pre-solidified 50% Matrigel in 24-well plates (100 ml per well). Cells were fed with 500 mL RPMI medium containing 10% FBS and grown for 14 days with a change
 of medium every 3 days. For the SA study, A375 and B16-F10 cells (500 cells/well)
 were grown on Matrigel and treated either with DMSO (vehicle) or SA in serum-fed
 conditions for the indicated days. Spheres were imaged and then manually quantified.

310 Cell senescence β-Gal assay

Cells were fixed and stained with Senescence β-Galactosidase Staining Kit (CST
#9860) following the manufacturer's instructions and imaged on a Leica DMIL LED
microscope under bright field settings.

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315 **3D Matrigel Invasion and Wound Healing Assays**

Cells (1x10⁵ cells per well) were seeded in a 24-well Boyden chamber with an 8-mm 316 filter coated with 20% growth factor reduced Matrigel. Cells were grown in RPMI 317 medium containing 10% FBS for 16, 24 and 48 h at 37° 318 С with 5% CO2. Cells the inner side of the chamber were gently removed by scraping with a wet cotton 319 on swab. Invaded cells at the outer side of the chamber were fixed with 4% formaldehyde 320 for 30 min at RT and rinsed twice with PBS. Cells were stained with 0.5% Crystal 321 322 Violet for 20 min at RT and then rinsed with tap water to remove excess dye. Analysis was performed based on the average number of stained cells per field from f 323 ive random fields at 20x magnification on a Leica DMIL LED microscope 324

Wound healing assays were performed by seeding cells in complete media on a 24-325 well plate for 24–48 h until a confluent monolayer had formed. Linear scratches were 326 µl pipette tip. Monolayers were washed three times with made using a sterile 200 327 PBS to remove detached cells, and then complete media was added. Images 328 of the wound were taken immediately and 24h following wound formation 329 on a Leica DMIL LED microscope under the phase contrast setting . Wound area was 330 measured over time using ImageJ. 331

332

333 RhoA activity assay

RhoA activities were measured in melanoma cells using RhoA G-LISA activation assay kit according to the manufacturer's protocol. Briefly, cells were lysed in ice-cold lysis buffer and quickly cleared by centrifugation. Precision Red Advanced Protein Assay Reagent (Part # GL50) was used to quantify protein contents. Equal amounts of proteins were loaded onto ELISA plates. After several antibody incubation and washing steps the active RhoA bound protein levels were evaluated colorimetricallyby OD490 nm absorbance measurement.

341

342 IMPDH2 activity assay

IMPDH2 activity was measured by monitoring the reduction of NAD⁺ to NADH and the subsequent increase in absorbance at 340 nm in buffer: 100 mM Tris-HCl, 100 mM KCl, 2 mM DTT pH 7.4. 2 µg of recombinant IMPDH2 protein was preincubated with 3 mM IMP and 2 µg of recombinant EZH2 and then with 10 mM GTP for an additional 10 min before the reaction was initiated by the addition of 1 mM NAD⁺. NADH production was measured 1 h after incubation in a FLUOstar Omega plate reader by 0D340 nm absorbance.

350

351 Histochemical and immunostaining

H&E staining was done to evaluate nucleolar sizes⁵. For Fontana Masson staining, 352 cells were sorted onto slides were fixed with 4% PFA for 20 minutes and washed twice 353 with distilled water for 5 minutes. Then slides were incubated in Fontana silver nitrate 354 working solution (2.5% Silver nitrate, 1% ammonium hydroxide) at 60°C for 2 hours. 355 356 Slides were rinsed in water three times and incubated with 0.2% gold chloride solution (Sigma-Aldrich) for 2 minutes. Rinsed slides were incubated with 5% sodium 357 thiosulfate for 2 minutes. After rinsing with water twice, slides were counterstained with 358 10 µg/mL DAPI solution for 5 minutes. Slides were rinsed and mounted with 359 fluorescence mounting medium (Dako). For Schmorl's staining, samples were 360 dewaxed with 3 x 5-minute histolene washes, and rehydrated in washes of 100%, then 361 95% and then 75% ethanol for 5 minutes each. Slides were washed with distilled water 362 and placed in Schmorl's Stain for 10 minutes. Slides were washed with water for 1 363 minute, placed in Eosin in water for 15 seconds, and returned to constant washing 364 with water for 3 minutes. The slides were finished with 4 x 2-minute washes of 100% 365 ethanol and 3 x 2-minute washes of Histolene. Slides were mounted with DPX 366 367 Mounting Medium (Thermo Fisher Scientific).

368

For melanoma patient sample IHC, slides were incubated at 60°C for 1h, dewaxed in histolene, and hydrated through graded alcohols and distilled water. Sections were subjected to antigen retrieval in Antigen Retrieval solution (Dako, pH6 for EZH2, IMPDH2 antibody) at 125° for 3 minutes heated by a pressure cooker. Primary antibody listed in Table S1 was diluted into blocking buffer and slides were incubated
overnight at 4°C. After washing with TBST, the slides were incubated with secondary
antibody using an ImmPRESS™ HRP Anti-Mouse IgG (Peroxidase) Polymer
Detection Kit (Vector Laboratories) for 60 min at RT. Sections were washed with TBST
and slides were developed by adding AEC+ High Sensitivity Substrate Chromogen
Ready to use (Dako K346111-2).

379

For immunofluorescence, cells were fixed with 4% PFA diluted in PBS for 15 min at 380 381 RT, rinsed three times with PBS, and blocked for 1h using blocking buffer (5% normal donkey serum containing 0.3% Triton X-100 in PBS). After blocking, slides were 382 incubated with primary antibody (Table S1) diluted in antibody buffer (5% bovine 383 serum albumin containing 0.3% Triton X100 in PBS) at 4 °C overnight. S 384 lides were washed three times with PBS and incubated with fluorescent secondary antibodies 385 indicated in Table S1. Slides were washed three times with PBS, stained with 10µg/ml 386 DAPI and coverslipped using Fluorescence Mounting Medium (Dako). Slides were 387 imaged using Leica DMIL LED inverted fluorescent microscope or Nikon A1r Plus si 388 confocal microscope. 389

390

391 **Proximity ligation assays**

Cells were seeded on round coverslips. After 24 h of seeding, cells were fixed with 4% 392 PFA for 15 min at RT, rinsed three times with PBS, and blocked for 1 h using blocking 393 394 buffer (5% normal goat serum containing 0.3% Triton X-100 in PBS). After blocking, slides were incubated with primary antibody diluted in antibody buffer (5% bovine 395 serum albumin containing 0.3% Triton X-100 in PBS) at 4°C overnight. Slides were 396 then washed three times with PBS and incubated with DuoLink PLA probes (Sigma, 397 Cat #DUO92101). The protocol for PLA secondary antibody incubation, ligation, 398 amplification, and washes were performed following the manufacturer's protocol. 399 Slides were imaged using a Nikon A1r Plus confocal microscope. Positive signals were 400 normalized to single-primary antibody control (EZH2 or IMPDH2) and image analysis 401 was performed using ImageJ. 402

403

404 **PDX Tumor Dissociation**

Mice were euthanized with CO2 and tumors were resected. Tumors were manually dissociated in Hank's Balanced Salt Solution (without Ca2+ and Mg2+, HBSS-/-),

followed by enzymatic tumour dissociation using the gentleMACS tissue dissociator in 407 Tissue Dissociating media (200 u/mL Collagenase IV, 5 mM CaCl₂ in HBSS -/-). Tissue 408 was washed with HBSS-/- and pelleted at 220g for 4 minutes at 4°C, and the 409 supernatant was removed. After this, the pellet was resuspended with 100units/g of 410 DNase and 5mL/g of warmed trypsinEDTA and incubated at 37°C for 2 minutes. Equal 411 volumes of cold staining media were added, and the samples were pelleted at 220g 412 for 4 minutes at 4°C. Supernatant was removed and the pellets were resuspended in 413 cold staining media and filtered with a 40-micron cell strainer. To separate the tumoral 414 415 cells from mouse stroma, cells were stained with an antibody cocktail of directly conjugated antibodies to mouseCD31 (endothelial cells), mouse CD45 (white blood 416 cells), mouse TER119 (red blood cells) and human HLA-A/B antibodyin staining media 417 on ice for 30 minutes. Labelled cells were resuspended in 2µg/ml DAPI in staining 418 media with 10% FBS and 10uL/mL of DNase. Cells were subsequently analyzed 419 and/or sorted on a FACSFusion (Becton Dickinson). 420

421

422 CD34+ bone marrow progenitor cell isolation and culturing

Donor CD34+ HSPC samples were obtained from normal patients after informed 423 424 consent in accordance with guidelines approved by The Alfred Health human research ethics committee. Cells from a leukapheresis sample were isolated using Ficoll-Paque 425 PLUS (GE Healthcare) and density centrifugation, followed by NH4CI lysis to remove 426 red blood cells. A secondary isolation step was completed using CD34 MicroBead Kit 427 (Miltenyi Biotec) performed according to the manufacturer's protocol for positive 428 selection of CD34+ cells from the mononuclear population. Isolated CD34+ cells were 429 cultured in expansion medium (Stemspan SFEM (Stem Cell Technologies 09650). 430 50ng/mL rhFLt3L (R&D 308-FKN), 50ng/mL rhSCF (R&D 255-SC), 10ng/mL rhIL-3 431 (R&D 203-IL), 10ng/mL rhIL-6 (R&D 206-IL), 35nM UM171 and 500nM Stemreginin) 432 with or without SA for 4 and 7 days. 433

434

Human skin acquisition, single cell suspension, isolation of melanocytes via FACS and melanocyte culture

Epidermal melanocytes were isolated from normal adult human breast skin. The skin samples were provided from Caucasoid donors (age 18 - 72) via The Victorian Cancer Biobank. Fat was removed from the skin and washed in PBS with Gentamycin (10 μ g/mL) and 80% EtOH. Then the skin was cut into small pieces (~5 mm²) and

incubated in Dispase (15 U/mL, Gibco/Thermo Fisher Scientific) with Gentamycin (10 441 µg/mL) at 4°C overnight. Epidermis was peeled from dermis by forceps and smashed 442 by scissors and incubated in Trypsin/EDTA (0.25%) at 37°C for 10 min to make a 443 single cell suspension of the epidermal cells. After pipetting and addition of fetal bovine 444 serum (FBS) to stop activity of trypsin (final concentration of FBS is 10%), the 445 epidermal single cell suspension was passed through cell-strainers (70 µm then 40 446 µm). After centrifugation (220g for 5 min) the collected epidermal cells were 447 suspended in the staining medium and viability was validated microscopically with 448 449 trypan blue.

450

The collected epidermal cells from skin were incubated with primary antibodies 451 including FITC anti-human CD326 (EpCAM) (1:100, mouse), FITC anti-human CD31 452 (1:100), FITC anti-human CD45 (1:100), FITC anti-human CD235a (1:100) and PE 453 anti-human CD117 (c-kit) (1:100) in the staining media for 30 min at 4°C. After a wash 454 and centrifuge the cells were suspended in DAPI (2.5 µg/mL) and subjected to FACS 455 analysis (BD FACSAria[™] Fusion flow cytometer, BD). Debris (by morphology plot: 456 FSC-A/SSC-A), doublets (by doublet plot: FSC-H/FSC-W and SSC-H/SSC-W) and 457 458 dead cells (DAPI+) were excluded. The Ckit+CD326-CD31-CD45-CD235a- fraction was sorted into 1.5 mL microcentrifuge tubes filled with Medium 254 (1 mL). 459

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The sorted primary human melanocytes were plated on HaCaT-derived ECM-coated culture dish and cultured in Medium 254 supplemented with Human Melanocyte Growth Supplement-2 (HMGS-2, including basic FGF, insulin, transferrin, bovine pituitary extract, endothelin-1, FBS, heparin and hydrocortisone, concentrations are proprietary, PMA-free) at 37°C with 10% O2 and 5% CO₂.

466

467 **RNA-Seq data analysis**

FASTQ files were processed using Laxy (https://zenodo.org/record/3767372) which 468 RNAsik pipeline 469 encompasses the (https://joss.theoj.org/papers/10.21105/joss.00583). Briefly, GRCh38 reference 470 genome was used for STAR alignment⁴⁵ and gene expression counts were performed 471 featureCounts⁴⁶. using Gene counts analysed using Degust 472 were (https://zenodo.org/record/3501067) for differential expression 473 analysis. Data 474 processing was performed on NeCTAR Cloud Servers, or MASSIVE High475 Performance Computing (HPC) cluster.

476

Differential gene expression analysis was performed using edgeR (v.3.32.1). Quasilikelihood F-test was performed with glmQLFit and glmQLFTest functions. Gene ontology (GO) enrichment test was performed using PANTHER (v16.0) fisher's exact test corrected by false discovery rate (FDR).

481

482 **TCGA survival analysis**

The clinical data and mRNA expression profiles for skin cutaneous melanoma samples 483 in TCGA PanCancer Atlas database were retrieved from MSKCC Cancer Genomics 484 Data Server (CGDS) (http://www.cbioportal.org)⁴⁷. The "high expression" and "low 485 expression" groups for each gene were defined as above or below the median 486 expression level for the cohort respectively. The overall survival (OS) curves were 487 calculated with the Kaplan-Meier method and the statistical significance were tested 488 with the log-rank test. The calculations were performed using the R package 'survival' 489 3.1-11 and the survival curves were plotted using the R package 'survminer' 0.4.4. 490

491

492 Sample preparation for GTP analysis

1 x10⁷ A375 cells were washed once with 0.9% NaCl and cell pellets were snap frozen 493 prior to LC-MS analysis. 200 µL of extraction solvent (2:6:1 CHCl₃: MeOH: H₂O) at 494 495 4°C was added to the washed cell pellets after which the samples were briefly vortexed, sonicated in an ice-water bath (10 minutes). Samples were then frozen in 496 liquid nitrogen and thawed three times before mixing on a vibrating mixer at 4°C for 10 497 minutes after which they were subjected to centrifugation (20,000 x g, 4°C, 10 min) 498 and the supernatant transferred to samples vials for prompt (same day) LC-MS 499 500 analysis.

501

502 LC-MS analysis for metabolomics

503 Samples were analyzed by hydrophilic interaction liquid chromatography coupled to 504 triple quadrupole mass spectrometry (LC–MS). In brief, the chromatography utilized a 505 ZIC-p(HILIC) column (Merck SeQuant ZIC-pHILIC 5um 150 x 4.6 mm, polymeric) and 506 guard (Merck SeQuant ZIC-pHILIC Guard, 20 x 2.1 mm, PEEK coated guard) with a 507 gradient elution of 20 mM ammonium carbonate (A) and acetonitrile (B) (linear gradient time-%B as follows: 0 min-80%, 15 min-50%, 18 min-5%, 21 min-5%, 24 min-80%, 32 min-80%) on a 1290 Infinity II (Agilent). The flow rate was maintained at 300 μ L/min and the column temperature 25°C. Samples were kept at 10°C in the autosampler and 5 μ L injected for analysis. The mass spectrometry was performed in multiple reaction monitoring (MRM) mode on an Agilent 6495 Triple Quadrupole. Full details are provided in supplementary material. Peak integration was carried out using MassHunter Qualitative Navigator B.08.00 (Agilent).

515

516 LC-MS analysis for proteomics

Immunoprecipitated proteins were reduced with 10 mM TCEP (Thermo Fisher),
alkylated with 40 mM iodoacetamide (Sigma Aldrich) and digested with sequencing
grade trypsin (Promega). Samples were acidified with 1% formic acid (FA) and purified
using OMIX C18 Mini-Bed tips (Agilent) prior to LC-MS/MS analysis.

521

Using a Dionex UltiMate 3000 RSLCnano system equipped with a Dionex UltiMate 522 3000 RS autosampler, an Acclaim PepMap RSLC analytical column (75 µm x 50 cm, 523 nanoViper, C18, 2 µm, 100Å; Thermo Scientific) and an Acclaim PepMap 100 trap 524 column (100 µm x 2 cm, nanoViper, C18, 5 µm, 100Å; Thermo Scientific), the tryptic 525 peptides were separated by increasing concentrations of 80% acetonitrile (ACN) / 526 0.1% formic acid at a flow of 250 nl/min for 128 min and analyzed with a QExactive 527 HF mass spectrometer (ThermoFisher Scientific). The instrument was operated in the 528 529 data dependent acquisition mode to automatically switch between full scan MS and MS/MS acquisition. Each survey full scan (m/z 375–1575) was acquired in the Orbitrap 530 with 60,000 resolution (at m/z 200) after accumulation of ions to a 3 x 10⁶ target value 531 with maximum injection time of 54 ms. Dynamic exclusion was set to 15 seconds. The 532 12 most intense multiply charged ions ($z \ge 2$) were sequentially isolated and 533 fragmented in the collision cell by higher-energy collisional dissociation (HCD) with a 534 fixed injection time of 54 ms, 30,000 resolution and automatic gain control (AGC) 535 target of 2×10^5 . 536

537

Raw data files were analyzed with the MaxQuant software suite v1.6.5.0⁴⁸ and its implemented Andromeda search engine⁴⁹ to obtain protein identifications and their respective label-free quantification (LFQ) values using standard parameters. The proteomics data were further analyzed using either Perseus⁵⁰ or LFQ-Analyst⁵¹.

542

543 Statistical Analysis

Analysis was performed using GraphPad Prism version 8. All analyses were performed using log-rank test, unpaired two-tailed t-tests, one-way or two-way ANOVA plus Tukey's multiple comparison tests as appropriate to the data type. The statistical parameters are reported in figure legends or text of the results. P values less than 0.05 were considered significant.

- 549
- 550 **RESULTS**
- 551

552 Methyltransferase-independent activity of EZH2 in melanoma

553

We recently reported that decreasing EZH2 abundance rather than EZH2 methyltransferase activity may be a key to realizing the therapeutic potential of EZH2 targeting in melanoma³⁵. Further to investigate methyltransferase-independent functions of EZH2 in melanoma, we compared melanoma cells subjected to EZH2 targeting by siEZH2 knockdown, or treatment with the EZH2 degrader DZNep or with EZH2-methyltransferase inhibitors GSK126 and EPZ-6438.

560

Although GSK126 and EPZ-6438 inhibited EZH2 methyltransferase activity as 561 measured by H3K27me3 levels (Fig. 1A, S1A and S1B), they had no effect on the 562 563 growth, clonogenicity, migration, invasion, or pigmentation of BRAFV600E mutant A375 and IGR37 melanoma cells, and only partial effects on NRASQ61K mutant 564 C006-M1 cells (Fig. 1A-1F and S1A-S1H). In contrast, EZH2 knockdown or DZNep 565 treatment displayed marked anti-melanoma effects and promoted melanocytic 566 differentiation in all lines tested (Figure 1A-D and S1A-1H). These findings provide 567 further evidence of methyltransferase independent functions of EZH2 in melanoma. 568

569

To examine methyltransferase-dependent and -independent transcriptional programs of EZH2, we performed global gene expression analysis in B16-F10 murine melanoma cells treated with siEzh2 knockdown vs siRNA control, and also with GSK126 vs DMSO control. 1370 genes were significantly increased by siEzh2 depletion (Figure 1E, 1F), of which 1226 (89.5%) were not upregulated by GSK126 treatment (Figure 1F). By gene ontology (GO) analysis, these 1226 genes were enriched in melanin and cholesterol biosynthesis pathways (Figure 1G). Of the 1119
genes that were downregulated by siEzh2 (Figure 1E-1H), 1087 (97.1 %) were not
changed upon GSK126 treatment (Figure 1H) and strongly enriched in DNA replication
and DNA repair pathways (Figure 1I). These data are consistent with regulation by
EZH2 of methyltransferase-dependent as well as -independent transcriptional
programs in melanoma.

582

Because siRNA might have off-target effects, we also tested a catalytically dead 583 584 mutant of EZH2, H689A, which lacks methyltransferase activity. A375 cells were treated with control (sh-control) or 3' UTR EZH2 region-targeting shRNA (Figure 2A 585 and S2A) to deplete endogenous EZH2, which was then rescued with either wild-type 586 (V5-EZH2-WT) or H689A-mutant EZH2 (V5-EZH2-H689A). In vitro and in vivo, 587 shEZH2 3'UTR knockdown reduced A375 and IGR37 clonogenicity, invasion, wound 588 healing and tumour formation, and increased pigmentation, in a manner that was 589 reversed similarly by ectopic expression of V5-EZH2-WT and V5-EZH2-H689A 590 591 (Figure 2B-F and S2B-D).

592

593 To complement these data, we generated deletion mutants to interrogate EZH2 domains that might rescue shEZH2 knockdown phenotypes (Figure 2G). Partial 594 rescue of A375 cell clonogenicity and full rescue of cell invasion were observed in all 595 shEZH2 3'UTR knockdown cells co-transfected with EZH2 deletion constructs that 596 597 lacked the SET domain, which encodes methyltransferase. In contrast, rescue was not consistently observed with N-terminal EED domain deletion mutants (Δ 1-169) 598 [clone 6] with intact SET domains (Figure S2E, Figure 2G-I), confirming that the N-599 terminal EED domain of EZH2, rather than the methyltransferase-containing SET 600 domain, is critical for the clonogenicity and invasion in melanoma. 601

602

603 Interactions between the EZH2 and IMPDH2

604

To characterize methyltransferase-independent actions of EZH2, we examined its interacting partners in melanoma by immunoprecipitating EZH2 from protein lysates and subjecting the precipitate to liquid chromatography mass spectrometry (LC-MS). Expected PRC2 complex proteins were identified as EZH2 binding partners in four $BRAF^{V600E}$ mutated cell lines, one $NRAS^{Q61K}$ mutated line, and in one $BRAF^{V600E}$ PDX melanoma (Figure 3A and Table S2). Additionally, ubiquitin degradation pathway
proteins UBR4 and NPLOC4, Kinesin 1 complex proteins including KIF5B, KLC1,
KLC2 and KLC4, and Inosine-5'-monophosphate dehydrogenase 2 (IMPDH2), were
all consistently co-immunoprecipitated with EZH2.

614

We focused on IMPDH2-EZH2 interactions because of IMPDH2's known role 615 in GTP metabolism⁵². Further, we found that IMPDH2 protein level is upregulated in 616 human melanoma cells compared to normal human melanocytes (Figure S3A) and 617 618 that high IMPDH2 expression is correlated with poor melanoma survival (p= 0.01, cbioportal) (Figure S3B). To investigate EZH2-IMPDH2 interactions, we verified them 619 endogenously in lysates from A375 cells and PDX tumors by reciprocal Co-IP (Figure 620 3B and S3C), finding. Interactions were reproducibly seen even after GSK126 or 621 EPZ6438 treatment, suggesting they occur independently of EZH2 methyltransferase 622 activity (Figure 3SD). Furthermore, we were unable to detect IMPDH2 methylation in 623 EZH2 Co-IPs by mass spectrometry (Table S3). 624

625

To define interacting domains, we exogenously co-expressed tagged EZH2 (HA-EZH2) with both full length IMPDH2 (MYC/FLAG-IMPDH2) in HEK293 cells (Figure S3E) and with the CBS domain of IMPDH2 [V5-IMPDH2-(1-187)] in A375 cells (Figure 3C). GST pull down assays showed that the CBS domain of IMPDH2 interacts with the N-terminal EED binding domain (1-170) of EZH2 (Figure 3D, 3E). EZH2 EED domain interactions with full length IMPDH2 were verified using exogenously expressed V5-EZH2 deletion mutants and MYC/FLAG-IMPDH2 (Figure 3F).

633

Further to characterize EZH2-IMPDH2 interactions, we also performed 634 proximity ligation assays (PLAs). 60% of A375 cells showed cytosolic EZH2-IMPDH2 635 interactions (<40nm apart, average 15 foci per cell), and 40% showed both cytosolic 636 and nuclear interactions (average 4 foci per cell were nuclear, Figure 3G). These 637 results were supported by western blotting of separated cytosolic and nuclear protein 638 fractions (Figure 3H) and multiplex immunofluorescence labelling of melanoma cell 639 lines (Figure S3F). In contrast, we did not detect cytosolic IMPDH2 interactions with 640 the PRC2 component SUZ12 by PLA, although 10% of cells showed PLA positive 641 nuclear foci (average 4 per cell, Figure 3G). 642

These data indicate that although EZH2 is mostly localized in nuclei, its Nterminal EED domain interacts directly with the IMPDH2-CBS domain predominantly in the cytosol and independently of PRC2 complex formation or EZH2 methyltransferase activity in melanoma cells.

648

649 Cytosolic EZH2 drives melanoma progression

650

To test a potential role for cytoplasmic EZH2 in melanoma progression, we developed an EZH2 mutant lacking a nuclear localization domain (EZH2- Δ NLS). We first generated A375 cells with stable 3'UTR EZH2 knockdown (to minimize endogenous EZH2) and then rescued EZH2 with full length (V5-EZH2-WT) or V5-EZH2-WT- Δ NLS lentiviral constructs. V5-EZH2-WT- Δ NLS expression was mostly cytoplasmic and depleted nuclear EZH2 methyltransferase activity on histone H3K27 (Figure S3G).

658

A375 shEZH2 melanoma cells and xenograft tumors displayed reduced 659 invasion and tumorigenicity that were restored similarly by WT-EZH2 and V5-EZH2-660 661 WT-ANLS (Figure 3I, 3J and S3H). Moreover, overexpression of cytosolic EZH2 lacking methyltransferase activity (V5-EZH2-H689A-ΔNLS) also restored the invasive 662 phenotype of shEZH2 A375 cells to levels comparable to those achieved with V5-663 EZH2-WT-ΔNLS. Although p38-dependent phosphorylation of EZH2 at its T367 664 residue was shown to induce cytosolic localization of EZH2 in breast cancer³⁸, analysis 665 of post-translational modifications of EZH2 in our EZH2-IP LC-MS data did not show 666 significantly different phosphorylation isoforms between cytosolic and nuclear EZH2 667 (Figure S3I, Table S4). Thus, cytoplasmic EZH2 expression is sufficient to promote 668 melanoma cell invasion and tumorigenicity irrespective of EZH2 nuclear or 669 methyltransferase function. 670

671

672 Increased cytosolic localization and activation of IMPDH2 by EZH2

673

We next investigated effects of EZH2-IMPDH2 interactions on IMPDH2 localization and tetramerization/activity. Stable EZH2 knockdown slightly decreased total IMPDH2 protein, but not mRNA expression, that was later rescued by overexpression of EZH2 (1-340) [clone 2] (Figure S3J). Fractionation and immunofluorescence experiments showed that stable or transient EZH2 knockdown
increased the nuclear localization of endogenous IMPDH2 and exogenously
expressed IMPDH2-CBS domain (Figure 3K, S3K, S3L). Conversely, overexpression
of cytosolic wild-type EZH2 (V5-EZH2-WT-ΔNLS) or of cytosolic EZH2 (1-340) [clone
2] increased cytosolic IMPDH2 compared to overexpression of wild type EZH2 (V5EZH2-WT) in endogenous EZH2- silenced A375 cells (Figure 3L, S3M), independently
of EZH2 methyltransferase activity (Figure 3L and S3G).

685

As tetramerization is an essential step in IMPDH2 activation ^{13,16,53}, we 686 investigated effects of EZH2 on IMPDH2 tetramerization and activation in A375 cells. 687 Cross-linked whole-cell extracts showed that IMPDH2 rather than IMPDH1 tetramers 688 were decreased by stable/transient EZH2 knockdown that was rescued by 689 overexpression with wild-type or methyltransferase-deficient EZH2 (full length and 690 EZH2 (1-340) [clone 2], Figures 3M, S3N and S3O). Additionally, co-incubation of 691 IMPDH2 with EZH2 increased basal IMPDH2 activity and reversed GTP-mediated 692 allosteric inhibition of IMPDH2 (Figure 3N). 693

694

695 **GTP-dependent regulation by IMPDH2 of ribosome biogenesis and actomyosin** 696 **contractility**

697

We next assessed pharmacological and genetic inhibition of IMPDH2 in 698 699 melanoma. Treatment with mycophenolic acid (MPA) or ribavirin, pan-IMPDH inhibitors^{12,54,55}, decreased cell proliferation, clonogenicity and invasion that was 700 701 rescued by guanosine addition regardless of B-Raf or N-Ras mutational status (Figure 4A-E and 4A-S4E). MPA and ribavirin also induced pigmentation and senescence 702 (Figure S4F, S4G). siRNA silencing of IMPDH2 also retarded melanoma cell 703 proliferation and invasion that was restored by guanosine addition (Figure 4F, 4G, S4H 704 and S4I). These results implicate a GTP-dependent role for IMPDH2 in melanoma 705 progression. 706

707

IMPDH2-dependent GTP biosynthesis was shown to support rRNA and tRNA
 synthesis⁷. Many tumor cells exhibit increased Pol I activity^{7,56-58}, and GTP-dependent
 Pol I activation has been shown in several cancers^{59,60}. We thus examined RNA
 synthesis by qRT-PCR in A375 cells and found that both MPA treatment and IMPDH2

silencing blunted pre-rRNA (Pol I transcript), and pre-tRNAI13 (Pol III transcript)
expression levels, but not pre-GAPDH mRNA (Pol II transcript), in a time dependent
manner (Figure 4G and S4I). This correlated with triggering nucleolar stress responses
characterized by delocalization of nucleolin and induction of p53 activity (Figure 4H,
4I, S4J, S4K and S4L), with both effects reversed by guanosine (Figure 4H, 4I). We
thus conclude that IMPDH2 regulates ribosome biogenesis in melanoma cells via *de novo* GTP synthesis.

719

GTP is also essential for G-protein activity¹¹, and Rho-GTPases regulate the actomyosin cytoskeleton via ROCK I/II activation and phosphorylation of MLC2 in myosin II to promote melanoma progression^{3,4,61}. MPA treatment and IMPDH2 silencing in melanoma cells reduced RhoA activity and phospho-MLC2/F-actin levels (Figure 4J and S4M), and guanosine again rescued these effects (Figure 4J and S4M), indicating that IMPDH2 regulates actomyosin contractility via GTP synthesis.

726

727 EZH2 promotes clonogenic and invasive phenotypes via IMPDH2 and cellular 728 GTP

729

IMPDH2 is the rate-limiting enzyme in the production of GTP⁶²⁻⁶⁴. Because 730 EZH2 regulated IMPDH2 tetramerization and activity (Figures 3M and 3N), we 731 checked the contribution of EZH2 to cellular GTP production in melanoma. Stable 732 733 EZH2 knockdown reduced GTP levels by 50% in A375 cells, an effect that was reversed by overexpression with N-terminal domain of EZH2 (1-340) [clone 2] (Figure 734 735 5A), which we previously identified as an interaction site for IMPDH2 (Figures 3D-F). Concurrently, EZH2 knockdown also reduced cell proliferation, migration and invasion 736 in a guanosine-reversible manner (Figure 5B and S5A-D). Moreover, EZH2-WT or -737 H689A overexpression induced cell proliferation and invasion, which were reduced to 738 shEZH2 levels by IMPDH2 silencing; again, these effects were rescued by guanosine. 739 These data implicate IMPDH2 as a key intermediary between EZH2 and GTP 740 synthesis in melanoma (Figure 5C, 5D and S5C-D), in which case EZH2 would be 741 expected to modulate critical, IMPDH2- and GTP-dependent functions in cancer cells, 742 such as rRNA metabolism and GTPase activity (Figures 4G-J). 743

Consistent with this, EZH2 knockdown reduced rRNA synthesis and ribosome 745 biogenesis and induces p53 in melanoma cells (Figures 5E, S5E and S5F) in a 746 guanosine-reversible manner (Figure S5E). Additionally, RhoA activity and phospho-747 MLC2 levels were lowered by EZH2 silencing (Figure 5F, 5G, S5G, S5H and S5I) and 748 similarly restored by guanosine (Figure 5H and S5G). Reduced phospho-MLC2 was 749 also seen using siEZH2 constructs or EZH2 degradation by DZNep or MS1943, but 750 not by use of EZH2 methyltransferase inhibitors GSK126 and EPZ-6438 (Figure S5J. 751 S5K), consistent with a methyltransferase-independent role for EZH2 in RhoA 752 753 dependent myosin II activation. In summary, EZH2 regulates levels and critical functions of cellular GTP in melanoma via IMPDH2. 754

755

Nucleolar size and the cytosolic localization of EZH2-IMPDH2 interactions are increased during melanoma progression in patients

758

If cytosolic EZH2-IMPDH2 interactions drive melanoma cell proliferation and 759 invasion as above, then increased cytosolic expression of these proteins might be 760 expected during melanoma progression in patients. To test this, we interrogated the 761 ENSG00000178035-762 Protein Atlas database (https://www.proteinatlas.org/ IMPDH2/tissue/skin#img), observing IMPDH2 expression but undetectable EZH2 763 764 labelling in the nuclei of normal human melanocytes. Consistent with this, in immunostaining of normal human melanocytes and melanoma samples, EZH2 and 765 766 IMPDH2 expression were either not detectable or nuclear in normal melanocytes and stage I melanoma samples (Figures 5I and S5M). In stage IV metastatic melanomas, 767 however, cytosolic EZH2 and IMPDH2 expression were significantly increased and 768 asociated with increased cellular nucleolar size, an indicator of ribosome biogenesis 769 770 (Figure 5I). A functional, methyltransferase-independent link between nucleolar size and EZH2 was verified in tumours from A375 cells in which EZH2 was depleted by 771 shEZH2-3'UTR and then re-expressed with either wild-type or methyltransferase-772 deficient EZH2 (Figure S5L). These data are consistent with the possibility that 773 cytosolic EZH2-IMPDH2 interactions drive ribosome biogenesis during melanoma 774 progression in patients. 775

776

Sappanone A impedes EZH2-IMPDH2 interactions and melanoma progression 778

As EZH2 interacts with CBS domain of IMPDH2 (Figure 3C), we sought to test 779 drugs that can inhibit this interaction. A small molecule called Sappanone A (SA) was 780 demonstrated specifically to inhibit IMPDH2 by directly targeting the conserved 781 cysteine residue 140 (Cys140) in the CBS domain of IMPDH2, inducing an allosteric 782 effect on the catalytic pocket that suppressed IMPDH2 activity. We thus examined 783 effects of SA on EZH2-IMPDH2 interactions and melanoma progression. SA inhibited 784 endogenous EZH2-IMPDH2 interactions. EZH2-IMPDH2-CBS 785 both domain interactions and IMPDH2 tetramerization in A375 and B16-F10 cells in a dose-786 787 dependent manner (Figure 6A-C and S6A-B). In addition, 10 to 20 µM of SA also promoted nuclear localization of IMPDH2 (Figure 6D and S6C). These data indicated 788 that EZH2-IMPDH2 interactions can be targeted by SA. 789

790

In vitro, SA reduced melanoma cell proliferation and clonogenicity dose-791 dependently and in a manner that was reversed by guanosine (Figure 6E and S6D, 792 S6E). Moreover, induction of cell proliferation and invasion by EZH2-WT 793 overexpression following shEZH2 transduction were reduced to those of shEZH2 by 794 5 µM SA treatment, an effect that was able to be rescued by guanosine (Figure 6F 795 796 and S6F). In 3D Matrigel spheroid assays, pre-treatment with SA for 10 days prior to seeding spheroid cultures or treatment of established spheroids demonstrated 797 798 profound anti-melanoma effects, even after 7 days of SA drug washout (Figures 6G-J and S6G-I). SA also induced ribosomal stress and reduced myosin II activation in 799 800 A375 cells (Figure 6M and S6L).

801

To examine normal cells that might be susceptible to targeting the EZH2-802 IMPDH2 interactome, we performed LC/MS on EZH2-immunoprecipitated lysates 803 from CD34⁺ human bone marrow progenitor cells. Although IMPDH2 was detected in 804 anti-EZH2 immunoprecipitates, this could not be verified by CoIP-coupled WB in the 805 same stringency conditions used previously (Table S2 and Figure 6N). Consistent with 806 this, SA has no observable cytotoxic effect on blood progenitors (Figure 6O). Similarly, 807 SA treatment of freshly isolated normal human melanocytes for 7 days at 2 to 5 µM 808 did not significantly attenuate cell growth (Figure 6P). These data indicate that 809 pharmacological inhibition of EZH2-IMPDH2 interactions by SA attenuates melanoma 810 progression without melanocyte and blood cell toxicity by impeding rRNA metabolism 811 and actomyosin contractility. 812

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EZH2-IMPDH2 interactions and Sappanone A treatment in uveal melanoma and non-melanoma cancers

816

Although increased EZH2 and IMPDH2 have been linked to many solid 817 cancers¹⁰, potential interactions between them have not been reported. We thus 818 extended the above studies, examining EZH2 and IMPDH2 levels and interactions in 819 uveal melanoma (92.1 and OMM1), and ovarian (OVCAR-3 and OVCAR-8), breast 820 821 (MCF7 and MDA-MB-231), and prostate (LNCaP and C4:2) cancer cell lines. Cytosolic EZH2/IMPDH2 interactions were seen in all lines tested (Figures 7A-E and S7A-E). 822 Moreover, total EZH2 degradation by MS1943 treatment, but not treatment with 823 GSK126, a EZH2 methyltransferase inhibitor, attenuated OMM1, OVCAR-8, MDA-824 MB-231, and C4-2 cell growth. We therefore also tested effects of SA on EZH2-825 IMPDH2 interactions and IMPDH2 activity in OMM1, OVCAR-8, MDA-MB-231, and 826 PC3 cells, observing reduced EZH2-IMPDH2 interactions (Figure 7F-G, S7F-G) and 827 IMPDH2 tetramerization in every case (Figure S7H-K). These data suggest that 828 cytosolic EZH2-IMPDH2 interactions may be a therapeutic target in a range of cancers 829 830 beyond cutaneous melanoma.

831

Broadly, our findings support a model of EZH2 function in which its canonical role in the repressive PRC2 complex is complemented in at least some cancers by methyltransferase independent cytosolic EZH2 sequestration and binding to IMPDH2 to activate GTP synthesis and facilitate ribosome biogenesis and actomyosin contractility, thereby promoting cancer progression (Figure 7L).

837

838 **DISCUSSION**

839

EZH2 is a bona-fide oncoprotein in cutaneous and uveal melanomas^{24-28,65}, and breast⁶⁶, prostate⁶⁷, and ovarian⁶⁸ cancers, imparting proliferative, migratory, and invasive cancer cell phenotypes. However, the mechanisms through which it imparts these properties are incompletely understood. Here, we reveal a previously unappreciated, methyltransferase-independent function of EZH2 that acts via cytosolic interactions with and activation of IMPDH2 to maintain cellular GTP.

Recently, uveal melanoma cells were shown to be resistant to EZH2 847 methyltransferase inhibition⁶⁹ unless exposed to supraphysiological doses⁷⁰. Further, 848 triple-negative breast cancer MDA-MB-231 cells and castrate-resistant prostate cancer 849 C4-2 and DU145 cells were also reported to be similarly resistant, although sensitive 850 to total EZH2 silencing, suggesting methyltransferase-independent functions of 851 EZH2^{38,67,71}. Using an unbiased 852 proteomics approach. we uncovered methyltransferase-independent binding partners of EZH2 in cutaneous melanoma to 853 identify IMPDH2 as a critical mediator of the oncogenic effects of EZH2. Furthermore, 854 855 we showed that EZH2-IMPDH2 interactions are commonly seen and functionally consequential in other cancers, suggesting that their targeting may represent a 856 common molecular target in human cancer. 857

858

Previous studies found that EZH2 controls melanoma growth and metastasis 859 through transcriptional repression of distinct tumor suppressors, such as ciliary genes 860 and AMD1 in N-Ras mutant tumors^{27,28} and also regulates mechanisms of adaptive 861 resistance to immunotherapy³⁰. Recently, the combination of EZH2 and MEK inhibition 862 was found to reduce tumor burden markedly in NRAS mutant cells, but not BRAF 863 864 mutant cells⁷². This is consistent with our observations of partial anti-melanoma effects following EZH2 methyltransferase inhibition in NRAS mutant melanoma cells. In 865 contrast, BRAF mutant melanoma cells were resistant to EZH2 methyltransferase 866 inhibition but sensitive to total EZH2 silencing. Methyltransferase-dependent functions 867 of EZH2 may be more prominent in NRAS mutant melanomas due to lower expression 868 levels of IMPDH2 and EZH2 in NRAS mutant cell lines compared with BRAF mutant 869 870 cells.

871

Cytosolic localization of EZH2 contributes to pro-metastatic behaviors (i.e. 872 invasion, migration), but this may involve methyltransferase dependent or independent 873 mechanisms in different cell types³⁴. Cytoplasmic localization of EZH2 was observed 874 in murine fibroblasts, where it retained methyltransferase activity and regulated actin 875 polymerization³⁶. In leukocytes, EZH2 methylated the cytoplasmic protein talin-1 to 876 enhance cell migration by inhibiting binding of talin-1 to F-actin⁷³. p38-dependent 877 phosphorylation of EZH2 at the T367 residue was shown to induce cytosolic 878 localization of EZH2 in breast cancer cells, where it interacted with cytoskeletal 879 proteins to promotes metastasis³⁸. Cytoplasmic EZH2 expression has also been 880

observed in prostate cancer cells³⁷. In this study, we identified EZH2 as a regulator of 881 RhoAGTPase activity and actomyosin contractility via RhoA/ROCK/myosin II 882 activation^{3,4,61,74}. Consistent with a role for EZH2 in metastasis, we observed cytosolic 883 localization of EZH2 in melanoma cells particularly in association with more 884 advanced/metastatic disease. However, this was not explained by differential 885 phosphorylation of EZH2. The mechanism of nuclear-cytosolic EZH2 shuttling in 886 melanoma cells remains to be elucidated, although our identification by LC-MS of 887 interactions between EZH2 and kinesin family protein components suggests a role for 888 889 the latter.

890

Our data suggest that by altering the subcellular localization of IMPDH2, 891 cytosolic EZH2 may switch differentiation-inducing nuclear IMPDH2 into proliferation-892 inducing cytosolic IMPDH2. Nuclear IMPDH accumulates during the G2 phase of the 893 cell cycle or following replicative/oxidative stress, and binds to single-stranded, CT-894 rich DNA sequences via its CBS domain⁷⁵⁻⁷⁹. Thus, in nuclei, IMPDH acts as a 895 transcriptional regulator of histones and E2F genes independently of its enzymatic 896 activity⁷⁹. Interestingly, in mouse melanoma cells, Impdh2 disruption by SA reduced 897 898 pigmentation via repression of tyrosinase gene expression⁸⁰.

899

900 In this study, we confirmed that EZH2 alters the subcellular localization of IMPDH, as EZH2 knockdown enhanced nuclear localization of IMPDH2. Functionally, 901 902 gene expression analysis showed that pigmentation-related genes (Tyr, Oca2, Trp1) were upregulated by Ezh2 knockdown independent of its methyltransferase function. 903 These lines of evidence suggest that EZH2 may regulate pigmentation-related gene 904 expression via regulation of IMPDH2 nuclear localization. We cannot exclude the 905 possibility that EZH2 manipulation may induce oxidative stress that induces nuclear 906 IMPDH2 localization in melanoma cells. However, in normal melanocytes where EZH2 907 was not observed, we observed nuclear IMPDH2. Thus, in the absence of EZH2, 908 nuclear IMPDH2 may stabilize melanocyte differentiation in normal melanocytes and 909 910 melanomas, whereas during melanoma progression, augmented cytosolic EZH2 may move IMPDH2 to the cytosol where its GTP-producing enzymatic activity supports cell 911 912 growth and invasion.

Pharmacological targeting of IMPDH2 may represent a tolerable therapeutic 914 strategy in EZH2-IMPDH2 activated cancers. Although trials of pan-IMPDH inhibitors, 915 such as MPA, tiazofurin and benzamide riboside, have been conducted in patients 916 with leukemia and multiple myeloma⁸¹⁻⁸⁴, these studies were terminated due to 917 neurotoxic side effects^{52,85,86}. Because IMPDH2 is mainly expressed in rapidly 918 proliferating immunocytes, in contrast to the IMPDH1 isoform in normal human 919 leukocytes and lymphocytes^{87,88}, MPA was shown recently to have more 920 hematological side effects than the IMPDH2 specific inhibitor, SA, in vivo¹⁶. Consistent 921 922 with this, we demonstrated low or no EZH2-IMPDH2 interactions in CD34⁺ human blood progenitors and SA demonstrated no anti-proliferative effects on these cells, in 923 contrast to melanoma cells. Although we found that MPA is anti-tumorigenic and anti-924 metastatic in melanoma cells, IMPDH2 specific inhibitors are likely to be better and 925 better tolerated treatment options. 926

927

In conclusion, we report a role for the EZH2 oncoprotein in promoting tumorigenesis and metastasis in melanoma cells by interacting with IMPDH2 to promote GTP generation, and thereby mechanisms such as upregulating rRNA metabolism and actomyosin contractility that support cancer progression. Discovery of this previously unappreciated, non-enzymatic, GTP-dependent function of EZH2 opens new avenues for EZH2-targeted therapeutics.

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1136 Main Figures

1137 Figure1.



1138

Figure 1. Methylation-independent functions of EZH2 are predominant in 1139 melanoma. A375 cells were treated with siEZH2, 2 µM DZNep, 2 µM GSK126, 2 µM 1140 EPZ6438 and scramble or DMSO (control) for 3 days prior to: (A) Western blot analysis 1141 of EZH2, H3K27me3, H3 and β -Actin protein level, (B) cell growth analysis done by 1142 Trypan Blue haemocytometer counting, (C) clonogenicity after low-density seeding 1143 (crystal violet stain). Clonogenicity was assessed in pre-treated (3 days) cells seeded 1144 at 2000 cells in 6-well plate followed by crystal violet staining (0.5% in methanol) after 1145 incubation for 10 days in drug-free media. Representative images after crystal violet-1146 1147 stained wells were shown above bars (D) Boyden chamber migration was assessed in pre-treated (3 days) cells seeded at 50,000 cells in 24-well plate after incubation for 1148 24h. Representative images after crystal violet-stained wells were shown above bars. 1149

1150 (E) B16-F10 cells were treated with either GSK126 versus vehicle control (DMSO) or siEzh2 versus siCtrl and then profiled in triplicate RNAseg experiments. Genes that 1151 were significantly up- or downregulated by siEzh2 compared with the control were 1152 clustered across all samples and are shown as heatmaps. Each row represents one 1153 gene and each column triplicate sample. The siEzh2-induced genes that were also 1154 induced by GSK126 were termed class I genes and those unchanged by GSK126 1155 class II genes. Genes that were activated by Ezh2 were defined as class III genes. (F) 1156 Venn diagram showing overlap among si-Ezh2 induced and GSK126-induced genes 1157 1158 compiled from RNAseq experiment in G. (G) GO biological process analysis of 1226 class II genes. (H) Venn diagram showing overlap among si-Ezh2 repressed and 1159 GSK126-repressed genes compiled from RNAseq experiment in G. (I) GO biological 1160 process analysis of 1087 class III genes. Data for B-D are from three independent 1161 experiments and are presented as mean ±SD, analyzed by one-way ANOVA plus 1162 Tukey's multiple comparison tests. ns: non-significant. 1163

1165 Figure 2.



1166

Figure 2. EZH2 has methyltransferase independent function in melanoma 1167 tumorigenicity and invasion. (A) Western blot analysis of A375 cells showing EZH2 1168 knockdown after lentiviral transduction with control shRNA (shControl) or 3' UTR 1169 EZH2-targeting shRNA (shEZH2) and rescue with V5-tagged WT-EZH2 or 1170 methyltransferase deficient H689A-EZH2. (B) Clonogenicity assay of cells described 1171 in A. Representative images after crystal violet-stained wells were shown above bars. 1172 (C) Matrigel-coated Boyden chamber invasion assay of cells described in A. 1173 1174 Representative images after crystal violet staining were shown above bars. (D) Wound healing assay of cells described in A. Representative images of the wound after 24 h 1175 shown above bars. (E) Tumor caliper of A375 xenografts as described in A. (F) Tumor 1176 weights of A375 xenografts at the end point. Representative tumors per group were 1177

shown above bars. (G) Western blot analysis, (H) invasion, (I) clonogenicity of A375 cells with EZH2 knockdown followed by rescue with V5-tagged EZH2 deletion mutant vectors. Data for B-D, H, I are from at least three independent experiments and are presented as mean \pm SD, analyzed by one-way ANOVA plus Tukey's multiple comparison tests. Data for E, F are from 7 mice per group and are presented as mean \pm SD, analyzed by two-way ANOVA plus Tukey's multiple comparison tests. ns: nonsignificant.

1186 Figure 3.



1187

1188 Figure 3. Cytosolic EZH2 interacts with IMPDH2 through the IMPDH2-CBS domain and moves IMPDH2 to cytoplasm/ increases its tetramerization-1189 1190 mediated activity. A) List of overlapping proteins co-immunoprecipitated (Co-IP) with EZH2 from C006-M1, LM-MEL-28:B4:F3, IGR37, A375 and LM-MEL-45 melanoma 1191 cells (all data derived from n=3 biological replicates). (B) The interaction between 1192 endogenous EZH2 and IMPDH2 was determined in A375 cells by immunoprecipitation 1193 (IP) with anti-IMPDH2 and anti-EZH2 antibody followed by western blotting with anti-1194 EZH2 and anti-IMPDH2 antibody. (C) HA-tagged EZH2-WT and V5-tagged IMPDH2 1195 (1-187) were co-expressed in A375 cells. The interaction between overexpressed 1196 EZH2 and IMPDH2 (1-187) was determined by immunoprecipitation with anti-HA 1197 antibody followed by western blotting with anti-V5 antibody. (D) GST-EZH2 deletion 1198

mutant constructs. (E) The binding of V5-IMPDH2-CBS protein to GST-EZH2 1199 peptides was probed with WB using the V5 specific antibody. Total cell lysate from 1200 HEK293 overexpressing V5-IMPDH2-CBS was used as a source of IMPDH2-CBS in 1201 GST-pull-down experiment. (F) The binding of Myc/Flag tagged full length IMPDH2 1202 protein to V5 tagged EZH2 deletion mutant peptides (shown in Fig. 2G) was shown by 1203 co-IP with anti-Flag antibody followed by probing with anti-V5 antibody. (G) Ligation 1204 1205 proximity images depicting co-localization with EZH2 and IMPDH2 by red fluorescent dots A375 cells. Scale bar=10 µm. Number of interaction loci depicted as red dots 1206 1207 were counted for cytoplasm and nucleus of total of 100 cell. (H) Cytosolic/Nuclear fractionation was done for A375, B4:F3, IGR37, B16-F10 cells followed by IP with anti-1208 EZH2 antibody followed by western blotting with anti-EZH2 and anti-IMPDH2 antibody 1209 (right). Lamin A/C is nuclear, and β -Actin is cytosolic marker. Inputs were shown on 1210 the left. (I) Matrigel-coated Boyden chamber invasion assay of A375 cells showing 1211 EZH2 knockdown after lentiviral transduction with control shRNA (shControl) or 3' UTR 1212 EZH2-targeting shRNA (shEZH2) and rescue with V5-WT-EZH2 or V5-EZH2-H689A, 1213 V5-EZH2-ANLS-WT and V5-EZH2-ANLS-H689A. Representative images after crystal 1214 violet staining were shown above bars. (J) Tumor weights of indicated A375 xenografts 1215 1216 (n=7) at the end point. (K) Cytosolic/Nuclear fractionation was done from A375 cells with control shRNA (shControl) or 3' UTR EZH2-targeting shRNA followed by IP with 1217 anti-EZH2 antibody followed by western blotting with anti-EZH2 and anti-IMPDH2 1218 antibody. Lamin A/C is nuclear, and β -Actin is cytosolic marker. (L) Cytosolic/Nuclear 1219 1220 fractionation was done from cells described in (I) followed by western blotting with anti-V5, anti-EZH2, anti-IMPDH2 antibody and β-Actin antibody. (M) The clusters of 1221 1222 IMPDH2 and IMPDH1 tetramer were detected from cross-linked whole-cell extracts isolated from cells described in Figure 2A. (N) Relative IMPDH2 activity measured by 1223 NADH absorbance at OD340nm. 2 µg of recombinant IMPDH2 was preincubated with 1224 and 3 mM IMP and 2 µg of recombinant EZH2 and then 10mM GTP was added for 10 1225 min prior to reaction initiation by with 1 mM NAD⁺. Data for I, J and N is from three 1226 independent experiments and are presented as mean ±SD, analyzed by one-way 1227 ANOVA plus Tukey's multiple comparison tests. ns: non-significant. 1228

1230 Figure 4.



1231

Figure 4. IMPDH2 induces clonogenecity/ invasion in melanoma cells by 1232 regulating ribosome biogenesis and actomyosin contractility via cellular GTP 1233 level regulation. (A) Time-dependent growth curves of A375 cells upon 25 µM 1234 Ribavirin or DMSO (control); 3 µM MPA, or methanol (Control). (B) A375 cell growth 1235 analysis done by Trypan Blue haemocytometer counting after treated with 3 µM MPA 1236 or methanol control with the addition of 100 µM guanosine or vehicle control for 3 days. 1237 (C) Clonogenicity assay of A375 cells described in A. Clonogenicity was assessed in 1238 pre-treated (2 and 3 days) cells seeded at 2000 cells in 6-well plate followed by crystal 1239 violet staining (0.5% in methanol) after incubation for 10 days in drug-free media. 1240 Representative images after crystal violet-stained wells were shown. (D) 3D matrigel 1241 clonogenicity assay of A375 and B16-F10 cells treated with 3 µM MPA or methanol 1242

1243 (control) for 10 days. (E) Matrigel-coated Boyden chamber invasion assay of cells described in B. (F) Cell growth analysis done by Trypan Blue haemocytometer 1244 counting after A375 cells were treated with siIMPDH2 or scramble control with the 1245 addition of 100 µM guanosine or vehicle control for 3 days. (G) Matrigel-coated Boyden 1246 chamber invasion assav of cells described in F. (H) Nascent transcripts of the 1247 indicated genes were analyzed by gRT-PCR after A375 cells were treated with 3 µM 1248 MPA for 0h. 4h and 8h. (I) IF staining with anti-NCL antibody in A375 cells treated with 1249 25 µM Ribavirin, 3 µM MPA, or methanol (Control) with the addition of 100 µM 1250 1251 guanosine or vehicle control for 24h. DAPI stains the nuclei. Scale bar: 20 µm (J) RhoA activity assay in A375 cells treated with 3 µM MPA or methanol (Control) with the 1252 addition of 100 µM guanosine for 24h. (K) IF staining of A375 cells described in J with 1253 anti-p-MLC2 (green) and phalloidin (red). DAPI stains the nuclei. Scale bar: 20 µm. % 1254 peripheral p-MLC2 positive cells were plotted on the right plot. Data for A, B, F, H, J 1255 and K are from three independent experiments and are presented as mean ±SD, 1256 analyzed by one-way or two-way ANOVA plus Tukey's multiple comparison test. ns: 1257 non-significant. 1258

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1260 Figure 5.



1261

Figure 5. EZH2 regulates clonogenicity/ invasion by regulating rRNA 1262 metabolism and Rho GTPase activity via GTP production in melanoma. (A) 1263 Relative GTP levels were quantified by HPLC in A375 cells with stable EZH2 1264 knockdown (shEZH2) and overexpression with V5-EZH2-clone2. (B) Cell growth 1265 analysis of A375 cells with stable EZH2 knockdown after 3 days of control or 100 µM 1266 1267 guanosine addition done by Trypan Blue haemocytometer counting. (C) Cell growth analysis done by Trypan Blue haemocytometer counting and (D) invasion assay 1268 1269 counting done by crystal violet staining after A375 cells with stable EZH2 knockdown were rescued by V5-tagged WT-EZH2 or methyltransferase deficient H689A-EZH2 1270 overexpression followed by scramble, si-IMPDH2#1, or si-IMPDH2#2 oligos and 100 1271 µM guanosine addition. (E) Nascent transcripts of the indicated genes were analyzed 1272

by qRT-PCR, (F) RhoA activity assay, (G) p-MLC2 IF in A375 cells showing EZH2 1273 knockdown after lentiviral transduction with control shRNA (shControl) or 3' UTR 1274 1275 EZH2-targeting shRNA (shEZH2) and rescue with V5-tagged WT-EZH2 or methyltransferase deficient H689A-EZH2. % Peripheral p-MLC2 positive cells were 1276 plotted below images. (H) % peripheral p-MLC2 positive cells were plotted. (I) Human 1277 melanoma samples from grade I to IV were stained with anti-EZH2 and anti-IMPDH2 1278 1279 antibodies. Grade I, II, III: n=31 grade IV: n=8. Nucleolar sizes were measured from HE stained samples. Scale bar: 50 µm. Data are presented as mean ±SD, analyzed 1280 by student t-test. Data for A-H are from three independent experiments and are 1281 presented as mean ±SD, analyzed by one-way or two-way ANOVA plus Tukey's 1282 multiple comparison test. ns: non-significant. 1283

1285 Figure 6.



1286

Figure 6. Pharmacological inhibition of EZH2/IMPDH2 interactions by SA 1287 attenuates the growth and invasion abilities of melanoma cells in vitro. (A) The 1288 1289 interaction between endogenous EZH2 and IMPDH2 upon 16h SA treatment (DMSO, 5, 10, 20 µM) was determined in A375 cells by IP with anti-EZH2 antibody followed by 1290 WB with anti-EZH2 and anti-IMPDH2 antibody. The inputs were shown on the left. (B) 1291 The interaction between HA-EZH2 and V5-IMPDH2-CBS upon 16h SA treatment 1292 (DMSO, 2, 5, 10 µM) was determined in A375 cells by IP with anti-HA antibody 1293 followed by WB with anti-V5 and anti-HA antibody. The inputs were shown above the 1294 IP blots. (C) The clusters of IMPDH2 tetramer were detected from cross-linked whole-1295 cell extracts isolated from A375 cells treated with SA for 16h. (D) Cytosolic versus 1296 1297 nuclear localizations of EZH2 and IMPDH2 were examined upon SA treatment in A375 cells by Co-IF. % Nuclear IMPDH2 positive cells were plotted on the group. (E) Dose 1298

1299 dependent cell growth curve of A375 cells treated with the indicated dose of SA and -/+ 100 µM guanosine for 3 days. Clonogenicity was shown in the inlet. (F) Matrigel-1300 1301 coated Boyden chamber invasion assay in A375 cells with stable EZH2 knockdown and later rescued by V5-tagged WT-EZH2 overexpression followed by scramble, si-1302 IMPDH2#1, or si-IMPDH2#2 oligos and 100 µM guanosine addition. (left), invaded cell 1303 numbers per field were plotted on the right graph. (G) Spheroid areas of 3D colonies 1304 grown for 4, 10, 14 days with DMSO or 5 µM SA containing culture medium were 1305 measured by Image J program. (H) Sphere formation in 3D Matrigel (therapeutic). 1306 1307 A375 cells were grown in Matrigel for 6 days in the absence of SA followed by 4d and 10d days with DMSO or 10 µM SA. Spheroid areas were measured by Image J 1308 program and presented in the graph (I). Invasive spheroid numbers were counted 1309 manually and presented in the graph (J). (K) Sphere formation in 3D Matrigel 1310 (preventive). A375 cells were grown in Matrigel for 10 days in presence of either 1311 DMSO (control) or 10 µM SA and then the colonies were grown 7 more days without 1312 SA or DMSO and spheres were counted manually and presented in the graph (L). (M) 1313 The effect of SA on ribosome biogenesis was measured in A375 cells treated with the 1314 indicated doses of SA by anti-NCL antibody. DAPI stains the nuclei. Scale bar: 20 µm. 1315 1316 (N) The effect of SA on EZH2 and IMPDH2 interaction was shown by Co-IP coupled WB in CD34⁺ BM cells. Cell growth analysis of CD34⁺ bone marrow progenitor cells 1317 (n=2 patients in triplicates) (O) and normal human melanocytes (n=4) (P) treated with 1318 DMSO (vehicle), 2 µM, or 5 µM SA for the indicated time points. Data for D, E, F, G, I, 1319 1320 J and L are from three independent experiments and are presented as mean ±SD, analyzed by one-way or two-way ANOVA plus Tukey's multiple comparison test. ns: 1321 non-significant. 1322

1324 Figure 7.



1325

Figure 7. EZH2-IMPDH2 interaction is commonly seen in uveal melanoma, 1326 breast, prostate, ovarian cancer, and SA attenuates their growth in vitro. EZH2 1327 and IMPDH2 interaction was shown by IP with anti-IMPDH2 antibody followed by WB 1328 with anti-EZH2 and anti-IMPDH2 antibody in (A) uveal melanoma, (B) ovarian cancer, 1329 (C) breast cancer and (D) prostate cancer cell lines. Inputs were shown at the top of 1330 each Co-IP blots. (E) Cytosolic/Nuclear fractionation was done for MDA-MB-231, C4-1331 2, PC3, OVCAR8 and OMM1 cells followed by IP with anti-EZH2 antibody followed by 1332 western blotting with anti-EZH2 and anti-IMPDH2 antibody. The effect of SA on EZH2 1333 and IMPDH2 interaction was shown by Co-IP coupled WB in (F) OMM1 and (G) 1334 OVCAR8 cells. Dose-dependent growth curves of (H) OMM1, (I) OVCAR8, (J) MDA-1335 MB-231 and (K) C4:2 cells upon SA treatment for 3 days. (L) Proposed model 1336

depicting both canonical nuclear and non-canonical cytosolic functions of EZH2 as an epigenetic silencer and as GTP regulator via IMPDH2 interaction, which can be blocked by SA. EZH2 induces tumorigenicity and metastasis in melanoma by upregulating rRNA metabolism and RhoA dependent actomyosin contractility via GTP production.

1343 Supplemental Figures

1344 Figure S1.



1345

Figure S1. Pharmacological inhibition of EZH2 abundance, but not its activity 1346 1347 reduces melanoma cell growth/ invasion and induces pigmentation. C006-M1 and IGR37 cells were treated with siEZH2, 2 µM DZNep, 2 µM GSK126, 2 µM 1348 EPZ6438 and scramble or DMSO (control) for 3 days prior to: (A, B) Western blot 1349 analysis of EZH2, H3K27me3, H3 and β-Actin protein level, (C, D) cell growth analysis 1350 done by Trypan Blue haemocytometer counting, (E) Matrigel-coated Boyden chamber 1351 invasion assay was assessed in pre-treated (3 days) cells seeded at 100,000 cells in 1352 24-well plate coated with matrigel after incubation for 24h. (F) Invaded cell counts per 1353 well were done by CV staining. (G) Cell pigmentation was assessed by Fontana 1354 Masson staining. (H) Pigmented cell percentages were calculated per well. Data for 1355 C, D, F and H are from three independent experiments and are presented as mean 1356

- 1357 ±SD, analyzed by one-way ANOVA plus Tukey's multiple comparison test. ns: non-
- 1358 significant.

1360 Figure S2.



Figure S2. EZH2 has methyltransferase independent function in melanoma 1362 1363 clonogenicity, invasion and pigmentation. (A) Western blot analysis of IGR37 cells showing EZH2 knockdown after lentiviral transduction with control shRNA (shControl) 1364 1365 or 3' UTR EZH2-targeting shRNA (shEZH2) and rescue with V5-tagged WT-EZH2 or methyltransferase deficient H689A-EZH2. (B) Clonogenicity assay of cells described 1366 in A. Representative images after crystal violet-stained wells were shown above bars 1367 1368 and representative images of cell pellets were shown below bars. (C) Matrigel-coated Boyden chamber invasion assay of cells described in A. Representative images after 1369 crystal violet staining were shown above bars. (D). Western blot analysis of EZH2, V5, 1370 H3K27me3, H3 and β-Actin from A375 xenograft tumor lysates. (E) V5-EZH2 deletion 1371 mutant constructs. Data for B, C are from three independent experiments and are 1372 presented as mean ±SD, analyzed by one-way ANOVA plus Tukey's multiple 1373 comparison test. 1374

1375

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1376 Figure S3.



1377

Figure S3. EZH2 interacts with IMPDH2 and induces its tetramerization 1378 methyltransferase independently. (A) Western blot analysis of EZH2, IMPDH2 and 1379 β-Actin in C006-M1 (NRASQ61K), IGR37 (BRAFV600E), LM-MEL28: B4:F3 1380 (BRAFV600E), C32 (BRAFV600E), SK-MEL28 (BRAFV600E), A375 (BRAFV600E), 1381 LM-MEL33 (BRAFV600E), LM-MEL45 (BRAFV600E), IGR39 1382 (BRAFV600E) melanoma cells and normal human melanocytes (NHM). (B) Kaplan-Meier curves of 1383 overall survival of TCGA PanCancer Atlas cutaneous melanoma patients (n = 392 1384 patients), stratified by IMPDH2 mRNA levels. Data were analyzed by log rank test. (C) 1385 The interaction between endogenous EZH2 and IMPDH2 was determined in PDX 1386 tumor lysates by IP with anti-IMPDH2 and anti-EZH2 antibody followed by WB with 1387 anti-EZH2 and anti-IMPDH2 antibody. (D) A375 cells were treated with DMSO 1388

(control), 2 µM GSK126 or 2 µM EPZ6438 for 2 days prior to: (A) Western blot analysis 1389 of EZH2, H3K27me3, H3 and β-Actin (left) and interaction of EZH2/ IMPDH2 were 1390 1391 shown by IP with anti-IMPDH2 antibody followed by WB with EZH2 antibody. (E) HAtagged EZH2-WT and MYC/FLAG-tagged IMPDH2-WT were co-expressed in 1392 HEK293 cells. The interaction between overexpressed EZH2 and IMPDH2 was 1393 determined by immunoprecipitation with anti-HA antibody followed by western blotting 1394 1395 with anti-Myc-tag antibody. (F) Co-immunofluorescence (Co-IF) staining with anti-EZH2 and anti-IMPDH2 antibodies. DAPI stains the nuclei. Scale bar: 20 µm (G) 1396 1397 Western blot analysis of EZH2, V5, H3K27me3, H3 and β -Actin in A375 cells described in Figure 3F. (H) Tumor calipers of indicated A375 xenografts (n=7) at the 1398 end point. Data are presented as mean ±SD, analyzed by two-way ANOVA plus 1399 Tukey's multiple comparison test. ns: non-significant. (I) Cytosolic and nuclear Ezh2 1400 phosphorylation sites and their percentages measured by LC-MS. Known kinases for 1401 the corresponding phospho-sites were also included on the last column of the table.?: 1402 Unknown kinases. (J) Western blot of of EZH2, V5, H3K27me3, H3 and β-Actin in 1403 A375 cells with control shRNA (shControl) or 3' UTR EZH2-targeting shRNA and V5-1404 1405 EZH2-clone2. (K) Co-IF staining with anti-EZH2 and anti-IMPDH2 antibodies in C006-1406 M1 cells treated with scramble control or siEZH2 for 3 days. DAPI stains the nuclei. Scale bar: 20 µm (L) IF staining with anti-V5 antibody in A375 cells harboring V5-1407 1408 IMPDH2 (1-187) that was treated with scramble control or siEZH2 for 3 days. DAPI stains the nuclei. Scale bar: 20 µm. (M) Cytosolic/Nuclear fractionation was done from 1409 1410 A375 cells shown in J. Lamin A/C is nuclear, and β -Actin is cytosolic marker. The clusters of IMPDH2 tetramer were measured after cross-linking of B16-F10 cells (N) 1411 1412 treated with or scramble control, siEzh2#1 orsiEzh2#2 for 3 days and of A375 cells shown in J (O). 1413

1415 Figure S4.



1416

Figure S4. Pharmacological or genetic inhibition of IMPDH2 reduces 1417 1418 clonogenicity, invasion by p53 induction and ROCK-myosin II pathway activation. Time-dependent growth curves of A375 (A), C006-M1 (B), B4:F3 (C) cells 1419 1420 upon 25 µM Ribavirin or DMSO (control); 3 µM MPA, or methanol (Control). (D) Clonogenicity assay of B16-F10 cells described in A. (E) Matrigel-coated Boyden 1421 chamber invasion assay of cells described in A, B. (F) Bright field images of cells 1422 described in A, B, C. (G) B16-F10 cell senescence determined by β-gal staining 1423 (green). (H) Cell growth analysis done by Trypan Blue haemocytometer counting, (I) 1424 Western blot analysis of IMPDH2, *β*-Actin, *p*-MLC2 and MLC2 in A375 cells after 1425 1426 treated with siIMPDH2#1, siIMPDH2#2 or scramble (control) for 3 days. (J, K) Western blot analysis of p53 and β-Actin in A375 cells with 0h, 4h, 8h 3 µM MPA treatment -/+ 1427

1428 100 μ M guanosine. (L) Co-IF staining of A375 cells treated with 25 μ M Ribavirin or 1429 DMSO (control); 3 μ M MPA with anti-p53 (green) and anti-p21 (red) or anti-NCL (red), 1430 anti-FBL (green). DAPI stains the nuclei. Scale bar: 100 μ m. (M) IF staining of A375 1431 cells described in H with anti-p-MLC2 (green) and phalloidin (red). DAPI stains the 1432 nuclei. Scale bar: 20 μ m. Data for A, B, C and H are from three independent 1433 experiments and are presented as mean ±SD, analyzed by one-way ANOVA plus 1434 Tukey's multiple comparison test.

1435

1437 Figure S5.



1438

Figure S5. EZH2 modulates ROCK-myosin II activity via Rho GTPase regulation 1439 in melanoma cells. (A) Western blot analysis of EZH2, IMPDH2, H3K27me3, H3, β-1440 1441 Actin, (B) Matrigel-coated Boyden chamber invasion assay in A375 cells with stable EZH2 knockdown (shEZH2) -/+ 100 µM guanosine for 3 days. (C) Western blot 1442 analysis of IMPDH2, EZH2 and β-Actin and (D) matrigel-coated Boyden chamber 1443 invasion assay in A375 cells with stable EZH2 knockdown and rescue with V5-tagged 1444 WT-EZH2 with scramble, si-IMPDH2#1, or si-IMPDH2#2 oligos and 100 µM 1445 guanosine addition. (E) IF staining with anti-NCL antibody in A375 cells with stable 1446 1447 EZH2 knockdown (shEZH2) -/+ 100 µM guanosine for 3 days. (F) Western blot analysis of p53, EZH2, H3K27me3, H3, β-Actin (top) and qRT-PCR of p53, MDM2, 1448 PUMA, CDKN2A (bottom) in A375 cells with stable EZH2 knockdown. (G) p-MLC2 IF 1449

1450 in A375 cells with stable EZH2 knockdown (shEZH2) -/+ 100 µM guanosine for 2 days. (H) Western blot analysis of p-MLC2, RhoA, EZH2, V5, β-Actin, H3K27me3 and H3 in 1451 A375 cells showing EZH2 knockdown shRNA (shControl) or 3' UTR EZH2-targeting 1452 1453 shRNA (shEZH2) and rescue with V5-tagged WT-EZH2 or methyltransferase deficient 1454 H689A-EZH2. (I) Western blot analysis of p-MLC2. MLC2. EZH2 and β-Actin in B16-F10 cells treated with scramble (control), siEzh2#1, or siEzh2#2 for 3 days. IF staining 1455 with anti-p-MLC2 antibody in B16-F10 (J) and A375 cells (K) treated with siEzh2#1. 1456 siEzh2#2, 2 µM DZNep, 2 µM GSK126, or 2 µM EPZ6438 for 2 days. DAPI stains the 1457 1458 nuclei. Scale bar: 20 µm. % peripheral p-MLC2 positive cells were plotted next to the images. (L) Nucleolar sizes were measured from HE stained xenograft samples (n=7) 1459 obtained in Fig. 2F. Scale bar: 50 µm. (M) Co-IF with anti-DCT (melanocyte marker), 1460 anti-EZH2 and anti-IMPDH2 in normal human skin samples. Data for F, J, K and L are 1461 from three independent experiments and are presented as mean ±SD, analyzed by 1462 one-way ANOVA plus Tukey's multiple comparison test. ns: non-significant. 1463

1465 **Figure S6.**



1466

Figure S6. SA reduces EZH2/IMPDH2 interaction, IMPDH2 tetramerization/ 1467 nuclear translocation and attenuates the growth and invasion abilities of 1468 melanoma cells in vitro. (A) The interaction between endogenous EZH2 and 1469 IMPDH2 upon 16h SA treatment (DMSO, 5, 10, 20 µM) was determined in B16-F10 1470 cells by Co-IP with anti-EZH2 antibody followed by WB with anti-EZH2 and anti-1471 IMPDH2 antibody. The inputs were shown on the left. (B) The clusters of IMPDH2 1472 tetramer were detected from cross-linked whole-cell extracts isolated from B16-F10 1473 cells treated with indicated dose of SA for 16h. (C) Cytosolic versus nuclear 1474 localizations of EZH2 and IMPDH2 were examined upon SA treatment in A375 cells 1475 by Co-IF using anti-EZH2 (green) and anti-IMPDH2 (red) antibodies. DAPI stains the 1476 nuclei. Scale bar: 20 µm. (D) Clonogenicity assay of A375 cells described in A. (E) 1477 Dose dependent cell growth curve of B16-F10 cells treated with the indicated doses 1478

1479 of SA and -/+ 100 µM guanosine for 3 days. (F) Cell growth analysis of A375 cells with stable EZH2 knockdown and rescue with V5-tagged WT-EZH2 or methyltransferase 1480 deficient H689A-EZH2 followed by DMSO or 5 µM SA and 100 µM guanosine addition. 1481 (G) Time dependent sphere formation in 3D Matrigel (preventive). A375 and B16-F10 1482 cells were grown in Matrigel for 4 d. 10 d and 14 d in the presence of either DMSO 1483 (control) or 5 µM SA. (H) Sphere formation in 3D Matrigel (therapeutic). B16-F10 cells 1484 were grown in Matrigel for 6 days in the absence of SA followed by 4d and 10d days 1485 with DMSO or 10 µM SA. Spheroid areas were measured by Image J program and 1486 1487 presented in the graph (I). (J) Sphere formation in 3D Matrigel (preventive). B16-F10 cells were grown in Matrigel for 10 days in presence of either DMSO (control) or 10 1488 µM SA and then the colonies were grown 7 more days without SA or DMSO and 1489 spheres were counted manually and presented in the graph (K). (L) The effect of SA 1490 on actomyosin contractility was measured in A375 cells treated with DMSO or 5 µM 1491 SA for 3 days with anti-p-MLC2 antibody (green)/ Phalloidin (red) IF. DAPI stains the 1492 nuclei. Scale bar: 20 µm. Data for E, F, I and K are from three independent 1493 experiments and are presented as mean ±SD, analyzed by one-way or two-way 1494 ANOVA plus Tukey's multiple comparison test. ns: non-significant. 1495

1497 **Figure S7.**



Figure S7. Cytosolic EZH2-IMPDH2 interaction is seen in uveal melanoma, 1499 breast, prostate, ovarian cancer, and SA attenuates IMPDH2 tetramerization. 1500 EZH2 and IMPDH2 interaction was shown by IP with anti-EZH2 antibody followed by 1501 WB with anti-EZH2 and anti-IMPDH2 antibody in (A) uveal melanoma, (B) ovarian 1502 cancer, (C) breast cancer and (D) prostate cancer cell lines. (E) Cytosolic/Nuclear 1503 fractionation was done for OMM1, OVCAR8, MDA-MD231 and C4:2 cells followed by 1504 IP with anti-IMPDH2 antibody followed by western blotting with anti-EZH2 and anti-1505 IMPDH2 antibody. Lamin A/C is nuclear, and β -Actin is cytosolic marker. Inputs were 1506 shown above the IP blot. The effect of SA on EZH2 and IMPDH2 interaction was 1507 1508 shown by Co-IP coupled WB in (F) MDA-MB-231 and (G) PC3 cells. The clusters of IMPDH2 tetramer were detected from cross-linked whole-cell extracts isolated from 1509 (H) OMM1, (I) OVCAR8, (J) MDA-MB-231 and (K) PC3 cells treated with the indicated 1510 dose of SA for 16h. 1511

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