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7 8	Control of topoisomerase II activity and chemotherapeutic inhibition by TCA cycle metabolites
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#### 27 **SUMMARY**

- 28 Topoisomerase II (topo II) is essential for disentangling newly replicated chromosomes.
- 29 DNA unlinking involves the physical passage of one DNA duplex through another and
- 30 depends on the transient formation of double-strand DNA breaks, a step exploited by
- 31 frontline chemotherapeutics to kill cancer cells. Although anti-topo II drugs are
- 32 efficacious, they also elicit cytotoxic side effects in normal cells; insights into how topo II
- is regulated in different cellular contexts is essential to improve their targeted use. Using
- 34 chemical fractionation and mass spectrometry, we have discovered that topo II is
- 35 subject to metabolic control through the TCA cycle. We show that TCA metabolites
- 36 stimulate topo II activity *in vitro* and that levels of TCA flux modulate cellular sensitivity
- to anti-topo II drugs *in vivo*. Our works reveals an unanticipated connection between the
- control of DNA topology and cellular metabolism, a finding with important ramifications
- 39 for the clinical use of anti-topo II therapies.
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- 41

## 42 Key Words

- 43 Topoisomerase, TCA cycle, cancer, metabolism, chemotherapy, etoposide,
- 44 dexrazoxane, DNA topology
- 45

#### 46 INTRODUCTION

The complement of chromosomes in a single human cell, laid out end-to-end, is 47 48 ~10,000 times longer than the diameter of the nucleus in which they reside. Efficient packaging, access, and duplication of this genetic material depends on the proper 49 50 control of DNA superstructure. Some of the largest physical rearrangements of the genome occur during DNA replication and mitosis as sister chromatids are synthesized. 51 52 condensed, and segregated into two daughter cells (Barrington et al., 2017). The double-helical structure of DNA presents a challenge to these transformations, resulting 53 54 in superhelical intertwinings and chromosomal entanglements (catenanes) (Peter et al., 1998; Postow et al., 2001; Sundin and Varshavsky, 1981). Enzymes known as 55 56 topoisomerases are required by cells to resolve topological stresses in DNA; of the two 57 major classes, only type II topoisomerases are able to unlink tangled double-stranded 58 segments to facilitate chromosome partitioning prior to cell division (Vos et al., 2011). 59

Beyond supporting supercoiling homeostasis and promoting DNA unlinking, 60 61 mounting evidence suggests that topo II has additional roles in regulating chromatin 62 structure throughout the eukaryotic cell cycle (Lee and Berger, 2019). Although most 63 DNA regions are decatenated by the end of replication in S phase (Charbin et al., 2014; 64 Lucas et al., 2001), the maintenance of some catenated regions, mostly in regions with 65 repetitive sequences, appears to be important for proper chromosome condensation and sister chromatid cohesion (Bauer et al., 2012; Daniloski et al., 2019). Loss or gain 66 67 of topo II function has been shown to disturb the precisely balanced catenation state of the genome and lead to detrimental effects on chromosome condensation (Cuvier and 68 69 Hirano, 2003; Samejima et al., 2012; Uemura et al., 1987). In vertebrates, the cell cycle-70 dependent expression of topo II $\alpha$ , one of two topo II isoforms found in such organisms, provides one means of calibrating levels of topoisomerase activity (Heck et al., 1988; 71 72 Kimura et al., 1994; Woessner et al., 1991); however, invertebrates and lower-order 73 eukaryotes (such as yeast and Drosophila) express only a single isoform of topo II for 74 which cell cycle-dependent expression has not been observed (Eser et al., 2011; Goto 75 and Wang, 1984; Spellman et al., 1998; Whalen et al., 1991). Post-translational 76 modifications such as SUMOvlation ubiquitylation, and phosphorylation – the majority of

which map to an unconserved, species-specific C-terminal domain of topo II – have
been reported to regulate enzyme stability, localization and activity, but how these
marks exert such functions is poorly understood (Lee and Berger, 2019). Collectively,
these and other observations demonstrate that there is requirement for precise
regulation of topo II activity. However, whether there exist fundamental regulatory
mechanisms that are conserved throughout eukaryotes has not been established.

Because topo II is essential for cell proliferation, it is a potent target for many 84 chemotherapeutics (Nitiss, 2009). A special class of topo II-targeting drugs, categorized 85 86 as topoisomerase 'poisons,' triggers cell death by inducing topo II to generate cytotoxic 87 DNA damage (Pommier et al., 2010; Wu et al., 2011). The major limitation of 88 topoisomerase poisons is their potential to generate DNA damage in noncancerous 89 cells that can lead to serious toxic side effects and therapy-related neoplasias (Felix, 90 1998; Turcotte et al., 2018). A greater understanding of the cellular mechanisms that 91 diminish or enhance topo II activity levels is necessary to more optimally match 92 topoisomerase-targeted therapies to specific cancer types and to design combinatorial 93 strategies that might selectively enhance the potency of anti-topoisomerase drugs in 94 cancer cells.

95

96 In a prior study, we found that a highly-conserved pocket in the ATPase domain of topo II that binds to ICRF-187, a clinically-approved drug, also associates with a 97 98 natural, plant-derived antagonist of topo II known as resveratrol (Lee et al., 2017). This 99 observation led us to wonder whether other small molecules found natively in cells 100 might also modulate topo II function. To test this idea, we prepared crude metabolite 101 extracts from S. cerevisiae and analyzed them for activity against topo II in vitro. We 102 discovered that these extracts were capable of stimulating topo II activity, an effect that 103 has not been observed previously for other agents that act on topo II. Using biochemical 104 fractionation and LC-MS/MS analysis, we determined that di- and tri- carboxylate TCA 105 cycle metabolites were responsible for the stimulatory effect, and that these compounds 106 increase the efficiency of the topo II strand passage reaction. By monitoring the 107 sensitivity of yeast to different classes of topo II inhibitors under different metabolic

108 states, we further found that TCA cycle flux can directly influence topo II activity in vivo. 109 Collectively, our results show that natural small molecules produced by metabolic processes of the cell can directly modulate topo II function and that manipulation of 110 111 these processes affects the efficacy of clinically-approved topo II-targeting drugs. These 112 findings in turn reveal an unanticipated link between cellular metabolism and the regulation of a key central dogma process, a discovery that provides new directions for 113 114 improving the safety and efficacy of commonly used anti-topo II chemotherapies. 115 116 117 RESULTS 118 Small-molecule metabolites from *S. cerevisiae* stimulate topoisomerase II activity 119 in vitro 120 To search for natural products that might regulate topo II, we first tested whether 121 122 crude S. cerevisiae metabolite extracts could impact enzyme activity in vitro. Metabolic 123 extracts were prepared from yeast cultures grown in minimal media without ammonium 124 sulfate, as sulfate ions were found to enhance topo II supercoil relaxation activity 125 (Figure S1A) (all other media components were confirmed to have no significant effect

on topo II activity (Figure S1B)). Small-molecule extracts were taken from log-phase
 cultures to capture metabolic profiles representative of actively dividing cells in all cell
 cycle stages (Figure 1A). A sample of the leftover spent media was also lyophilized as
 a control for media components and any secreted compounds.

130

131 We next assessed the effects of the crude metabolite extract and the spent 132 media control on topo II using decatenation assays. Purified S. cerevisiae topo II 133 (ScTop2) was incubated with kinetoplast DNA (kDNA) – a highly interlocked network of 134 small circular DNAs present in certain protozoa – and ATP to catalyze strand passage. 135 All experiments were conducted with excess ATP (5 mM) to eliminate any response that 136 might have been elicited from the presence of nucleotide in the extracts. Although the 137 spent media sample had little to no effect on DNA unlinking by ScTop2, the crude 138 metabolite extract stimulated decatenation activity in a dose-dependent manner

(Figures 1B-D). Similar to results obtained from decatenation assays, the crude extract
 also stimulated DNA supercoil relaxation by *Sc*Top2, as evidenced by the conversion of
 a negatively supercoiled plasmid substrate into a distribution of relaxed topoisomers

- 142 (Figures 1E-G).
- 143

144 Biochemical characterization of active compounds from yeast metabolite extracts 145 Upon discovering a stimulatory activity in crude extracts, we set out to define the chemical properties of the active agent(s). Liquid-liquid extraction with butanol was first 146 147 conducted under both acidic and basic conditions. In both cases, the stimulatory activity 148 was recovered in the aqueous phase, indicating that the agent was highly polar 149 (Figures S2A and S2B). Based on this finding and the nucleotide-binding capabilities of 150 topo II, we surmised that the factor might contain phosphate groups or phosphodiester 151 bonds. However, the stimulatory activity was resistant to treatment with Antarctic 152 phosphatase or snake venom phosphodiesterase, suggesting that neither phosphates 153 nor phosphodiester bonds were present in the stimulatory metabolite (Figures S2C and 154 **S2D**).

155

156 To further characterize the active compound and prepare samples for mass 157 spectrometry analysis, we developed a purification protocol for the agent (Figure 2A). 158 Solid-phase extraction (SPE) was used as a first-pass, bulk-fractionation method to 159 remove contaminants and enrich the stimulatory activity roughly two-fold (Figure 2B). 160 The active fraction from the SPE step was then further fractionated using reverse-phase 161 HPLC, followed by normal phase HPLC (Figures 2C-F). Interestingly, this procedure 162 not only enriched the stimulatory agent, but also revealed two separate fractions (the 163 gray and pink fractions in Figures 2D and 2F) that inhibited DNA supercoil relaxation by 164 ScTop2 (Figure 2F): these latter activities appear to have been masked by the 165 stimulatory activity in previous purification steps. The appearance of these distinct 166 activities indicates that cells contain more than one class of small molecules that can 167 act on topo II.

168

#### 169 Identification of topo II-stimulating metabolites

170 To identify the stimulatory agent, the stimulatory fraction (green in Figures 2D 171 and 2F) and flanking fractions (pink and blue in Figures 2D and 2F) from the last HPLC 172 purification step were analyzed by LC-MS/MS. Spectra were first collected from a 173 pooled sample to generate a peak-identification library that was representative of all 174 metabolites present in the three samples. Data were processed in Compound Discoverer<sup>TM</sup> by referencing mass spectral databases of natural products (Figure 3A). 175 176 Next, the samples were analyzed individually and spectra from each sample were 177 indexed based on the pooled library. By comparing peak intensities across samples, we 178 generated a candidate list of compounds that were enriched in the stimulatory fraction 179 as compared to its neighboring fractions (Figure 3B, Table S1).

180

Thirty-two candidate compounds were initially tested in eight different pools for 181 stimulatory activity in supercoil relaxation assays. Although most pools showed no effect 182 183 on topo II, some pools did impact enzymatic activity (Figure 3C); compounds from pools displaying activity were then tested individually. Several compounds from the 184 initial candidate list showed an ability to stimulate DNA supercoil relaxation by ScTop2. 185 186 To better understand the chemical properties necessary for this stimulation, we compared the chemical structures of active compounds to those of inactive compounds 187 188 with similar features (Figure 3D). These structure-activity relationship (SAR) analyses 189 revealed that short-chain dicarboxylic acids were able to stimulate topo II activity. 190 Compounds with an additional hydroxyl group on the methylene chain maintained 191 activity, but an amine or a methyl group in the same position ablated the stimulatory 192 effect. Tricarboxylate compounds were also able to stimulate topo II activity, whereas 193 extension of the connecting linker beyond three carbons ablated compound activity 194 (Figure 3D). Together, these tolerable and intolerable chemical changes outlined the 195 specificity of the interaction between the stimulatory compound and topo II. Based on 196 these observations, we identified two chemical motifs – succinic and glutaric acids – 197 that confer stimulatory activity and conducted a substructure search of these motifs against the Human Metabolome Database. This analysis revealed that the motifs are 198 199 most strongly represented in TCA cycle intermediates (Figure 3E and Table S2). Four

TCA cycle intermediates (citrate, isocitrate, succinate, and malate) were identified as stimulatory compounds from our original candidate list (**Table S1**).

202

## 203 TCA cycle intermediates stimulate topo II activity in vitro

204 Based on our observations, we predicted that other TCA cycle intermediates 205 would have similar effects on topo II activity. All TCA cycle intermediates, with the 206 exception of succinyl-CoA, have succinic and/or glutaric acid substructures (Figure 3E). 207 Having ascertained that such a moiety is present in compounds capable of stimulating 208 topo II activity, we proceeded to evaluate the effects of all TCA cycle intermediates on 209 topo II. Consistent with our previous observations, the addition of these intermediates 210 led to as much as a 4- to 12-fold stimulation of decatenation activity and to a 3- to 5-fold 211 stimulation of supercoil relaxation activity at compound concentrations ranging from 25-212 40 mM (Figures 4A-J and S3). Glutamate, an inactive compound (Figure 3D), was 213 added as a non-TCA metabolite control to further validate the specificity of the 214 stimulatory effect.

215

Of the seven TCA metabolites tested, citrate and isocitrate are distinguished by 216 the presence of a third carboxylic acid group. Citrate is a known Mg<sup>2+</sup> ion chelator 217 (Yamagami et al., 2018) and Mg<sup>2+</sup> ions are required for topo II catalytic activity 218 (Osheroff, 1987). To evaluate the potential role of Mg<sup>2+</sup>-ion chelation in the observed 219 220 stimulation of topo II, we assessed the ability of TCA metabolites to stimulate 221 decatenation activity at varying concentrations of Mg(OAc)<sub>2</sub> (Figure S4). Increasing the concentration of  $Ma^{2+}$  alone led to a small but detectable increase in decatenation. 222 223 However, the addition of 30mM citrate or isocitrate significantly stimulated decatenation beyond the effect of adding Mg<sup>2+</sup> alone (Figure S4). These data indicate that the 224 chelation of free Mg<sup>2+</sup> ions by citrate or isocitrate does not appreciably contribute to the 225 226 stimulation of topo II strand passage activity.

227

We next biochemically characterized the ability of TCA metabolites to modulate the ATPase activity of *Sc*Top2 (**Figures 4K, 4L and S5**). Oxaloacetate was excluded from the ATPase activity experiment because as a substrate for lactate dehydrogenase,

it interferes with the coupled reaction. For the evaluated metabolites, we observed that,

with the exception of citrate, all led to only a very slight increase in the maximum rate of

- ATP hydrolysis (< 40%) as compared to the glutamate control. The dicarboxylate
- 234 intermediates had no significant effect on the K<sub>m</sub> of enzyme affinity, whereas an
- increase in  $K_m$  (~2-3 fold) was seen with tricarboxylate metabolites (a response that
- may relate to the chelating activity of these compounds, as Mg<sup>2+</sup> is a co-factor for the
- 237 topo II ATPase reaction). Overall, our data indicate that TCA metabolites have relatively
- little effect on ATP turnover by topo II as compared to the degree of stimulation they
- exert on the enzyme's strand passage activity.
- 240

# The interaction between topo II and TCA cycle metabolites is specific to

## 242 eukaryotic type II topoisomerases

243 Having established a stimulatory interaction between TCA intermediates and 244 budding yeast topo II, we next assessed whether this effect was maintained in human 245 homologs. We tested citrate as a representative tricarboxylic acid metabolite and 246 succinate as representative dicarboxylic acid metabolite against human topo 247 II $\alpha$  (*Hs*Top2A) and topo II $\beta$  (*Hs*Top2B). Both compounds stimulated DNA decatenation 248 by the human topo IIs (Figures 5A-H). HsTop2B decatenation activity was increased by 249 ~10- to 15-fold by both compounds (Figures 5A-D). Succinate had a similar effect on 250 HsTop2A, whereas stimulation of HsTop2A by citrate was biphasic (Figures 5E-H), 251 reaching a peak level of decatenation activity at 30mM citrate that was enhanced ~20-252 fold compared to the lowest amount of metabolite tested (5 mM) (Figure 5F). Because 253 the TCA cycle is present in prokaryotes as well as eukaryotes, we also tested the 254 activity of citrate and succinate against *E. coli* topo IV (Figures 5I-L). Interestingly, both 255 compounds behaved like glutamate and had little to no effect on kDNA decatenation by 256 topo IV. These findings demonstrate that the action of TCA metabolites is specific for 257 eukaryotic type II topoisomerases and evolutionarily conserved from yeast to humans. 258

259 Altering TCA cycle flux leads to changes in levels of topo II activity in vivo

Having established that TCA cycle metabolites exert a stimulatory biochemical effect on topo II, we proceeded to explore the physiological relevance of this activity.

Our *in vitro* data predicted that an increase or decrease in the abundance of TCA cycle intermediates should lead to commensurate changes in topo II activity in the cell. To test this hypothesis, we devised two approaches to modulate TCA cycle flux and assess topo II function *in vivo* using drug sensitivity. Experiments were conducted with a drugefflux deficient (ED) strain of *S. cerevisiae* to improve the dynamic range of the growth assays (Stepanov et al., 2008).

268

269 We first assessed the effects of lowered TCA metabolite abundance by deleting 270 *mpc1*, a subunit of the mitochondrial pyruvate carrier (MPC), in the ED background. 271 Transport of pyruvate into the mitochondrial matrix is necessary to replenish TCA cycle 272 intermediates; loss of mpc1 leads to a significant decrease in the concentration of TCA 273 intermediates (Herzig et al., 2012; Morita et al., 2019). Given our biochemical data, we 274 hypothesized that topo II activity levels should be lower in the  $mpc1\Delta$  strain as 275 compared to the wildtype *MPC1* strain on account of the diminished metabolite levels 276 (Figure 6A). To test this idea, we evaluated the sensitivity of the two strains to chemical 277 agents that antagonize topo II. Type II topoisomerase antagonists are broadly classified 278 as either poisons or general catalytic inhibitors (Nitiss, 2009). Poisons stabilize the 279 cleavage complex of topo II bound to substrate DNA, leading the enzyme to form 280 persistent protein-DNA adducts and DNA-strand breaks. As a result, cells that 281 overexpress topo II are hypersensitive to poisoning agents (Nitiss et al., 1992). In 282 contrast, general catalytic inhibitors reduce enzyme activity without directly stimulating 283 DNA breakage. As a consequence of these different modes of action, changes in topo II 284 activity will have opposing effects on the sensitivity of cells to poisons or catalytic 285 inhibitors: increasing topo II activity is expected to lead to hypersensitivity to poisons 286 (due to the elevated formation of toxic cleavage complexes) and resistance to catalytic 287 inhibitors (because stimulation of enzyme activity overcomes general inhibition), 288 whereas decreasing topo II activity should display the converse (Figure 6A). 289 Consequently, changes in drug cytotoxicity indicate changes in topo II activity. For our 290 experiments, we chose etoposide as a topo II poison and ICRF-187 as a catalytic 291 inhibitor (Ross et al., 1984; Tanabe et al., 1991). The effects of these drugs on the 292 growth of the  $mpc1\Delta$  strain and the MPC1 strain were observed by measuring optical

density over the course of 35 hours. Interestingly, the *mpc1*∆ strain proved resistant to
etoposide and sensitized to ICRF-187 as compared to the control strain, consistent with
the idea that decreasing TCA intermediate abundance leads to a corresponding
decrease in topo II activity (Figures 6B, 6C, S6A, and S6B).

297

298 To test the effect of increasing TCA metabolite abundance, we developed a 299 strategy to enhance TCA cycle flux by adding non-fermentable carbon sources to the 300 growth media. As described by the Crabtree effect (Crabtree, 1928), yeast will opt to 301 generate ATP through glycolysis rather than oxidative phosphorylation under high 302 glucose conditions, regardless of oxygen availability. However, when fermentable 303 carbon sources are limited, yeast will generate ATP by oxidative phosphorylation 304 (OXPHOS), which is coupled to the TCA cycle. To induce different metabolic states, we 305 cultured yeast in minimal media either with glucose only, or with glucose supplemented 306 with lactate and glycerol (LG). Lactate and glycerol are non-fermentable and must be 307 metabolized through the TCA cycle to generate ATP. By LC-MS analysis, we were able 308 to qualitatively compare the levels of five TCA metabolites in yeast grown glucose 309 media with and without LG. We detected significant increases in succinate and malate 310 in the presence of LG and no significant changes in the other metabolites (Figure S6C). 311 To further check that the addition of LG increases TCA flux, we compared the effect of 312 UK-5099, a chemical inhibitor of the MPC that inhibits pyruvate entry into the TCA cycle, 313 on colony growth in the two different nutrient environments. Yeast cells proved more 314 resistant to UK-5099 in media containing LG as compared to the glucose-only condition, 315 confirming that the addition of non-fermentable carbon sources increases TCA flux to 316 counteract the effects of UK-5099 (Figure S6D).

317

We initially examined the effects of adding LG on the sensitivity of yeast cells to anti-topo II agents using a spot-growth assay. The addition of LG sensitized yeast to etoposide as expected (**Figure S6E**) but we were unable to reach cytotoxic concentrations of ICRF-187 necessary to see rescue. We therefore designed a liquid media growth assay that incorporates a diauxic shift to assess the impact of TCA metabolism on cellular sensitivity to topo II antagonists (**Figure 6D**). Starter cultures

324 were first grown to saturation in media containing either glucose alone, or in LG alone to 325 allow the yeast to adjust to either a glycolysis-dominant or an OXPHOS-dominant 326 expression profile. These cells were then shifted into glucose-containing media with and 327 without LG to monitor growth in the presence of anti-topo II drugs. Because glucose is 328 the preferred carbon source, yeast from the glucose-only starter condition will primarily 329 rely on glycolysis and show only a slight increase in TCA metabolism in the presence of 330 LG (Figure 6D). By contrast, yeast grown with only non-fermentable carbon sources will 331 undergo a diauxic shift as glucose is introduced. If LG are present, this diauxic shift will 332 be delayed and the concentration of TCA intermediates will be elevated as the cells 333 attune to metabolizing non-glycolytic carbon substrates (Figure 6D). In agreement with 334 the spot test, the addition of LG sensitized cells to etoposide (Figures 6E, 6F, S6F, and S6G). The addition of LG did not rescue cells that were acclimated to glucose only 335 336 nutrient conditions from ICRF-187 (Figures 6G and S6H). However, the addition of LG 337 did partially rescue yeast from ICRF-187 toxicity when the starter culture was grown 338 without glucose (Figures 6H and S6I). Collectively, these results show that cellular topo 339 Il activity correlates with changes in TCA cycle flux.

340

341 To confirm that the changes in sensitivity to topo II-targeted drugs were due to 342 catalytic stimulation and not a result of differences in protein expression levels, we 343 generated two yeast strains with a 3xHA-tag at either the N-terminal or C-terminal end 344 of endogenous topo II. These strains were then cultured in glucose-only media or 345 glucose media supplemented with LG to compare topo II expression levels under these 346 different growth conditions (Figure S7). Cells were harvested in mid-log phase to 347 capture topo II expression in actively dividing cells and assessed for topo II levels by 348 Western blot. These data show that the change in relative protein abundance across the 349 different growth conditions was insignificant, supporting the idea that the differences in 350 topo II activity observed from the drug screen derive from biochemical stimulation of the 351 enzyme and not due to alterations in protein concentration.

- 352
- 353
- 354 **Discussion**

#### 355 **Regulatory interactions between endogenous metabolites and DNA machinery**

356 In the present work, we show that DNA topoisomerase II (topo II) is subject to 357 regulatory control by small-molecule intermediates of the TCA cycle. Short-chain (2-358 and 3-carbon spaced) di- and tricarboxylic acids are found to stimulate both DNA 359 supercoil removal and DNA decatenation by topo II *in vitro* (Figure 4). Interestingly, 360 TCA cycle intermediates stimulate strand passage without increasing ATP turnover rate 361 (Figures 4 and S5), indicating that these agents increase enzyme efficiency. Human 362 topo IIs are also stimulated by TCA intermediates, whereas homologous bacterial type II 363 topoisomerases are not, suggesting that topo II evolved a response to metabolic control 364 to meet a specific regulatory need in eukaryotes (Figure 5).

365

366 TCA cycle intermediates are some of the most abundant metabolites in the cell, 367 with some intermediates such as citrate, succinate, and malate each reaching 368 concentrations in the 1-5 mM range (Park et al., 2016; Wittmann et al., 2005). Given 369 that seven out of eight TCA cycle intermediates stimulate to topo II activity, a global shift 370 in the collective concentration of this metabolite pool would be expected to exert a 371 functional effect. In vitro, supercoil relaxation was stimulated ~5-fold and DNA 372 decatenation ~12-fold stimulation at the higher metabolite concentrations tested ( $\geq 20$ 373 mM) (Figure 4). However, up to 3-fold increases in activity were still evident even at 374 lower, more physiological levels (between 5-15mM). Many biological systems are 375 maintained at homeostatic setpoints where modest (sub-twofold) biochemical changes 376 can have profound physiological consequences. For example, a < 2-fold increase in  $K_m$ 377 and < 20% decrease in  $k_{cat}$  of the tetracycline resistance protein, TetX2, is sufficient to 378 impart robust resistance to this antibiotic in bacteria (Walkiewicz et al., 2012). Similarly, 379 a ~15% decrease in the expression of mismatch repair protein MSH2 is reported to 380 confer chemoresistance to the chemotherapeutic, temozolomide, in glioblastoma cells 381 (McFaline-Figueroa et al., 2015). Regulatory processes can also be highly sensitive to 382 minor changes in protein activity. A < 10% difference in the concentration of Bicoid, a 383 morphogen, determines cell fate during *D. melanogaster* embryonic development 384 through transcriptional regulation (Driever and Nüsslein-Volhard, 1988; Fradin, 2017). 385 Based on these examples, it is not unreasonable that even a two-fold change in topo II

activity resulting from differing TCA cycle intermediate concentrations would have
physiologic consequences. Consistent with this idea, we show that TCA cycle status
indeed directly impacts topo II function *in vivo*, with changes in TCA flux altering the
relative sensitivity of budding yeast to two different classes of topo II-targeted drugs
(Figure 6).

391

392 Why link topo II activity to metabolic status? One possibility is that it may provide 393 a means to coordinate enzyme function with the cell cycle. Cell cycle-dependent 394 variation in topo II concentration is already known to occur for topo II  $\alpha$  in eukaryotes 395 that express two isoforms of the enzyme (Heck et al., 1988; Kimura et al., 1994; 396 Woessner et al., 1991). Too little or too much topo II activity throughout the cell cycle 397 can impair chromosome condensation and sister chromatid segregation, potentially 398 leading to cell-cycle arrest (Andrews et al., 2006; Cuvier and Hirano, 2003; Downes et 399 al., 1994; Giménez-Abián et al., 1995; Samejima et al., 2012; Uemura et al., 1987). 400 Metabolic sensing by topo II may provide a means for fine-tuning activity, as metabolism 401 is highly responsive to environmental changes (Brauer et al., 2008; Lee and Finkel, 2013; Wellen and Thompson, 2010; Zhu and Thompson, 2019). 402

403

404 The cell cycle is a controlled sequence of events resulting in the duplication of 405 cellular biomass and faithful segregation of genetic material (Elliott and McLaughlin, 406 1978; Goranov et al., 2009; Johnston et al., 1977; Jorgensen et al., 2002; Kalucka et al., 407 2015). This process is not only energetically demanding but also requires the activation 408 of anabolic pathways to generate macromolecular building blocks necessary to 409 accommodate increased transcription, translation, and DNA replication. As such, 410 metabolism has been observed to fluctuate in accordance with cell-cycle progression, 411 and cell-cycle regulators such as cyclin/CDK complexes and the APC-C/Cdh1 have 412 been shown to coordinate metabolic shifts (Almeida et al., 2010; Buchakjian and Kornbluth, 2010; Colombo et al., 2010; Liu et al., 2020; Salazar-Roa and Malumbres, 413 414 2017; Tudzarova et al., 2011; Wang et al., 2014). Proliferating cells strongly up-regulate 415 glycolysis in G1 to rapidly generate ATP and carbon substrates for anabolic pathways 416 (Buchakjian and Kornbluth, 2010; Diaz-Moralli et al., 2013; Pavlova and Thompson,

417 2016), but as cells enter S phase, glycolysis is down-regulated and metabolism shifts 418 towards the pentose phosphate pathway to support DNA replication (Diaz-Moralli et al., 419 2013; Da Veiga Moreira et al., 2015). As transcription ramps down upon chromosome 420 condensation (Johnson and Holland, 1965; Littau et al., 1964; Prescott and Bender, 421 1962; Taylor, 1960), the cell shifts towards the TCA cycle and OXPHOS as a more 422 efficient source of ATP to fuel mitotic progression (Wang et al., 2014). The shift in 423 energy metabolism from glycolysis towards TCA cycle/OXPHOS at mitotic entry mirrors 424 the upregulated expression pattern of vertebrate topo II $\alpha$  that occurs at the same cell 425 cycle transition point. Based on these observations we hypothesize that TCA cycle 426 fluctuations coordinate an increase of topo II activity at the G2-to-mitosis transition. 427 likely to help ensure that chromosomes are fully decatenated. This control mechanism 428 may be particularly important in lower-order eukaryotes that do not display cell cycle-429 dependent expression of topo II and could have been preserved in vertebrates that are capable of upregulating topo II $\alpha$  expression as means of contending with the large 430 431 chromosomal content.

432

#### 433 **Cancer specific metabolic profiles may impact efficacy of topo II-targeted**

#### 434 chemotherapeutics

435 Cancer cells have distinguishing metabolic features, one of the first of which was observed by Otto Warburg (Warburg, 1925). The Warburg Effect, characterized by 436 437 increased aerobic glycolysis and lactic acid fermentation in cancer cells, was initially 438 thought to indicate a shift away from TCA cycle metabolism and OXPHOS to favor the rapid production of ATP through glycolysis; however, growing evidence shows that 439 440 increased rates of glycolysis serve to provide anabolic intermediates for biosynthetic 441 pathways, and that the TCA cycle and OXPHOS both remain active in cancer cells 442 (Locasale et al., 2011; Pavlova and Thompson, 2016). Diverting pyruvate away from the 443 TCA cycle through lactic acid fermentation replenishes NAD+ to maintain high rates of 444 glycolysis (Luengo et al., 2020; Pavlova and Thompson, 2016). TCA cycle metabolites 445 are key anabolic precursors and citrate, in particular, is critical to support elevated levels 446 of lipid biosynthesis in cancer cells (Li and Cheng, 2014; Menendez and Lupu, 2007). Recent studies have also found that the impairment of TCA cycle metabolism and 447

OXPHOS can decrease tumorigenic and metastatic potential (Cai et al., 2020; Cavalli et
al., 1997; Lebleu et al., 2014; Morais et al., 1994; Tan et al., 2015). These findings
demonstrate that cancer cells not only maintain active TCA cycle metabolism but are
dependent on it.

452

453 Different cancer cell types have been shown to have distinct metabolic 454 dependencies that may influence their response to various treatment modalities (Diaz-455 Ruiz et al., 2011; Guppy et al., 2002; Martin et al., 1998; Pasdois et al., 2003). With 456 respect to the TCA cycle, mutations in several key TCA cycle enzymes have been observed in specific cancers. Isocitrate dehydrogenase (IDH1/2) mutations are 457 458 frequently found in gliomas, cholangiocarcinomas, and a subset of acute myeloid leukemias (Balss et al., 2008; Borger et al., 2012; Yan et al., 2009; Yang et al., 2012). In 459 460 such cases, a single amino acid substitution at Arg132 causes IDH to generate 2-461 hydroxyglutarate (2HG) rather than its normal product,  $\alpha$ -ketoglutarate (Dang et al., 2009; Ward et al., 2010; Yang et al., 2012). Interestingly, 2HG was identified as a topo 462 463 II-stimulating metabolite in our studies (Table S1), suggesting that cancers that 464 accumulate 2HG may have elevated levels of topo II activity. Inactivating mutations in succinate dehydrogenase (SDH) and fumarate hydratase (FH) have also been shown to 465 466 cause an accumulation of succinate and fumarate in paragangliomas and 467 pheochromocytomas, as well as in some gastrointestinal and renal cell cancers (Astuti 468 et al., 2001; Baysal et al., 2000; Janeway et al., 2011; King et al., 2006; Letouzé et al., 469 2013; Selak et al., 2005; Tomlinson et al., 2002). The accumulation of oncometabolites 470 (e.g., 2HG, succinate, and fumarate) are thought to affect cell fate determination 471 pathways by disrupting chromatin modification status and DNA repair processes (Baksh 472 and Finley, 2020; Intlekofer and Finley, 2019; Sulkowski et al., 2020; Xiao et al., 2012). 473 Our demonstration that elevated levels of TCA intermediates sensitize yeast cells to 474 topo poisons (Figure 7) suggests that some topo II-targeted therapies may be more 475 effective and selective against cancers with that accumulate oncometabolites. Future 476 efforts will be needed to test this supposition.

- 477
- 478

#### 479 Concluding remarks

480 The present work reports on the ability of native small-molecule metabolites to 481 directly activate eukaryotic topo II in vitro and in vivo. Although there exist many different classes of topo II antagonists, this is to our knowledge the first report of a small 482 483 molecule capable of stimulating this enzyme. Polyamines, such as spermine and 484 spermidine, have been shown to enhance topo II supercoil relaxation activity by 485 condensing DNA but these compounds do not directly bind topo II (Pommier et al., 1989; Srivenugopal et al., 1987). Because polyamines act on the DNA substrate and 486 487 not the enzyme, they also affect the activity of type I topoisomerases and bacterial type 488 II topoisomerases (Srivenugopal and Morris, 1985; Srivenugopal et al., 1987). By 489 contrast, the specificity of TCA metabolites for stimulating eukaryotic topo II indicates 490 that the enhancement of strand passage activity is due to a direct interaction between 491 the enzyme and the metabolites (Figure 5). These findings also represent the first 492 example of an allosteric interaction between a non-nucleotide metabolite and an 493 enzyme involved in controlling a 'central dogma' process. Small molecule 'alarmones,' 494 such as (p)ppGpp and cGAMP, do exist that can regulate translation, transcription, and 495 DNA replication (Srivatsan and Wang, 2008; Sun et al., 2013); however, these agents 496 are nucleotide analogs and are produced in response to stress, as opposed to the 497 normal operations of the cell. Metabolites such as  $\alpha$ -ketoglutarate, succinate, fumarate, 498 and 2HG can act as cofactors or competitive inhibitors for chromatin modifying enzymes 499 (Baksh and Finley, 2020; Intlekofer and Finley, 2019), but prior to this study, none were 500 known the allosterically regulate an enzyme involved in nucleic acid transactions. The 501 discovery of a regulatory topo II-TCA cycle interaction opens up the possibility that other 502 such interactions between nuclear enzymes and endogenous small molecules may 503 exist for the purposes of modulating events such as transcription and/or DNA 504 replication. Our approach for isolating metabolite-enzyme interactions is readily 505 transferable to any system where enzyme activity can be monitored by a biochemical 506 assay. We expect that future applications of the approach may be useful for revealing 507 other regulatory connections between cellular metabolism and DNA-dependent 508 machineries.

509

510 With regard to anti-topo II agents as cancer therapeutics, the off-target poisoning 511 of topo II $\beta$  appears to be at least one major source of negative side effects, including 512 cardiotoxicity and development of secondary malignancies (Azarova et al., 2007: Felix, 513 1998; Turcotte et al., 2018; Yi et al., 2007). Because of the high degree of similarity 514 between the two human isoforms of topo II, designing strategies to increase the 515 specificity of topo II inhibitors is challenging. Our data show that there is a difference in 516 the biochemical response of topo II $\alpha$  and topo II $\beta$  to di- and tricarboxylic acids (Figure 517 5); defining the biochemical underpinnings of these differences may provide new 518 insights toward increasing the specificity of topo II-targeted drugs. To this end, we note 519 that some fractions of our metabolic extracts also showed inhibitory activity against topo 520 II, indicating that there exist other biologically relevant metabolite interactions that have 521 yet to be uncovered. The identification of these agents could serve as a starting point 522 for such investigations.

523

In closing, the relationship between TCA cycle metabolism and topo II activity gives rise to many new questions about how topo II drugs interact with cancerous and noncancerous cells. A deeper understanding of the metabolic regulation of topo II activity may be useful for improving for patient selection strategies and topo II-targeted chemotherapies. Future studies expanding into mammalian systems will clarify the conserved link between cellular metabolism and topo II regulation with the ultimate goal of improving the safety and efficacy of patient-specific cancer treatments.

#### 532 ACKNOWLEDGMENTS

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541

### 542 AUTHOR CONTRIBUTIONS

- 543 JHL conducted *in vitro* assays, fractionated yeast metabolites, and performed yeast cell
- 544 growth assays. EPM conducted LC-MS/MS analysis of metabolite samples. Data
- analysis was performed by JHL and EPM. YSL contributed to the development of the
- 546 yeast metabolite extraction method. JHL, EPM, NNB, and JMB contributed to
- 547 conceptual planning, experimental design, and data interpretation. JHL, EPM, and JMB
- 548 prepared the manuscript. All authors critically reviewed the manuscript and approved
- 549 the final version.
- 550

### 551 **DECLARATION OF INTERESTS**

552 The authors declare no competing interests

553

#### 555 **FIGURE LEGENDS**

Figure 1. Crude metabolite extracts from log-phase yeast stimulate ScTop2 strand 556 557 passage activity. (A) Schematic of metabolite extraction procedure from yeast cultures. 558 (B-C) Representative gels of kinetoplast DNA (kDNA) decatenation assays with 559 metabolite extracts and spent media control samples. Bands representing nicked 560 minicircles and closed minicircles are indicated to the left of the gels. No enzyme (-topo) 561 and no ATP (-ATP) negative controls show the starting substrate. Lyophilized samples 562 of the metabolite extract and spent media were titrated from 0 to 50 mg/ml in 10 mg/ml 563 increments. (D) Decatenation assay data represented as mean  $\pm$  SD of n=3 independent experiments. (E-F) Representative gels of supercoil relaxation assays with 564 565 metabolite extracts and spent media control samples. The bands representing 566 unrelaxed substrate (SC), the relaxed topoisomer distribution, and nicked/open circle 567 (OC) plasmids are indicated to the left of the gels. No enzyme (-topo) and no ATP (-ATP) negative controls show the starting substrate. Lyophilized samples of the 568 569 metabolite extract and spent media were titrated from 0 to 50 mg/ml in 10 mg/ml 570 increments. (G) Supercoil relaxation assay data represented as mean  $\pm$  SD of n=3

571 independent experiments.

572

576

#### 573 **Figure 2. Purification of stimulatory metabolites from crude metabolite extracts.**

574 (A) Schematic of purification steps to prepare samples with enriched stimulatory activity

575 for LC-MS/MS analysis. (B) Supercoil relaxation assay with metabolite samples before

('Crude') and after solid phase extraction ('SPE'). Crude and SPE purified metabolite

577 extracts were added from 0 to 40 mg/ml in two-fold increments (0, 2.5, 5, 10, 20, 40

578 mg/ml). (C-D) Chromatograms of reverse phase (C) and normal phase (D) HPLC

579 purification runs. The HPLC method is depicted by the purple line showing fraction of

580 phase B as indicated by the right y-axis. Red and green traces show absorbance values

- at 260 nm and 280 nm wavelengths respectively as indicated on the left y-axis. (E-F)
- 582 Supercoil relaxation assays of reverse phase fractions (E) and normal phase fractions
- 583 (F) indicated by the red brackets in (C) and (D). Fractions of interest are highlighted by
- 584 corresponding colors in the chromatograms and relaxation assay gels. Lyophilized

material from each fraction was solubilized in equal volumes of 12.5% DMSO and
titrated down from the maximum possible concentration in two-fold dilution steps

588 Figure 3. Identification and structure activity relationship (SAR) analysis of 589 stimulatory compounds (A) Schematic depicting how three sequential fractions 590 (indicated by pink, green, and blue in Figure 2) were pooled and analyzed by LC-591 MS/MS to generate an ion peak identification library. (B) Schematic depicting how 592 different samples (colored as pink, green, and blue) were analyzed individually. 593 Intensities of metabolite peaks were compared across the three samples to determine 594 relative abundance of each ion. Ions that were enriched in the stimulatory fraction 595 (green) were added to a candidate list for stimulatory compounds (see **Table S1**) (C) Example supercoil relaxation assays from testing an inactive and an active pool of five 596 597 candidate compounds. Candidate pools were prepared from commercial compounds. 598 The concentration of each candidate compound was titrated from 0 to 20 mg/ml in two-599 fold steps (0, 0.31, 0.63, 1.25, 2.5, 5, 10, 20 mg/ml). (D) SAR analysis comparing 600 stimulatory and inactive compounds identified from candidate pools. Chemical 601 properties that are necessary for activity (green circles) and chemical changes that ablate activity (blue circles) are highlighted in gray. (E) TCA cycle metabolites. Succinic 602

- acid (tan) and glutaric acid (gold) motifs are highlighted.
- 604

**Figure 4. TCA cycle intermediates stimulate strand passage by topo II without** 

606 significant effects on ATP hydrolysis. (A-H) Stimulation of topo II decatenation

607 activity (A-D) and supercoil relaxation activity (E-H) by citrate and oxaloacetate (OAA).

608 Citrate, OAA, and glutamate (negative control) were titrated from 0 to 40mM in 5mM

increments. Graphs (B, D, F, and H) represent mean ± SD of n=3-5 independent

- experiments. (I-J) Stimulation of ScTop2 decatenation (I) and relaxation (J) activity by
- TCA metabolites and glutamate (negative control). Stimulation values represent mean
- of n=3-6 independent experiments. (See also **Figure S3**). (K-L) Effects of TCA
- 613 metabolites on ATP hydrolysis activity. Experiments were performed at 0, 10, 20, and
- 614 30mM concentrations of the metabolites indicated in the legend to the right of the

615 graphs.  $K_m$  (K) and  $V_{max}$  (µmol ATP • min<sup>-1</sup> • nmol topo II<sup>-1</sup>) (L) values were derived from 616 n=3 independent experiments. (See also **Figure S5**).

617

Figure 5. Stimulatory effect of TCA metabolites is eukaryotic specific. Citrate and
succinate were titrated from 0 to 40 mM in 5 mM increments and added to decatenation
assays with *Hs*Top2B (A-D), *Hs*Top2A (E-H) and *Ec* topo IV (I-L). Graphs represent
mean ± SD of n=3-5 independent experiments.

622

**Figure 6. Changes in TCA cycle flux affect sensitivity of yeast to topo II inhibitors.** 

624 Green indicates conditions in which TCA flux is decreased, while orange indicates

625 conditions in which TCA flux is increased. (A) Predicted correlation between ATP-

626 generating metabolic pathways and topo II activity. Cells generate ATP by glycolysis

627 (green) and oxidative phosphorylation (TCA/OXPHOS, orange). Our model predicts that

as TCA metabolism increases, topo II activity will also increase (black). As topo II

629 activity increases, etoposide toxicity (red) increases and ICRF-187 toxicity (blue)

630 decreases; hence, etoposide toxicity should directly correlate with TCA flux and ICRF-

187 toxicity should inversely correlate with TCA flux. (B-C) Growth curves of *mpc1*⊿

632 (green) compared to MPC1 (orange) in the presence of etoposide (B) and ICRF-187

633 (C). Orange arrows show the effect of increasing TCA flux (downward indicates

634 sensitization and upward indicates rescue). (See also **Figure S6A-B**) (D) Schematic

depiction of the effects of nutrient changes on flux through glycolysis and TCA cycle.

Arrows are colored to show shifts in metabolism over time. Cultures initially grown in

637 glucose conditions (E, G) will continue to have high levels of glycolysis, but the addition

of lactate and glycerol will cause an increase in TCA flux. Cultures initially grown in

639 media with only lactate and glycerol (F, H) will generate ATP solely through

640 TCA/OXPHOS. Introduction of glucose will cause a diauxic shift towards glycolysis, but

if lactate and glycerol are maintained in the media, this shift will be delayed and TCA

642 flux will stay elevated. Letters to the right of (D) indicate the nutrient conditions of the

643 growth curves shown in (E-H). (E-H) Green lines show growth in glucose only media

and orange lines show growth in media with glucose, lactate, and glycerol. Cultures

645 were inoculated from starters grown in glucose only media (E and G) or lactate and

- 646 glycerol only media (F and H). Orange arrows show the effect of increasing TCA flux on
- 647 cytotoxicity of etoposide (E and F) and ICRF-187 (G and H) (downward indicates
- sensitization and upward indicates rescue). (See also **Figure S6F-I**).
- 649
- 650

### **Figure 7. Schematic depicting how sensitivity to topo II targeting drugs is**

652 influenced by metabolic state. When TCA cycle flux is low (green) topo II strand

653 passage activity is not stimulated. General catalytic inhibitors (e.g., ICRF-187) that work

by decreasing topo II activity will be more effective in this state, as compared to a high

TCA cycle flux state (orange). By contrast, topo II poisons (e.g., etoposide) are more

toxic when topo II activity is stimulated in the high TCA cycle flux state because

- 657 elevated topo II strand passage activity leads to increased formation of cytotoxic DNA
- 658 double strand breaks.
- 659

## 660 SUPPLEMENTAL FIGURE LEGENDS

## 661 Figure S1. Effects of synthetic media components on topo II activity. (A)

662 Schematic for acid/base butanol liquid-liquid extraction of crude metabolite extracts. 'Aq' 663 indicates the aqueous fraction, and 'Org' indicates the organic fraction. (B) Supercoil 664 relaxation assay with fractions from acid/base butanol extraction as indicated by colors. 665 Solid material left after solvent removal from each fraction was solubilized in equal volumes of 12.5% DMSO and titrated into relaxation assays in 2-fold dilution steps. (C-666 667 D) Activity of crude metabolites after enzymatic treatment by Antarctic phosphatase (C) 668 or snake venom phosphodiesterase (D). Lyophilized material from treated and 669 untreated samples were solubilized in equal volumes of 12.5% DMSO and titrated into

- 670 relaxation assays in 2-fold dilution steps.
- 671

Figure S2. Chemical properties of stimulatory metabolites. (A) Schematic for
acid/base butanol liquid-liquid extraction of crude metabolite extracts. 'Aq' indicates the
aqueous fraction, and 'Org' indicates the organic fraction. (B) Supercoil relaxation assay
with fractions from acid/base butanol extraction as indicated by colors. Solid material left
after solvent removal from each fraction was solubilized in equal volumes of 12.5%

DMSO and titrated into relaxation assays in 2-fold dilution steps. (C-D) Activity of crude
metabolites after enzymatic treatment by Antarctic phosphatase (C) or snake venom
phosphodiesterase (D). Lyophilized material from treated and untreated samples were
solubilized in equal volumes of 12.5% DMSO and titrated into relaxation assays in 2-fold
dilution steps.

- 682
- 683 Figure S3. Stimulation of ScTop2 supercoil relaxation activity (A-G) and
- 684 decatenation activity (H-N) by TCA cycle intermediates (blue) and glutamate
- 685 (negative control, orange). Graphs show mean ± SD of n=3-6 independent
- 686 experiments.
- 687

```
Figure S4. Chelation of Mg<sup>2+</sup> ions by tricarboxylate TCA cycle intermediates does
```

- 689 **not affect stimulatory topo II-metabolite interaction.** (A) Representative gels of
- 690 decatenation assays performed at different concentrations of Mg(OAc)<sub>2</sub> with no added
- 691 metabolite, 30mM citrate, and 30mM isocitrate. Mg(OAc)<sub>2</sub> was titrated from 10 to 50 mM
- in 10 mM steps. (B) Graphs show mean  $\pm$  SD of n=3 independent experiments to
- 693 compare the fraction decatenated under each reaction condition.
- 694

**Figure S5. Michaelis-Menten curves of ATP hydrolysis by ScTop2 in the presence** 

696 of TCA cycle intermediates and glutamate (negative control). Rates of ATP

- 697 hydrolysis were measured in a coupled assay by monitoring NADH consumption as a
- 698 function of light absorbance at 340 nm wavelength (y-axis). Graphs represent mean ±
- 699 SD of n=3 independent experiments.
- 700

## 701 Figure S6. Changes in TCA cycle flux affect sensitivity of yeast to topo II

- inhibitors extended data. (A-B) Growth curves of MPC1 (orange) and  $mpc1\Delta$  (green)
- with etoposide (A) and ICRF-187 (B). (C) Changes in TCA cycle metabolite abundance
- vupon addition of lactate and glycerol to media. Peak areas were normalized to the
- average of the control condition (glucose only). Bar graphs and error bars indicate mean
- $\pm$  SD of n=3. An unpaired, two-tailed t-test was used to analyze significance of changes
- peak area. One asterisk (\*) indicates p < 0.05, and two asterisks (\*\*) indicates p < 0.01.

708 (D-E) Serially diluted cultures of yeast were spotted on glucose containing agar plates 709 with and without lactate and glycerol. Spot growth at 0-25 µM UK-5099 (D) and 300 µM 710 etoposide (E) was observed after 2-3 days. (F-H) Growth curves of yeast in glucose-711 only media (green) or glucose media supplemented with lactate and glycerol (orange). 712 Cultures were inoculated from starters grown in glucose only media (F and H) or lactate 713 and glycerol only media (G and H). Orange arrows show the effect of increasing TCA 714 flux on cytotoxicity of etoposide (F and G) and ICRF-187 (H and I) (downward indicates 715 sensitization and upward indicates rescue).

716

## 717 Figure S7. ScTop2 expression levels in different nutrient conditions (A)

718 Representative western blot of 3xHA-tagged endogenous topo II in an N-terminally

tagged (N-tag) or a C-terminally tagged (C-tag) topo II strain from cells grown in media

with and without lactate and glycerol (LG). The wildtype (WT) yeast strain (untagged

topo II) is shown as a negative control. (B) Quantification of topo II expression levels in

the N-tag and C-tag strains. Intensities of the topo II bands were normalized to the

723 corresponding tubulin band. Relative amounts of topo II were calculated by dividing the

normalized topo II intensity in the +L/G condition to that of the - L/G condition. Data are represented as mean  $\pm$  SD of n=3 independent experiments.

726

Table S1. Relative abundance and activities of metabolites identified in LC-MS/MS
 analysis of inactive, stimulatory, and inhibitory fractions of yeast metabolite
 extracts.

730

731 Table S2. Results of substructure search of succinic and glutaric acid motifs on

- 732 the Human Metabolome Database
- 733

734	STAR METHODS
735	
736	Resource availability
737	
738	Lead Contact
739	Further information and requests for resources and reagents should be directed to and
740	will be fulfilled by the Lead Contact, James M. Berger (jmberger@jhmi.edu).
741	
742	Materials Availability
743	All reagents generated for this study are available on request through the Lead Contact.
744	
745	Data and Code Availability
746	The data obtained in this study will be accessible at the NIH Common Fund's NMDR
747	(supported by NIH grant, U01-DK097430) website, the Metabolomics
748	Workbench, https://www.metabolomicsworkbench.org
749	
750	Experimental Model and Subject Details
751	
752	Yeast strains and growth conditions
753	Crude metabolite samples were extracted from cultures of BY4741 yeast (MAT $lpha$
754	<i>his3<math>\Delta</math>1 leu2<math>\Delta</math> met15<math>\Delta</math>0 ura3<math>\Delta</math>0).</i> Growth experiments to assess the sensitivity of yeast to
755	topo II-targeted drugs were performed with a drug-efflux deficient (ED) strain
756	(pdr1_]: pdr1: Cyc8 LEU2) described in (Stepanov et al., 2008). To generate the ED-
757	<i>mpc1</i> $\Delta$ strain, the KANMX resistance marker was PCR amplified from the pUG6
758	plasmid (Güldener et al., 1996) with the following primers containing homology regions
759	outside of the MPC1 gene:
760	(5')CAGCAAACGTCAATACATCTACATATATACGTATAGATTTTATTGCACTGTGATC
761	GACATGGAGGCCCAGAATACC and
762	(5')GTTTCCATCTAGTCACCTACTTCAGGTTCTTAGACTGCTCGTTTTACCAGTATAG
763	CGACCAGCATTC. The deletion construct was then transformed into ED yeast. After

recovery on YPD plates followed by selection on G418 containing replica plates,  $mpc1\Delta$ mutants were verified by PCR.

766

767 CRISPR/Cas9 technology was used to incorporate a 3xHA-tag at the N-terminus
 768 and C-terminus of endogenous topo II in the BY4741 background. The following gRNA

- 769 sequences were integrated in the CRISPR/Cas9 vector, pJH2972 (Anand et al., 2017):
- 770 (5')GCAGTGAAAGATAAATGATCGTTGACATGGTTAGCCGTGCGTTTTAGAGCTAGA
- 771 AATAGC (N-terminal gRNA) and
- 772 (5')GCAGTGAAAGATAAATGATCAAAAAGAATGGCGCTTTCTCGTTTTAGAGCTAGA
- AATAGC (C-terminal gRNA). The CRISPR/Cas9 constructs were then transformed into
- 774 BY4741 yeast with their corresponding repair templates:
- 775 (5')TTTCAGTTAAAGGAGTTTATAACGACGAGCACGCCTAACCATGTACCCATACGA
- 776 TGTTCCTGACTATGCGGGCTATCCCTATGACGTCCCGGACTATGCAGGATCCTATC
- 777 CATATGACGTTCCAGATTACGCTGGTACGATGTCAACTGAACCGGTAAGCGCCTCT
- 778 GATAAATATCAGA (N-tag) and
- 780 CGATGTTCCTGACTATGCGGGCTATCCCTATGACGTCCCGGACTATGCAGGATCCT
- 781 ATCCATATGACGTTCCAGATTACGCTTGAATAATATTTATCGAGAGAAAGCGCCATT
- 782 CTTTTTATA (C-tag). Integration of the repair constructs was verified by colony PCR
- followed by BamHI digest of the product (only the PCR product from a strain with
- correct integration of the 3xHA-tag will have a BamHI cut site).
- 785

Cultures for metabolite extraction were grown in synthetic media in which ammonium sulfate was replaced with monosodium glutamate as the nitrogen source (SC-sulfate). Per liter, SC-sulfate media contains 75 mg of each amino acid, 20 mg of adenine, 75 mg of uracil, 25 mg of inositol, 1.71 g of YNB-sulfate (Sunrise Science), 1 g monosodium glutamate, and 20 g dextrose. Growth assays were conducted in SC media (Sunrise Science). Glucose (20 g/L), lactate (2%) and glycerol (1.5%) were added to the media after autoclaving as required for each experiment.

793

794 Method Details

#### 795 S. cerevisiae metabolite extraction

BY4741 cultures grown in SC-sulfate media were harvested at early log phase. 796 797  $OD_{600} = 0.4-0.8$ , by vacuum filtration through a surfactant-free cellulose acetate 798 membrane with a 0.2 µm pore size (ThermoFisher Scientific). When ~5 mL of media 799 was left in the filter, 30 mL of 100 mM NH₄OAc – TEA (pH 8.0) was added to wash the 800 cells and remove any remaining extracellular material. Immediately after the wash 801 solution was passed through the filter, the filter apparatus was released from vacuum 802 and 10 mL of 90% MeOH, 10 mM NH₄OAc – TEA (pH 8.0) (chilled to -80 °C) was 803 added to the filter for rapid quenching and extraction (Boer et al., 2010; Crutchfield et 804 al., 2010). The extract was then centrifuged at 4000g for 10-15 min to remove cell 805 debris. The extract supernatant was passed through a 3 kDa MWCO filter (Cytiva) to 806 remove large macromolecules. The filtrate was diluted 2-fold with water and lyophilized 807 to remove the extraction solvent. The remaining extracts were stored at -80 °C for up to 808 one month.

809

#### 810 Supercoil relaxation, decatenation, and ATPase activity assays

811 Recombinant eukaryotic proteins were prepared as described previously (Lee et al., 2017). Briefly, ScTop2, HsTop2A, and HsTop2B were expressed in the S. cerevisiae 812 813 strain BCY123 (Wasserman and Wang, 1994), and cell pellets were lysed by cryogenic grinding. Proteins were first purified by Ni-affinity purification (HisTrap FF, GE) followed 814 815 by cation exchange (HiTrap SP, GE). Affinity tags were removed by His<sub>6</sub>-tagged TEV 816 protease (QB3 MacroLab). The digested proteins were passed over a second Ni-affinity 817 column to remove the cleaved His<sub>6</sub>-tag and the His<sub>6</sub>-TEV protease. For the final 818 purification step, proteins were run on a gel-filtration column (S-400, GE). Recombinant 819 Ec topolV was prepared as described previously (Vos et al., 2013). His<sub>6</sub>-tagged 820 subunits of Ec topolV (ParE and ParC) were expressed in BL21 codon-plus (DE3) RIL 821 cells and cells were lysed by sonication. Proteins were first purified by Ni-affinity purification (HiTrap Ni<sup>2+</sup>, GE). After removing the affinity tags by TEV cleavage and a 822 823 second Ni-affinity step as described above, the proteins were purified by gel filtration (S-824 300, GE).

Negatively supercoiled pSG483, a derivative of pBluescript SK, was prepared
from XL1-Blue *E. coli* cultures by maxi prep as described previously (Lee et al., 2017).
Metabolite extracts were solubilized in 12.5% DMSO and metabolite stocks prepared
from commercial compounds were solubilized in water and adjusted to pH 6.5-7.5 with
HOAc or KOH.

831

832 For strand passage activity assays, enzyme stocks were first diluted in two-fold 833 steps in 30 mM Tris-HCI (pH 7.9), 500 mM KOAc, 10% glycerol and 0.5 mM TCEP. Ec 834 topo IV subunits were incubated together on ice at high concentrations (200 - 300 µM of 835 each subunit) for 10 min prior to dilution to allow formation of the holoenzyme. The 836 enzyme was then added to the appropriate DNA substrate, negatively supercoiled 837 pSG483 plasmid for supercoil relaxation assays or kDNA (Inspiralis) for decatenation 838 assays, followed by metabolites. Reactions were initiated by the addition of ATP (5 mM 839 for assays with extracts and 1 mM for assays with commercial metabolites) and shifted 840 to 30 °C for ScTop2 or 37 °C for HsTop2A/B and topo IV. The final reaction conditions 841 contained 32 mM Tris-HCI (pH 7.9), 100 mM KOAc, 20 mM Mg(OAc)<sub>2</sub>, 0.05 mg/ml BSA, 842 0.6 mM TCEP, and 10% glycerol. Assays with metabolite extracts also contained 2.5% 843 DMSO. Decatenation reactions were incubated for 3 min and supercoil relaxation 844 assays were incubated for 5 min. Reactions were guenched with 20 mM EDTA and 1% 845 SDS and treated with 0.2mg/mL Proteinase K to remove any remaining protein bound to 846 DNA. Final products were resolved and visualized by native gel electrophoresis (1.4%) 847 agarose, 1X TAE) run at 2-2.5 V/cm for 15-20h. Supercoil relaxation products were run 848 on native gels and decatenation products were run on gels containing 0.4 µg/mL 849 ethidium bromide.

850

ATP hydrolysis by topo II was measured by an NADH-coupled ATPase assay.
Reactions were prepared with 50 nM *Sc*Top2, 200 ng/μL sheared salmon sperm DNA
(Fisher Scientific), 0.5 mM NADH, 0.2% rabbit muscle pyruvate kinase/lactate
dehydrogenase (Sigma- Aldrich), 2 mM PEP, 36 mM Tris-HCI (pH 7.9), 100 mM KOAc,
10 mM Mg(OAc)<sub>2</sub>, 0.05 mg/mL BSA, 0.6 mM TCEP, and 2% glycerol in a final reaction
volume of 75 μL. ATP concentrations were titrated from 0-4 mM. Depletion of NADH

was observed over the course of one hour at 30°C by monitoring absorbance at 340 nm
in a CLARIOStar microplate reader (BMB LAB TECH). ATP hydrolysis rates were
calculated based on an NADH standard curve and fit for K<sub>m</sub> and V<sub>max</sub> values in PRISM
(GraphPad).

861

862 Fractionation and purification of active compounds from crude metabolite extracts

863 Lyophilized samples of metabolite extracts were resuspended with water and 864 adjusted to pH 3.5 with formic acid or pH 10 with TEA. The metabolite solution was 865 transferred to a separatory funnel and mixed with an equal volume of butanol. The 866 mixture was then left until the separation of aqueous and organic layers was complete. 867 The aqueous solvent was removed by lyophilization and the organic solvent was 868 removed in a rotary evaporator. The activities of the aqueous and organic fractions from 869 acid- or base- extraction were assessed in a supercoil relaxation assay to determine the 870 hydrophilic nature of the active compound.

871

Enrichment and purification of the stimulatory metabolite from crude extracts for LC-MS/MS analysis began with solid phase extraction (SPE). Crude extracts were solubilized in water and adjusted to pH 10 with TEA. A mixed-mode anion exchange cartridge was equilibrated as per the manufacturer's instructions (Oasis MAX, Waters). The flow-through from the passage of the crude extract through the SPE column (i.e., unbound material) was lyophilized for further purification.

878

879 The extract was then further purified by reverse-phase HPLC on a C18 column 880 (Atlantis T3, Waters). One column volume of water with 0.1% formic acid followed by 881 eight column volumes of MeOH were passed through the column over the course of the 882 HPLC run. Absorbance at 260 nm and 280 nm wavelengths was observed by a Waters 883 2996 Photodiode array. Fractions from the aqueous phase were lyophilized and 884 assessed for stimulation of supercoil relaxation. Active fractions were then further 885 purified through normal-phase HPLC on an amide column (XBridge BEH Amide, 886 Waters). One column volume of 80% acetonitrile with 0.1% formic acid followed by a 887 three-column volume gradient down to 40% acetonitrile with 0.1% formic acid were

passed through the column. Afterwards, one column volume of water with 0.1% formic
acid was passed over the column as the last step. The separated fractions were

- 890 lyophilized and assessed similarly to all prior purification steps.
- 891
- 892 Antarctic phosphatase and phosphodiesterase treatment of metabolites

893 Snake venom phosphodiesterase I (SVPD, Sigma-Aldrich) was reconstituted at 1 894 mg/mL in 20 mM Tris-HCI (pH 7.9), 100 mM NH<sub>4</sub>OAc, 20 mM Mg(OAc)<sub>2</sub>, 50% glycerol (Konokhova et al., 2016). Crude extracts were first fractionated by SPE as described 895 896 above. The lyophilized sample was then resuspended in water and split into four 897 samples of 44 µL each. Next, 5µl of 10X reaction buffer was added to each sample: 10x 898 Antarctic phosphatase (AP) reaction buffer (New England BioLabs) was added to two 899 samples and 200 mM Tris-HCI (pH 7.9), 1 M NaCl, 200 mM MgCl<sub>2</sub> was added to the 900 remaining two samples for SVPD treatment. For each pair of samples, 1 µl of water was added to the no enzyme control and 1 µl of enzyme (AP or SVPD) was added to the 901 902 second sample. All reactions were incubated at 37 °C for two hours. Samples were then 903 diluted to a 1 mL final volume and passed through a 3 kDa MWCO filter. The filtrate was 904 lyophilized and assessed for activity against topo II in a supercoil relaxation assay. 905

### 906 Untargeted LC-MS/MS analysis of metabolite samples

Lyophilized samples were reconstituted in water plus 0.1% (v/v) formic acid such
that samples reached a final concentration of 20 mg/ml. A pooled sample was
generated by combining equal volume aliquots from each sample. Untargeted
metabolomics uHPLC-MS/MS data acquisition was performed in both positive and
negative ion modes and using both reversed-phase and normal-phase separations.

Liquid chromatographic separation was performed using a Dionex Ultimate 3000
uHPLC system (Thermo) for reversed-phase and normal-phase methods. For both
methods, mobile phase A was 1mM ammonium acetate + 0.1% (v/v) formic acid in
water and mobile phase B was 1mM ammonium acetate + 0.1% (v/v) formic acid in 20%
water / 80% acetonitrile (v/v). Reversed-phase experiments were performed with a
Hypersil GOLD aQ polar endcapped C18 column (150 mm x 2.1 mm, 1.9 µm particle

919 size) (Thermo) using the following gradient of mobile phases: 0% B from 0-2 min, 0-920 60% B from 2-13 min, 60-90% B from 13-15 min, 90-0% B from 15-16 min, and 0% B 921 from 16-20 minutes. Normal-phase experiments were performed with a 922 Polyhydroxyethyl A hydrophilic interaction chromatography column (200 mm x 2.1 mm, 923 5 µm particle size) (PolyLC) using the following gradient of mobile phases: 80% B from 924 0-2 min, 80-40% B from 2-13 min, 40-10% B from 13-15 min, 10-80% B from 15-16 min, 925 and 80% B from 16-20 min. For both uHPLC methods, the flow rate was set to 0.2 926 mL/min. Samples were each injected 3 times under each unique set of conditions. A 927 blank injection of 0.1% formic acid in water (v/v) was run in between each sample 928 injection to minimize carry over. 929

After separation by uHPLC, metabolites were ionized and detected by a Q-930 931 Exactive quadrupole-orbitrap mass spectrometer (Thermo) using a heated electrospray 932 ionization source. Samples were run separately in negative and positive ion modes. In 933 positive mode, ionization was performed with a spray voltage of 4000 V, capillary 934 temperature of 275°C, sheath gas flow rate of 35 Arb, and auxiliary gas flow rate of 8 935 Arb. Negative mode ionization was performed with a spray voltage of 4000 V, capillary 936 temperature of 350°C, sheath gas flow rate of 25 Arb, and auxiliary gas flow rate of 2 937 Arb.

938

For each set of separation and ion modes, mass spectra were acquired by scans in data-dependent MS/MS mode for the pooled samples and full-MS mode for individual fraction replicates. Full-MS scans were performed at a resolution of 140,000 and a mass range of m/z 65-850. Data-dependent MS/MS scans were carried out for the top-5 abundant ions at a resolution of 140,000 FWHM with a dynamic exclusion of 6.0 seconds and stepped normalized collision energy of 20, 40, and 100. MS/MS fragments were observed at a resolution of 17,500 FWHM.

946

947 The LC-MS and LC-MS/MS data were processed using Compound Discoverer
948 3.1 software (ThermoFisher Scientific). Spectral features retention times were aligned
949 with a permitted mass deviation of 5 ppm. Background peaks were subtracted based on

features in a blank sample. The data-dependent MS/MS files were analyzed for
compound identification using mzCloud and the ChemSpider yeast metabolome
database. The full-MS scan data was used for comparison of metabolite abundance by
average peak area. Analysis of metabolite enrichment was performed by comparing
replicates from fractions with an effect on topo II activity to those with no effect on topo
II activity.

956

### 957 Targeted Metabolomics Analysis of TCA Metabolites

958 Lyophilized samples of crude extracts from yeast grown in different nutrient 959 conditions were reconstituted in water + 0.1% formic acid, such that the ratio of 960 milligrams of cell mass to microliters of acidified water was 0.5. Samples were analyzed 961 on a Dionex Ultimate 3000 uHPLC (Thermo) coupled to a TSQ Vantage triple 962 guadrupole mass spectrometer (Thermo). Analytes were separated on a Hypersil GOLD 963 aQ polar endcapped C18 column (150 mm x 2.1 mm, 1.9 µm particle size) using an 964 isocratic flow of 100% mobile phase A (water +0.1% formic acid) at a flow rate of 0.2 965 mL/min over five minutes. Analytes were ionized by heated electrospray ionization with 966 a spray voltage of 3000 V, capillary temperature of 204°C, sheath gas flow rate of 50 967 Arb, and auxiliary gas flow rate of 55 Arb. Metabolites were detected in negative ion 968 mode using distinct single reaction monitoring scans with the following transitions and 969 collision energies - citric acid: m/z 191  $\rightarrow$  111 (CE 12), alpha-ketoglutaric acid: m/z 145 970  $\rightarrow$  101 (CE 10), succinic acid: 117  $\rightarrow$  73 (CE 12), fumaric acid: 115  $\rightarrow$  71 (CE 10), malic 971 acid:  $133 \rightarrow 71$  (CE 15). Chromatographic peak areas were used for comparisons of 972 metabolite abundance. Data were normalized to the average of the control condition 973 and unpaired, two-tailed t-tests were performed in PRISM (GraphPad) to determine 974 significant differences.

975

### 976 S. cerevisiae growth assays

977 Stock solutions of etoposide (Sigma-Aldrich) and ICRF-187 (TCI) were prepared 978 at 100 mM concentration in 100% DMSO then aliquoted and stored at -20 °C. DMSO 979 was added to all no drug experimental controls to match the DMSO content of the drug 980 containing conditions. For spot growth assays, ED yeast were first grown to saturation

981 at 30 °C overnight in SC media. The starter culture was then diluted to  $OD_{600} = 0.1$  in 982 sterile water. In a sterile 96-well plate, the culture was serially diluted down from OD<sub>600</sub> 983 = 0.1 in 5-fold dilution steps. A multichannel pipette was then used to spot 5  $\mu$ L of each 984 dilution on an SC agar plate with the appropriate drug condition for each experiment. 985 Plates were incubated for 2-3 days at 30 °C and imaged by normal photography. For 986 liquid media growth assays, ED or ED-mpc1 $\Delta$  yeast were first grown to saturation at 30 987 °C overnight in SC media with the appropriate carbon source for each experiment. The 988 cultures were then diluted to OD<sub>600</sub> 0.01 in the growth assay media condition and placed 989 in a sterile 24-well plate with 1.5ml of culture in each well. The plates were incubated at 990 30 °C with constant shaking inside a Bio-Tek Synergy HT plate reader for 30 - 35 h. 991 OD<sub>600</sub> measurements were taken every 15 min to generate a growth curve for each 992 well.

993

#### 994 Western Blot

Approximately 1.5 x  $10^8$  cells or 15 OD<sub>600</sub> units of log phase yeast (OD<sub>600</sub> = 0.8-995 996 1.2) were harvested by centrifugation. Pellets were then resuspended in 200 µL of 10% 997 trichloroacetic acid (TCA) and incubated at room temperature for 30 min. The TCA 998 solution was then removed by centrifugation and the pellets were washed with 1 mL of 1 999 M HEPES•KOH (pH 7.5). After removal of the wash solution by centrifugation, the 1000 pellets were then resuspended in 50 µL 2x SDS-PAGE loading buffer with ~50 µL of 0.5 mM glass beads and vortexed for 3 min. An additional 50 µL of 2x SDS-PAGE loading 1001 1002 buffer was added before the samples were boiled for 5 min and then vortexed for 15 sec. After centrifugation to pellet the beads, 20 µL of each supernatant sample were run 1003 1004 on an SDS-PAGE gradient gel and transferred onto a PVDF membrane. The 1005 membranes were blotted with rat anti-HA (1:5000, Roche 11867423001) and rabbit antitubulin (1:5000, abcam ab184970) followed by IRDye 800CW goat anti-Rat (1:15,000, 1006 Li-Cor 926-32219) and IRDye 680CW goat anti-rabbit (1:15,000, Li-Cor 926-68071). 1007 Blots were imaged on a Li-Cor Odyssey system and band intensities were analyzed in 1008 1009 ImageJ.

1010

#### 1011 **Quantification and Statistical Analysis**

- 1012 Data are presented as the mean ± SD of a minimum of three independent experiments,
- 1013 indicated by the n value described in the figure legends.

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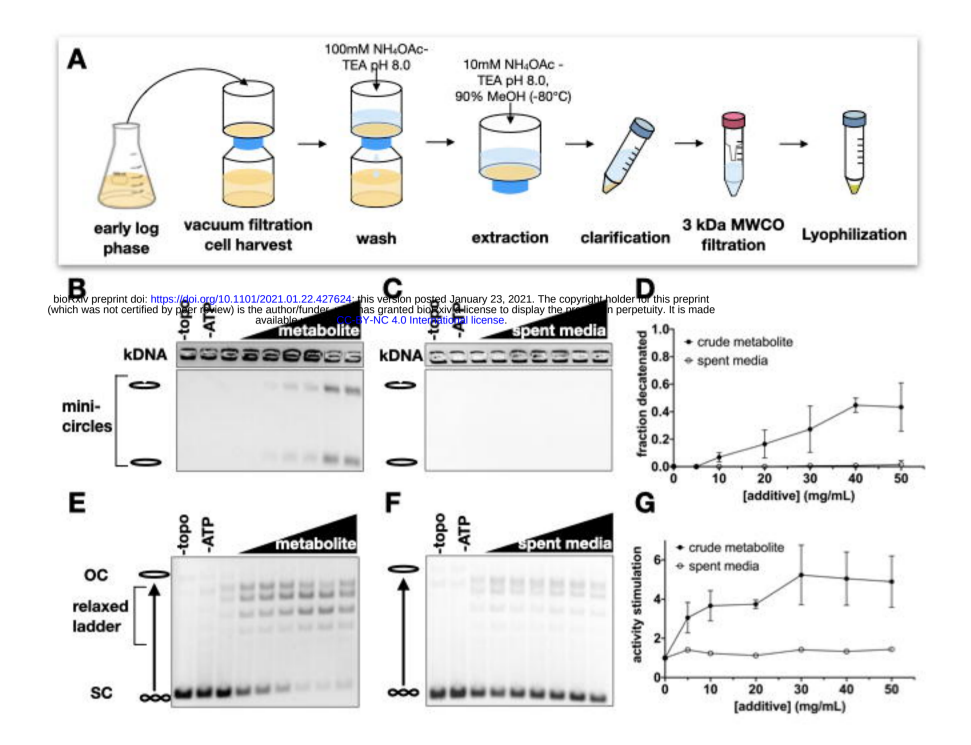
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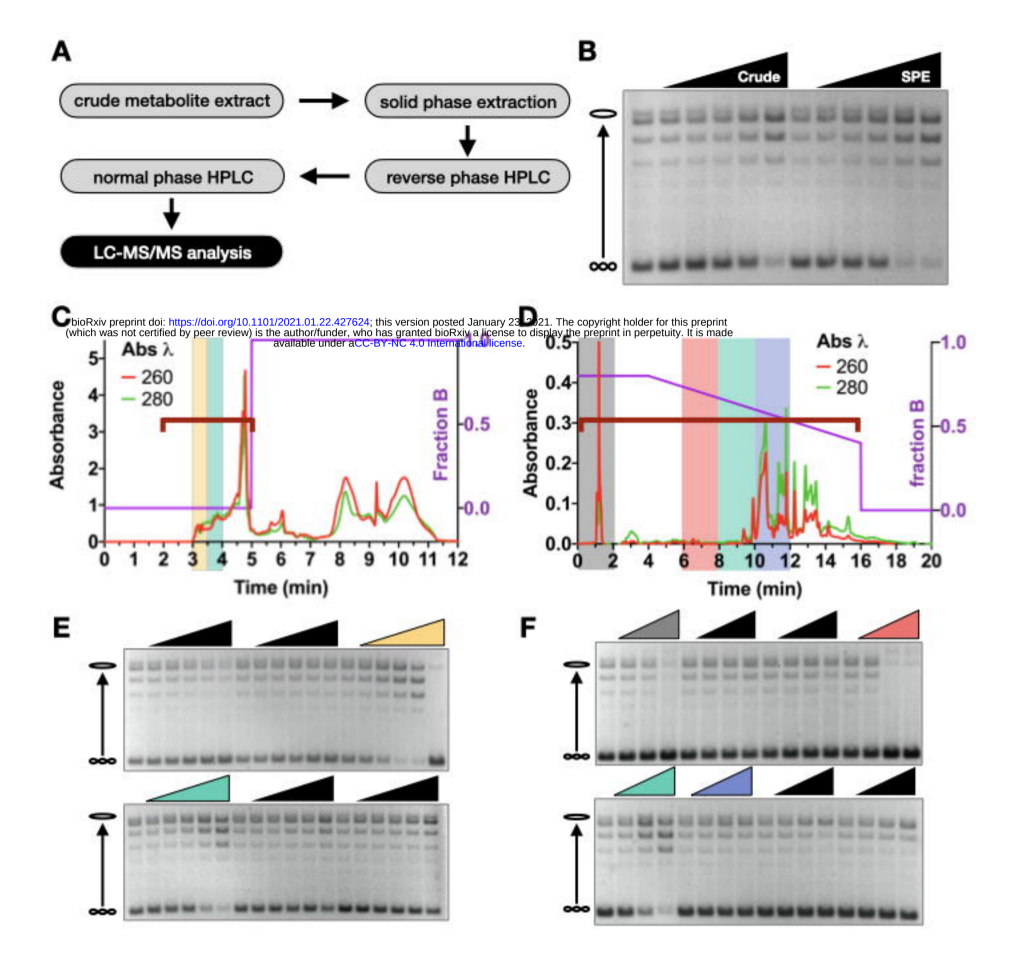
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**Figure 1. Crude metabolite extracts from log-phase yeast stimulate** *Sc***Top2 strand passage activity.** (A) Schematic of metabolite extraction procedure from yeast cultures. (B-C) Representative gels of kinetoplast DNA (kDNA) decatenation assays with metabolite extracts and spent media control samples. Bands representing nicked minicircles and closed minicircles are indicated to the left of the gels. No enzyme (-topo) and no ATP (-ATP) negative controls show the starting substrate. Lyophilized samples of the metabolite extract and spent media were titrated from 0 to 50 mg/ml in 10 mg/ml increments. (D) Decatenation assays with metabolite extracts and spent media control samples. The bands representative gels of supercoil relaxation assays with metabolite extracts and spent media control samples. The bands representing unrelaxed substrate (SC), the relaxed topoisomer distribution, and nicked/open circle (OC) plasmids are indicated to the left of the gels. No enzyme (-topo) and no ATP (-ATP) negative controls show the starting substrate from 0 to 50 mg/ml in 10 mg/ml increments. (SC), the relaxed topoisomer distribution, and nicked/open circle (OC) plasmids are indicated to the left of the gels. No enzyme (-topo) and no ATP (-ATP) negative controls show the starting substrate. Lyophilized samples of the metabolite extract and spent media were titrated from 0 to 50 mg/ml in 10 mg/ml increments. (SC), the relaxed topoisomer distribution, and nicked/open circle (OC) plasmids are indicated to the left of the gels. No enzyme (-topo) and no ATP (-ATP) negative controls show the starting substrate. Lyophilized samples of the metabolite extract and spent media were titrated from 0 to 50 mg/ml in 10 mg/ml increments. (G) Supercoil relaxation assay data represented as mean ± SD of n=3 independent experiments.



**Figure 2. Purification of stimulatory metabolites from crude metabolite extracts.** (A) Schematic of purification steps to prepare samples with enriched stimulatory activity for LC-MS/MS analysis. (B) Supercoil relaxation assay with metabolite samples before ('Crude') and after solid phase extraction ('SPE'). Crude and SPE purified metabolite extracts were added from 0 to 40 mg/ml in two-fold increments (0, 2.5, 5, 10, 20, 40 mg/ml). (C-D) Chromatograms of reverse phase (C) and normal phase (D) HPLC purification runs. The HPLC method is depicted by the purple line showing fraction of phase B as indicated by the right y-axis. Red and green traces show absorbance values at 260 nm and 280 nm wavelengths respectively as indicated on the left y-axis. (E-F) Supercoil relaxation assays of reverse phase fractions (E) and normal phase fractions (F) indicated by the red brackets in (C) and (D). Fractions of interest are highlighted by corresponding colors in the chromatograms and relaxation assay gels. Lyophilized material from each fraction was solubilized in equal volumes of 12.5% DMSO and titrated down from the maximum possible concentration in two-fold dilution steps

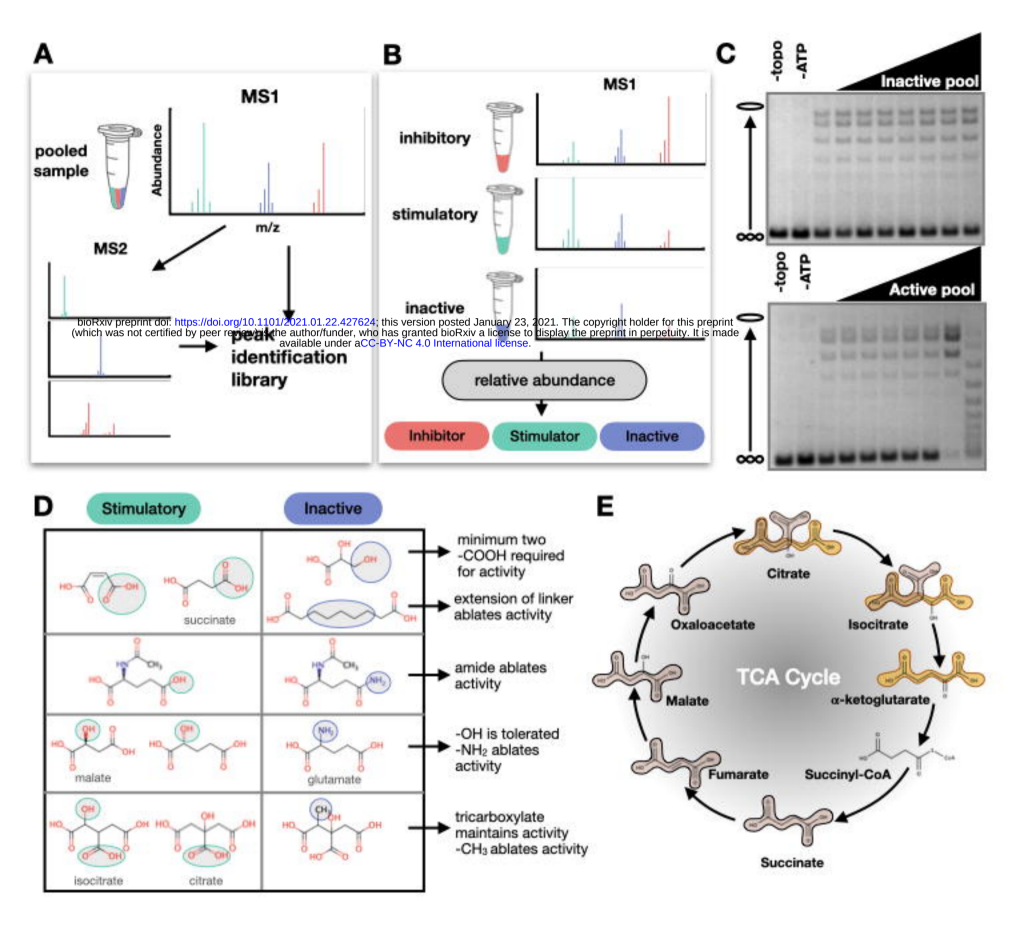
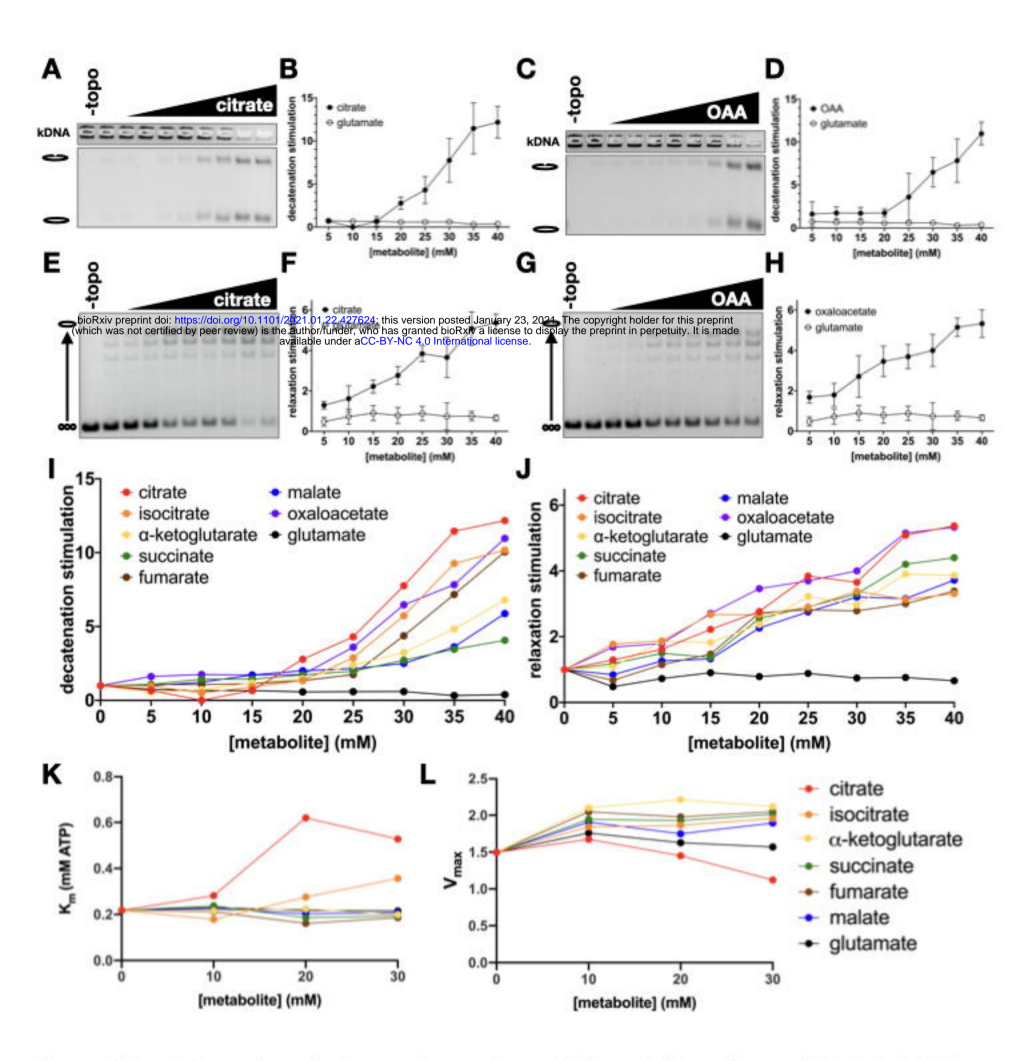


Figure 3. Identification and structure activity relationship (SAR) analysis of stimulatory compounds (A) Schematic depicting how three sequential fractions (indicated by pink, green, and blue in Figure 2) were pooled and analyzed by LC-MS/MS to generate an ion peak identification library. (B) Schematic depicting how different samples (colored as pink, green, and blue) were analyzed individually. Intensities of metabolite peaks were compared across the three samples to determine relative abundance of each ion. Ions that were enriched in the stimulatory fraction (green) were added to a candidate list for stimulatory compounds (see **Table S1**) (C) Example supercoil relaxation assays from testing an inactive and an active pool of five candidate compounds. Candidate pools were prepared from commercial compounds. The concentration of each candidate compound was titrated from 0 to 20 mg/ml in two-fold steps (0, 0.31, 0.63, 1.25, 2.5, 5, 10, 20 mg/ml). (D) SAR analysis comparing stimulatory and inactive compounds identified from candidate pools. Chemical properties that are necessary for activity (green circles) and chemical changes that ablate activity (blue circles) are highlighted in gray. (E) TCA cycle metabolites. Succinic acid (tan) and glutaric acid (gold) motifs are highlighted.



**Figure 4. TCA cycle intermediates stimulate strand passage by topo II without significant effects on ATP hydrolysis.** (A-H) Stimulation of topo II decatenation activity (A-D) and supercoil relaxation activity (E-H) by citrate and oxaloacetate (OAA). Citrate, OAA, and glutamate (negative control) were titrated from 0 to 40mM in 5mM increments. Graphs (B, D, F, and H) represent mean ± SD of n=3-5 independent experiments. (I-J) Stimulation of *Sc*Top2 decatenation (I) and relaxation (J) activity by TCA metabolites and glutamate (negative control). Stimulation values represent mean of n=3-6 independent experiments. (See also **Figure S3**). (K-L) Effects of TCA metabolites on ATP hydrolysis activity. Experiments were performed at 0, 10, 20, and 30mM concentrations of the metabolites indicated in the legend to the right of the graphs. K<sub>m</sub> (K) and V<sub>max</sub> (µmol ATP • min<sup>-1</sup> • nmol topo II<sup>-1</sup>) (L) values were derived from n=3 independent experiments. (See also **Figure S5**).

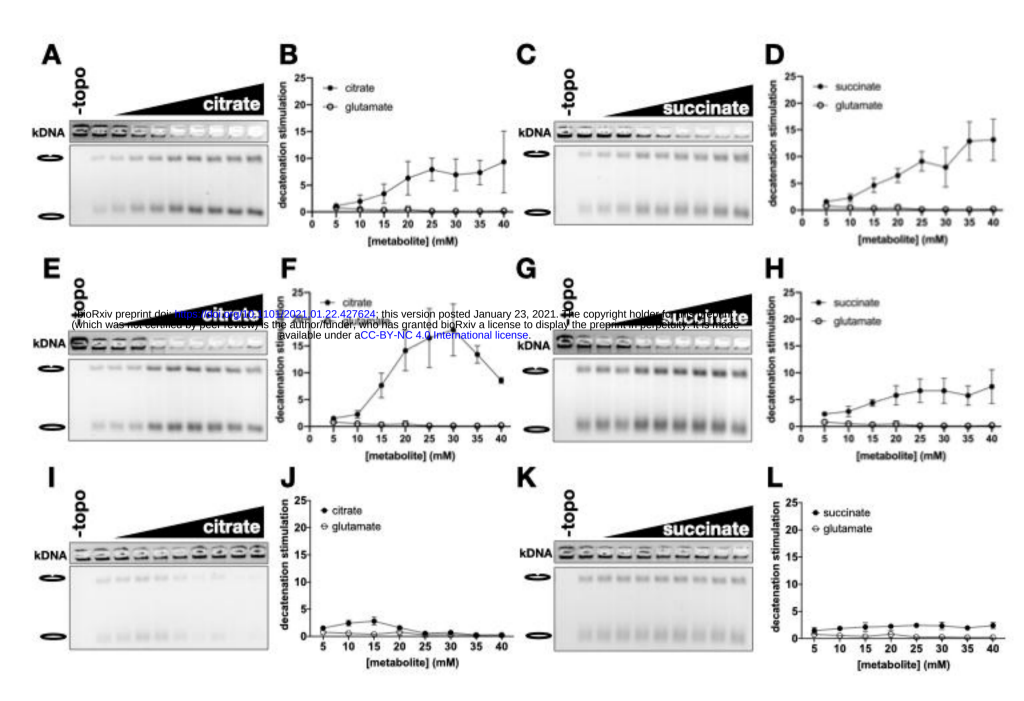


Figure 5. Stimulatory effect of TCA metabolites is eukaryotic specific. Citrate and succinate were titrated from 0 to 40 mM in 5 mM increments and added to decatenation assays with HsTop2B (A-D), HsTop2A (E-H) and Ec topo IV (I-L). Graphs represent mean ± SD of n=3-5 independent experiments.

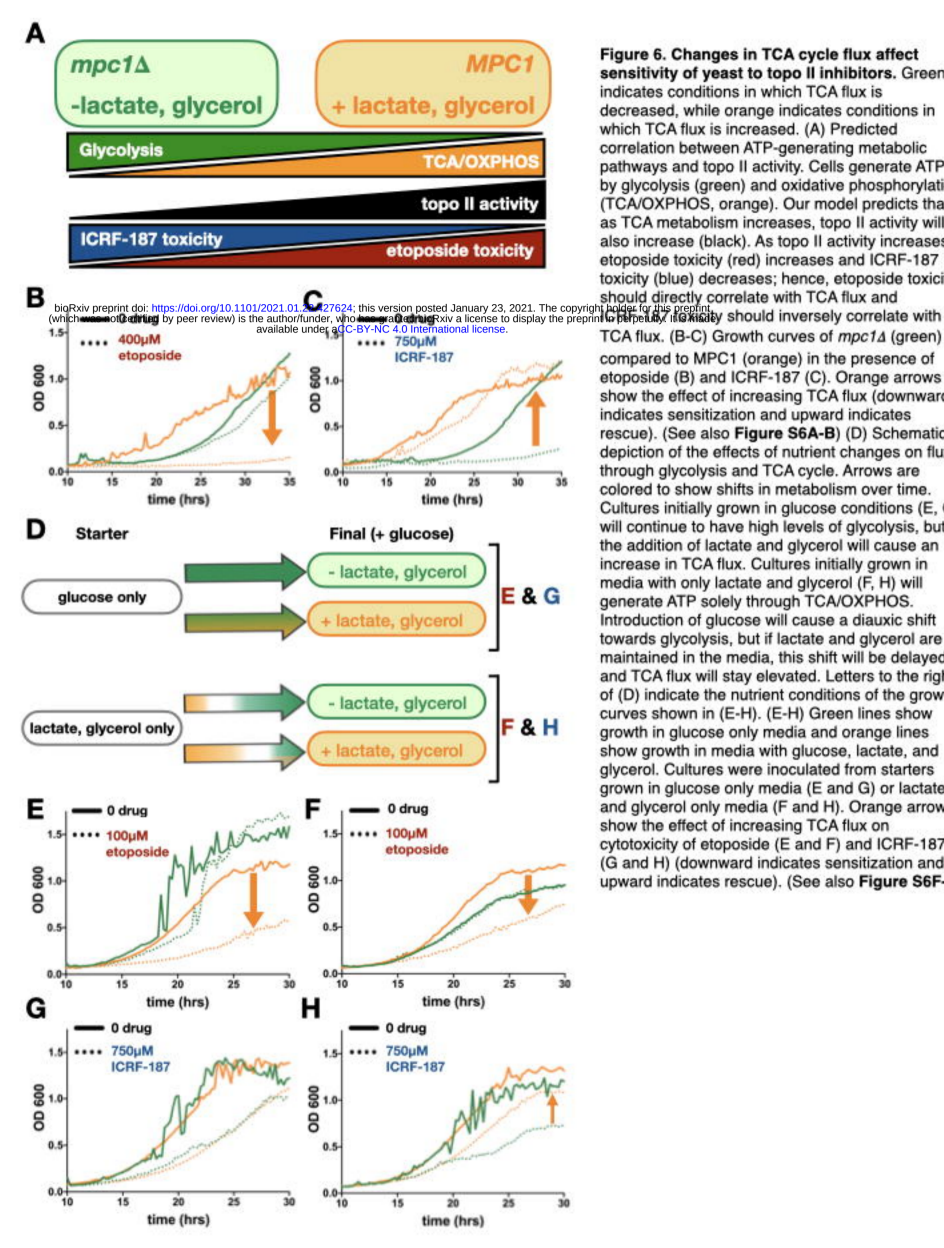


Figure 6. Changes in TCA cycle flux affect sensitivity of yeast to topo II inhibitors. Green indicates conditions in which TCA flux is decreased, while orange indicates conditions in which TCA flux is increased. (A) Predicted correlation between ATP-generating metabolic pathways and topo II activity. Cells generate ATP by glycolysis (green) and oxidative phosphorylation (TCA/OXPHOS, orange). Our model predicts that as TCA metabolism increases, topo II activity will also increase (black). As topo II activity increases, etoposide toxicity (red) increases and ICRF-187 toxicity (blue) decreases; hence, etoposide toxicity

compared to MPC1 (orange) in the presence of etoposide (B) and ICRF-187 (C). Orange arrows show the effect of increasing TCA flux (downward indicates sensitization and upward indicates rescue). (See also Figure S6A-B) (D) Schematic depiction of the effects of nutrient changes on flux through glycolysis and TCA cycle. Arrows are colored to show shifts in metabolism over time. Cultures initially grown in glucose conditions (E, G) will continue to have high levels of glycolysis, but the addition of lactate and glycerol will cause an increase in TCA flux. Cultures initially grown in media with only lactate and glycerol (F, H) will generate ATP solely through TCA/OXPHOS. Introduction of glucose will cause a diauxic shift towards glycolysis, but if lactate and glycerol are maintained in the media, this shift will be delayed and TCA flux will stay elevated. Letters to the right of (D) indicate the nutrient conditions of the growth curves shown in (E-H). (E-H) Green lines show growth in glucose only media and orange lines show growth in media with glucose, lactate, and glycerol. Cultures were inoculated from starters grown in glucose only media (E and G) or lactate and glycerol only media (F and H). Orange arrows show the effect of increasing TCA flux on cytotoxicity of etoposide (E and F) and ICRF-187 (G and H) (downward indicates sensitization and upward indicates rescue). (See also Figure S6F-I).

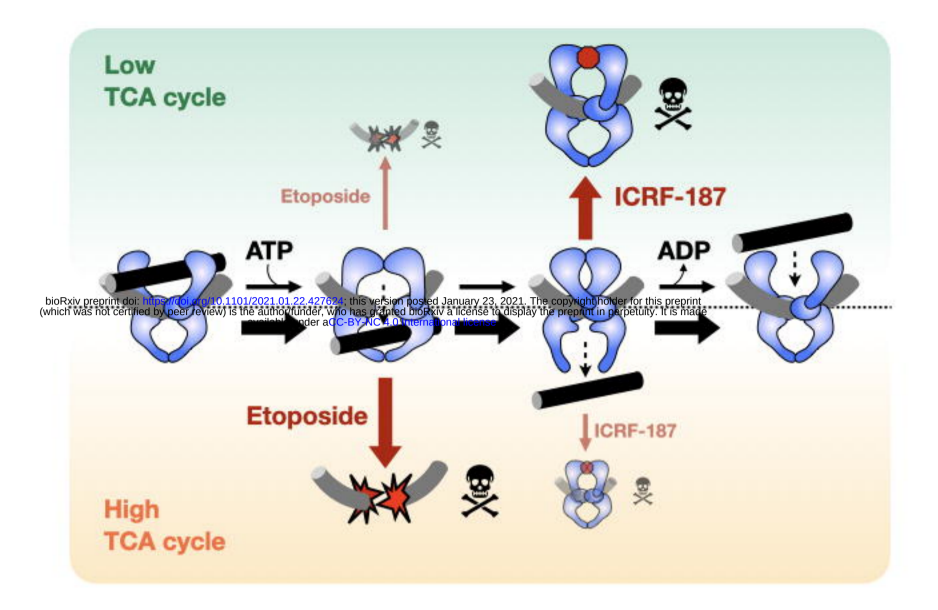


Figure 7. Schematic depicting how sensitivity to topo II targeting drugs is influenced by metabolic state. When TCA cycle flux is low (green) topo II strand passage activity is not stimulated. General catalytic inhibitors (e.g., ICRF-187) that work by decreasing topo II activity will be more effective in this state, as compared to a high TCA cycle flux state (orange). By contrast, topo II poisons (e.g., etoposide) are more toxic when topo II activity is stimulated in the high TCA cycle flux state because elevated topo II strand passage activity leads to increased formation of cytotoxic DNA double strand breaks.

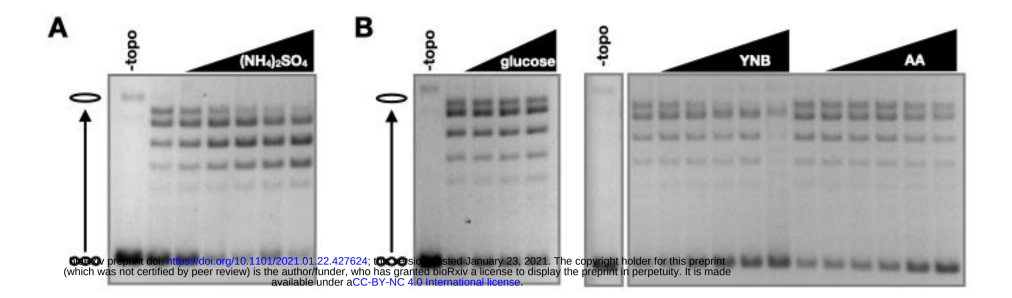


Figure S1. Effects of synthetic media components on topo II activity. Gels show supercoil relaxation activity of *Sc*Top2 at increasing concentrations of the four components of synthetic media. Negative controls with no enzyme (-topo) show the unrelaxed plasmid substrate. (A) (NH4)<sub>2</sub>SO<sub>4</sub> was titrated from 0 to 50 mM in 10 mM increments (0, 10, 20, 30, 40, 50 mM). (B) Glucose was titrated from 0 to 40 mg/ml in two-fold increments (0, 10, 20, 40 mg/ml). Yeast nitrogen base without added sulfate (YNB) and an amino acid mixture containing all 20 amino acids in equal amounts (AA) were titrated from 0 to 10 mg/ml in two-fold increments (0, 0.625, 1.25, 5, 10 mg/ml).

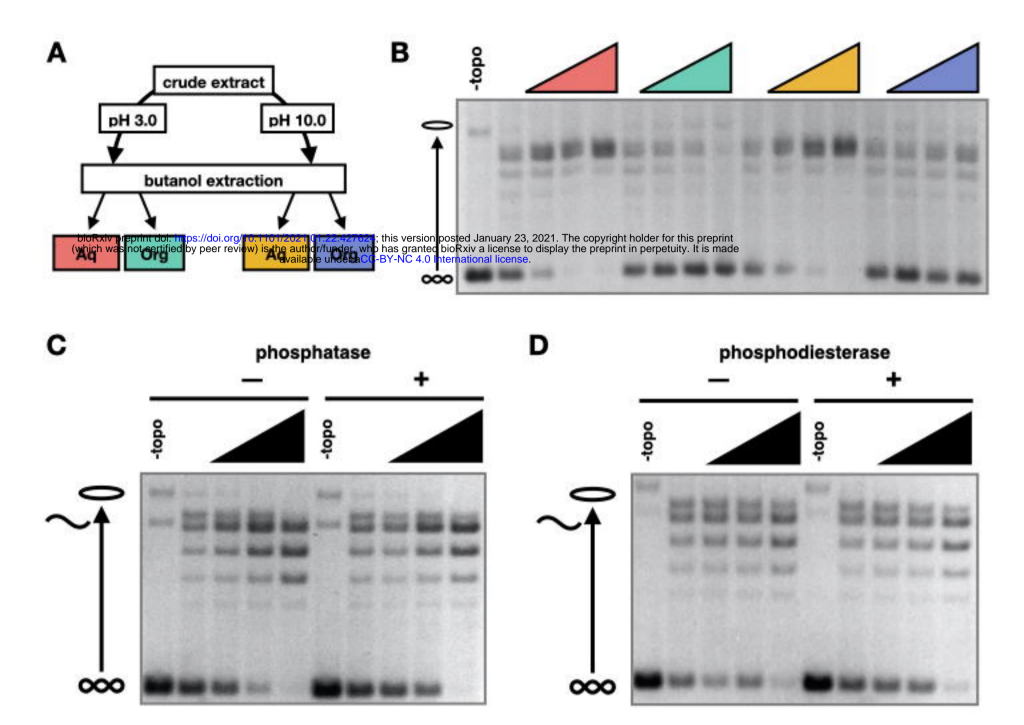


Figure S2. Chemical properties of stimulatory metabolites. (A) Schematic for acid/base butanol liquid-liquid extraction of crude metabolite extracts. 'Aq' indicates the aqueous fraction, and 'Org' indicates the organic fraction. (B) Supercoil relaxation assay with fractions from acid/base butanol extraction as indicated by colors. Solid material left after solvent removal from each fraction was solubilized in equal volumes of 12.5% DMSO and titrated into relaxation assays in 2-fold dilution steps. (C-D) Activity of crude metabolites after enzymatic treatment by Antarctic phosphatase (C) or snake venom phosphodiesterase (D). Lyophilized material from treated and untreated samples were solubilized in equal volumes of 12.5% DMSO and titrated into relaxation assays in 2-fold dilution steps.

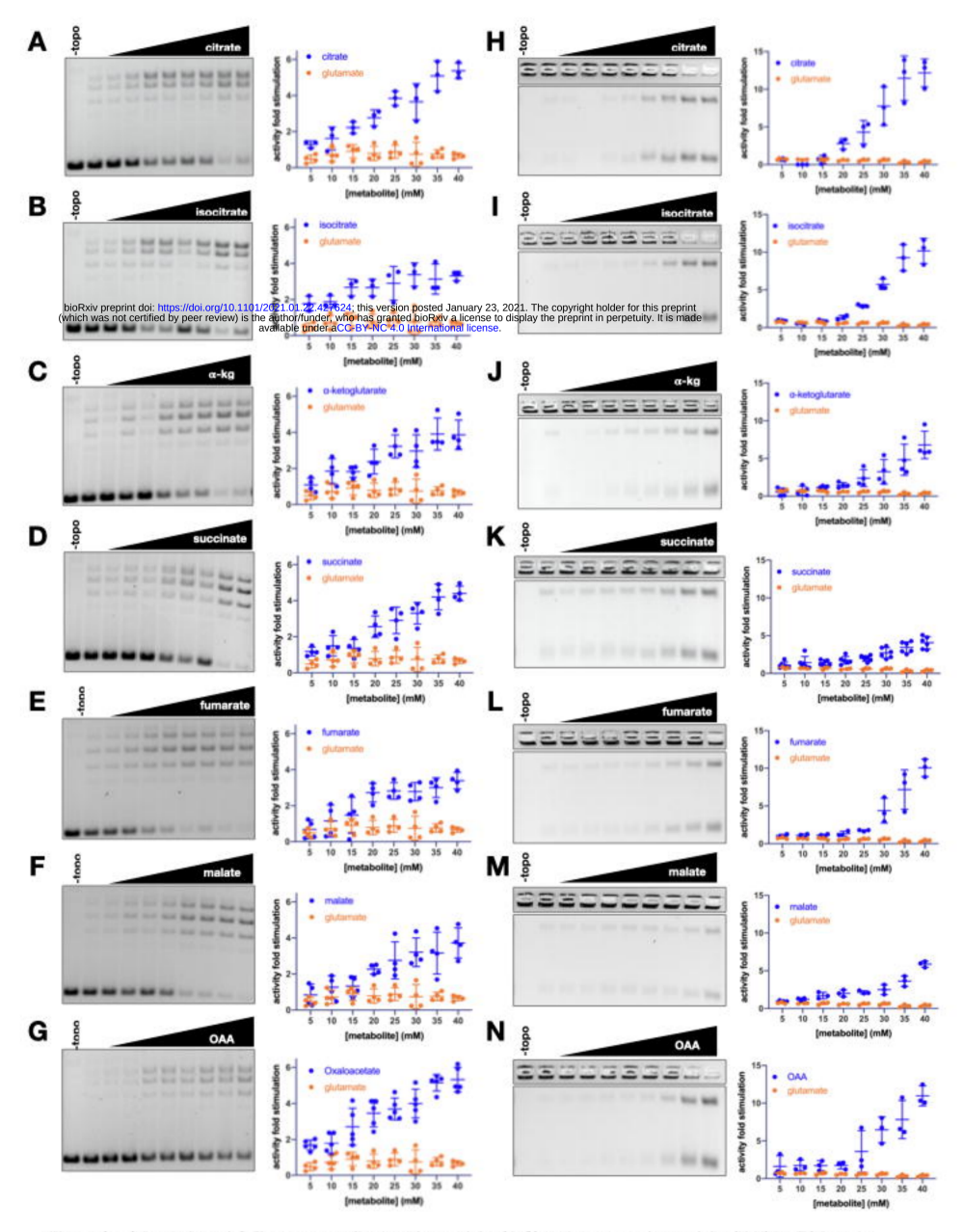


Figure S3. Stimulation of *Sc*Top2 supercoil relaxation activity (A-G) and decatenation activity (H-N) by TCA cycle intermediates (blue) and glutamate (negative control, orange). Graphs show mean ± SD of n=3-6 independent experiments.

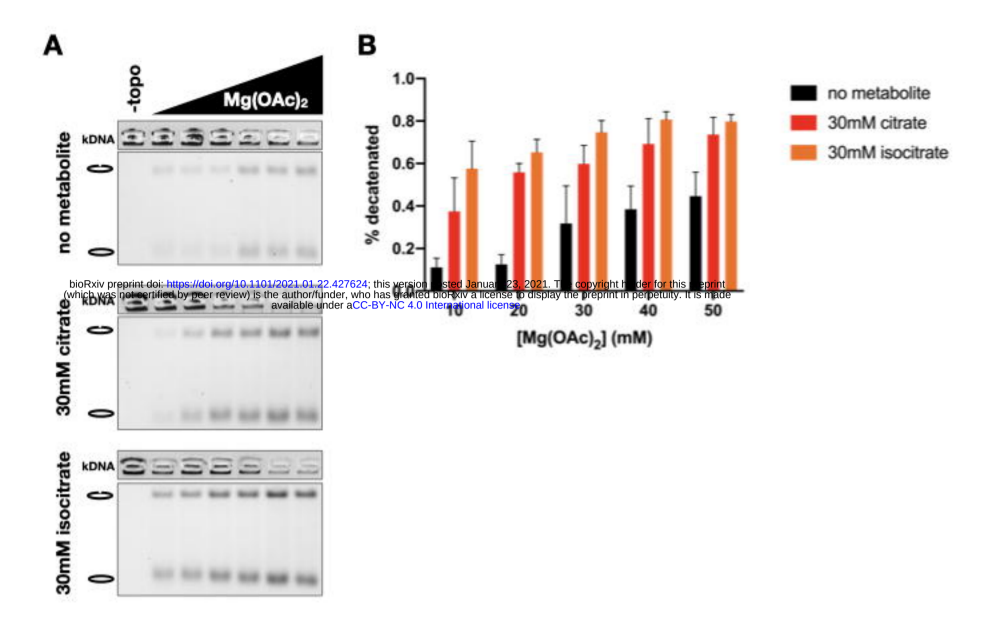


Figure S4. Chelation of Mg<sup>2+</sup> ions by tricarboxylate TCA cycle intermediates does not affect stimulatory topo II-metabolite interaction. (A) Representative gels of decatenation assays performed at different concentrations of Mg(OAc)<sub>2</sub> with no added metabolite, 30mM citrate, and 30mM isocitrate. Mg(OAc)<sub>2</sub> was titrated from 10 to 50 mM in 10 mM steps. (B) Graphs show mean ± SD of n=3 independent experiments to compare the fraction decatenated under each reaction condition.

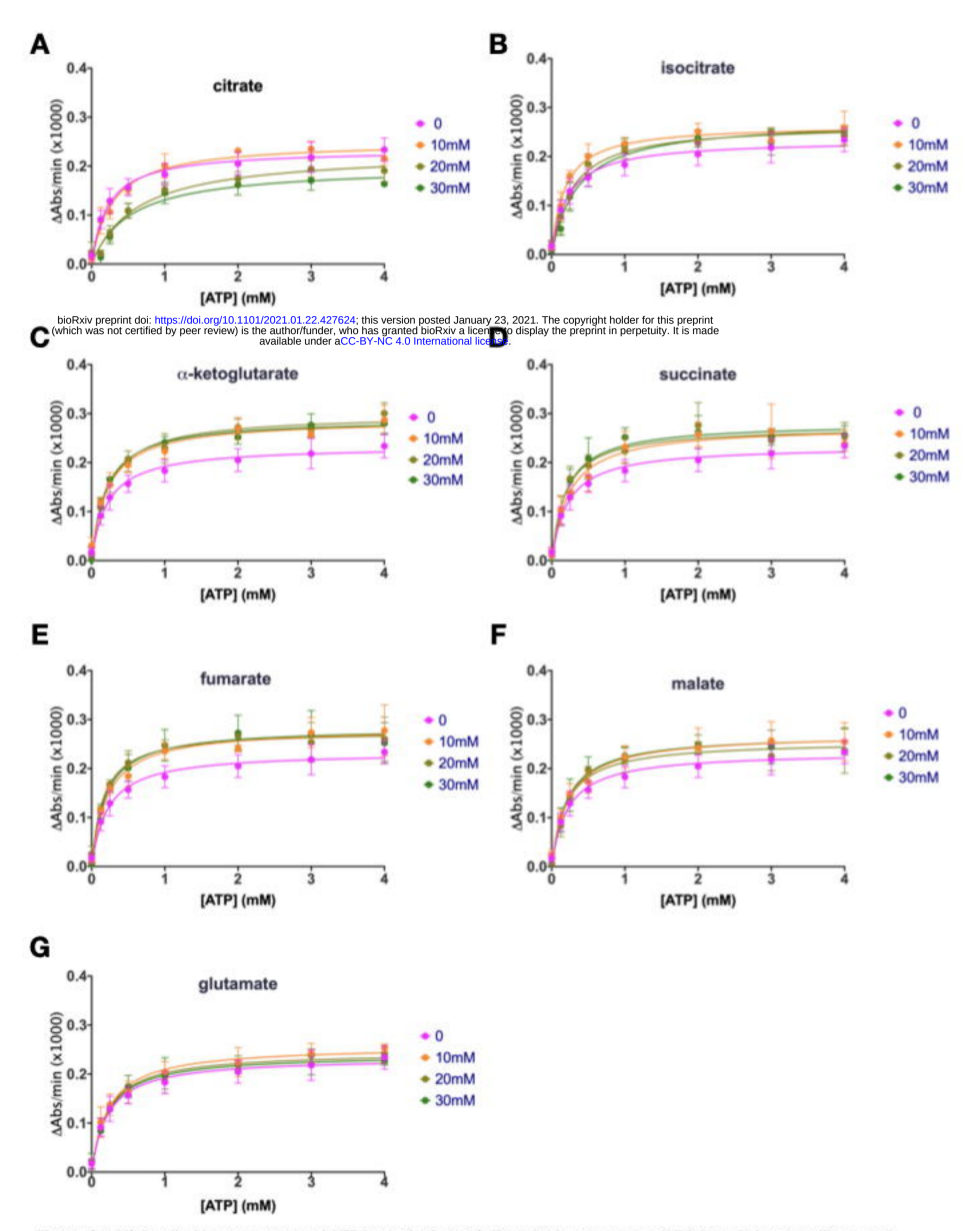


Figure S5. Michaelis-Menten curves of ATP hydrolysis by ScTop2 in the presence of TCA cycle intermediates and glutamate (negative control). Rates of ATP hydrolysis were measured in a coupled assay by monitoring NADH consumption as a function of light absorbance at 340 nm wavelength (y-axis). Graphs represent mean ± SD of n=3 independent experiments.

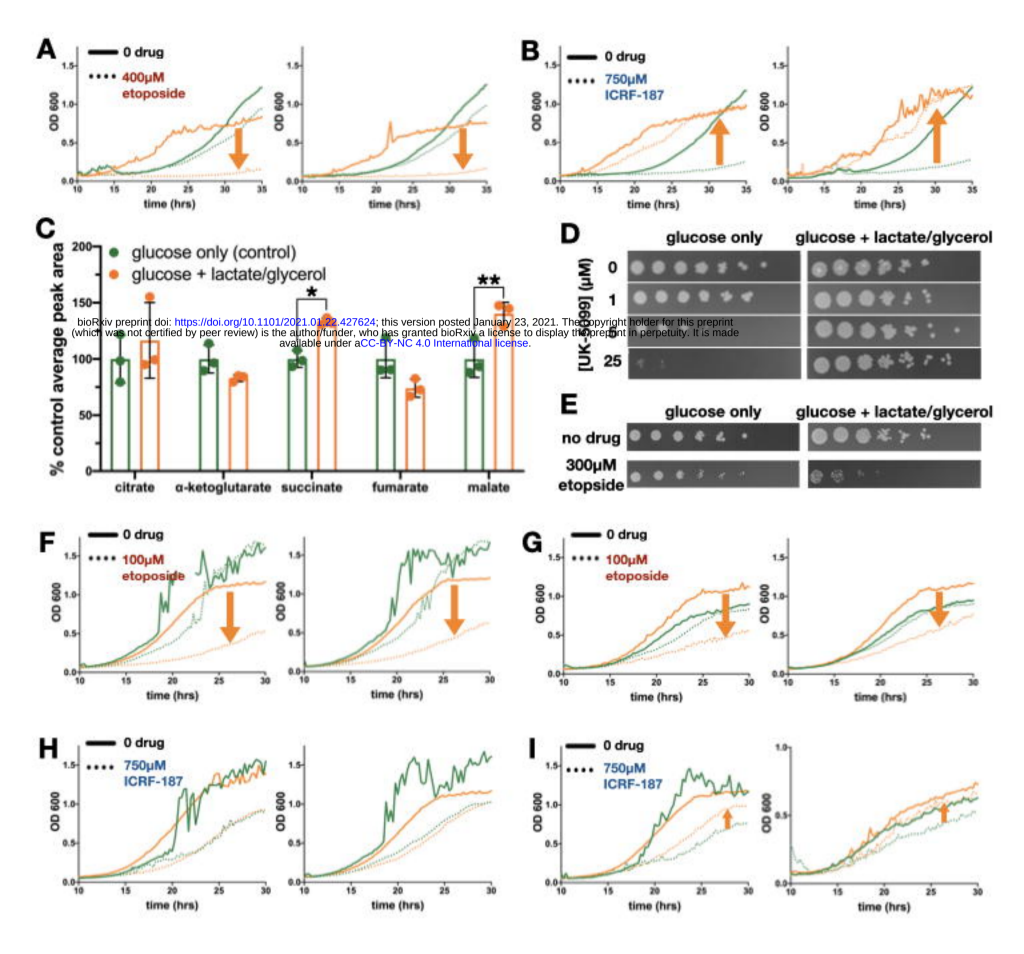


Figure S6. Changes in TCA cycle flux affect sensitivity of yeast to topo II inhibitors - extended data. (A-B) Growth curves of MPC1 (orange) and *mpc1*<sub>Δ</sub> (green) with etoposide (A) and ICRF-187 (B). (C) Changes in TCA cycle metabolite abundance upon addition of lactate and glycerol to media. Peak areas were normalized to the average of the control condition (glucose only). Bar graphs and error bars indicate mean  $\pm$  SD of n=3. An unpaired, two-tailed t-test was used to analyze significance of changes peak area. One asterisk (\*) indicates p < 0.05, and two asterisks (\*\*) indicates p < 0.01. (D-E) Serially diluted cultures of yeast were spotted on glucose containing agar plates with and without lactate and glycerol. Spot growth at 0-25  $\mu$ M UK-5099 (D) and 300  $\mu$ M etoposide (E) was observed after 2-3 days. (F-H) Growth curves of yeast in glucose-only media (green) or glucose media supplemented with lactate and glycerol (orange). Cultures were inoculated from starters grown in glucose only media (F and H) or lactate and glycerol only media (G and H). Orange arrows show the effect of increasing TCA flux on cytotoxicity of etoposide (F and G) and ICRF-187 (H and I) (downward indicates sensitization and upward indicates rescue).

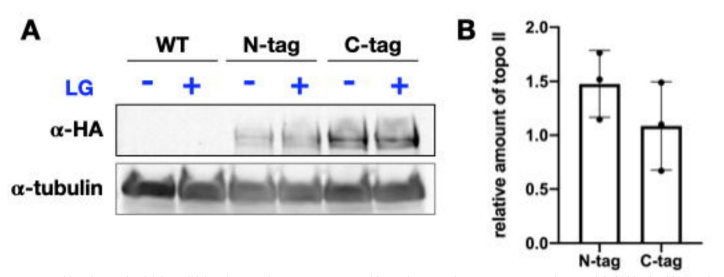


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