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1 **Rapid degradation of *C. elegans* proteins at single-cell resolution with a synthetic**
2 **auxin**

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39 **ABSTRACT**

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41 As developmental biologists in the age of genome editing, we now have access to an
42 ever-increasing array of tools to manipulate endogenous gene expression. The auxin-
43 inducible degradation system, allows for spatial and temporal control of protein
44 degradation, functioning through the activity of a hormone-inducible *Arabidopsis* F-box
45 protein, transport inhibitor response 1 (TIR1). In the presence of auxin, TIR1 serves as a
46 substrate recognition component of the E3 ubiquitin ligase complex SKP1-CUL1-F-box
47 (SCF), ubiquitinating auxin-inducible degron (AID)-tagged proteins for proteasomal
48 degradation. Here, we optimize the *Caenorhabditis elegans* AID method, utilizing 1-
49 naphthaleneacetic acid (NAA), an indole-free synthetic analog of the natural auxin indole-
50 3-acetic acid (IAA). We take advantage of the photostability of NAA to demonstrate via
51 quantitative high-resolution microscopy that rapid degradation of target proteins can be
52 detected in single cells within 30 minutes of exposure. Additionally, we show that NAA
53 works robustly in both standard growth media and physiological buffer. We also
54 demonstrate that K-NAA, the water-soluble, potassium salt of NAA, can be combined with
55 microfluidics for targeted protein degradation in *C. elegans* larvae. We provide insight into
56 how the AID system functions in *C. elegans* by determining that TIR1 interacts with *C.*
57 *elegans* SKR-1/2, CUL-1, and RBX-1 to degrade target proteins. Finally, we present
58 highly penetrant defects from NAA-mediated degradation of the Ftz-F1 nuclear hormone
59 receptor, NHR-25, during *C. elegans* uterine-vulval development. Together, this work
60 provides a conceptual improvement to the AID system for dissecting gene function at the
61 single-cell level during *C. elegans* development.

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62 INTRODUCTION

63

64 *In situ* techniques for targeted protein degradation enable a detailed analysis of
65 developmental events, mechanisms, and functions. RNAi and Cre or Flp-mediated
66 recombination (Qadota *et al.* 2007; Hubbard 2014; Shen *et al.* 2014) allow tissue-specific
67 study of gene products, but the persistence of the target protein following recombination
68 or RNA depletion can delay the manifestation of an otherwise acute phenotype. Several
69 methods have been described recently to enable tissue-specific protein degradation in
70 *Caenorhabditis elegans*, including ZF1 tagging (Armenti *et al.* 2014), a GFP nanobody
71 approach (Wang *et al.* 2017), sortase A (Wu *et al.* 2017), and auxin-mediated degradation
72 (Zhang *et al.* 2015).

73

74 The auxin-inducible degradation system allows for rapid and conditional
75 degradation of auxin-inducible degron (AID)-tagged proteins in *C. elegans* as well as in
76 other commonly used model systems including yeast (Nishimura *et al.* 2009), *Drosophila*
77 (Trost *et al.* 2016), zebrafish (Daniel *et al.* 2018), cultured mammalian cells (Nishimura *et*
78 *al.* 2009; Holland *et al.* 2012; Natsume *et al.* 2016), and mouse oocytes (Camlin and
79 Evans 2019). This protein degradation system relies on the expression of an *Arabidopsis*
80 F-box protein called transport inhibitor response 1 (TIR1). As a substrate-recognition
81 component of the SKP1-CUL1-F-box (SCF) E3 ubiquitin ligase complex, TIR1 carries out
82 its function only in the presence of the hormone auxin. Once bound to auxin, TIR1 targets
83 AID-tagged proteins for ubiquitin-dependent proteasomal degradation (**Figure 1A**).

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84 The *C. elegans* version of the AID system is robust and specific with minimal off-
85 target effects (Zhang *et al.* 2015). However, re-evaluation of the system is needed to
86 assess its utility among *C. elegans* researchers conducting microscopy-based single-cell
87 biology within a narrow developmental time frame. Here, we use 1-naphthaleneacetic
88 acid (NAA) and its water-soluble potassium salt analog (K-NAA), indole-free synthetic
89 analogs of the natural auxin indole-3-acetic acid (IAA), to degrade target proteins at
90 single-cell resolution in *C. elegans* larvae in standard growth media and physiological
91 buffer. Given the ability to solubilize K-NAA solely in water or physiological buffer (M9),
92 we also demonstrated rapid degradation kinetics of an AID-tagged transgene in a *C.*
93 *elegans*-based microfluidics for the first time (Keil *et al.* 2017). Next, we sought to gain
94 insight into which SCF complex members interact with TIR1, identifying *skr-1/2*, *cul-1* and
95 *rbx-1* as putative TIR1 interactors through RNAi depletion experiments. Finally, we
96 demonstrate potent temporal effects on uterine and vulval development following targeted
97 degradation of endogenous NHR-25, the single *C. elegans* homolog of *Drosophila* Ftz-
98 F1 and human SF-1 and LRH-1 (Chen *et al.* 2004; Ward *et al.* 2013). It is our hope that
99 this synthetic auxin analog will be applied at all stages of *C. elegans* development,
100 allowing for precise, rapid degradation of target proteins in a high-resolution and
101 quantitative fashion.

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106 MATERIALS AND METHODS

107

108 ***C. elegans strains and culture conditions***

109 Animals were maintained using standard culture conditions at 25°C (Brenner 1974) and
110 were synchronized through alkaline hypochlorite treatment of gravid adults to isolate eggs
111 (Porta-de-la-Riva *et al.* 2012). In the main text and figure legends, we designate linkage
112 to a promoter using a greater than symbol (>) and fusion to a protein using a double colon
113 (::). The following alleles and transgenes were used in this manuscript for experimental
114 purposes: LG I: *kry61[nhr-23::AID]*; LG II: *ieSi57[eft-3>TIR1::mRuby]*; LG IV: *ieSi58[eft-*
115 *3>AID::GFP]*, *syIs49 [zmp-1>GFP]*; LG X: *wrd10[nhr-25::GFP::AID]*.

116

117 ***Constructs and microinjection***

118 SapTrap was used to construct the 30xlinker::GFP^SEC^TEV::AID degron::3xFLAG
119 repair template (pJW1747) for generating the knock-in into the 3' end of the *nhr-25* gene
120 (Schwartz and Jorgensen 2016). DH10β competent *E. coli* cells, made in-house, were
121 used for generating the plasmid. The following reagents were used to assemble the final
122 repair template: pDD379 (backbone with F+E sgRNA), annealed oligos 3482+3483
123 (sgRNA), pJW1779 (5' homology arm), 3' homology arm PCR product, pJW1347 (30x
124 linker for CT slot), pDD372 (GFP for FP slot), pDD363 (SEC with LoxP sites), and
125 pJW1759 (TEV::AID degron::3xFLAG for NT slot).

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127 The pJW1747 repair template was purified using the Invitrogen PureLink HQ Mini
128 Plasmid DNA Purification Kit (K210001). The optional wash step in the protocol using a
129 4 M guanidine-HCl + 40% isopropanol solution is highly recommended, as excluding it
130 dramatically reduced injection efficiency in our hands. N2 animals were injected with a
131 mix consisting of 10 ng/μl of pJW1747, 50 ng/μl of pDD121 (Cas9 vector), and co-injection
132 markers (10 ng/μl pGH8, 5 ng/μl pCFJ104, 2.5 ng/μl pCFJ90) as previously described
133 (Frøkjær-Jensen *et al.* 2012; Dickinson *et al.* 2013, 2015). Knock-ins were isolated as
134 previously described (Dickinson *et al.* 2015). Each knock-in junction was verified via PCR
135 using a primer that bound outside the homology arm paired with a primer binding within
136 pJW1747. The knock-in was backcrossed five times against wild-type N2 animals to
137 produce JDW58. The SEC was then excised by heat-shock (Dickinson *et al.* 2015) to
138 produce JDW59; the knock-in sequence was re-confirmed by PCR amplification and
139 sequencing, using the oligos flanking the homology arms. JDW58 was crossed to CA1200
140 (*eft-3>TIR1::mRuby*) to generate JDW70. The SEC was then excised (Dickinson *et al.*
141 2015) to produce JDW71.

142

143 pDD121, pDD363, pDD372, and pDD379 (Dickinson *et al.* 2018) were gifts from
144 Bob Goldstein (Addgene plasmid numbers are 91833, 91829, 91824, and 91834,
145 respectively). pJW1347 and pJW1759 will be deposited into Addgene's repository and
146 are also available upon request. pJW1347 and pJW1759 were generated by TOPO blunt
147 cloning of PCR products. pJW1779 was generated by Gateway cloning into pDONR221
148 (Invitrogen). Oligo sequences used to generate these plasmids, the sgRNA, the 3'
149 homology arm, and for genotyping are in the Reagent Table.

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150 ***Auxin experiments***

151 For all auxin experiments, synchronized L1 larval stage animals were first transferred to
152 standard nematode growth media (NGM) agar plates seeded with *E. coli* OP50 and then
153 transferred at the P6.p 2-cell stage (mid-L3 stage) to either OP50-seeded NGM agar
154 plates treated with IAA, NAA, or K-NAA, or M9 buffer treated with NAA in the absence of
155 bacteria, NAA plus *E. coli* NA22, or K-NAA plus *E. coli* NA22.

156

157 For IAA and NAA experiments on plates, a 250 mM stock solution in 95% ethanol
158 was prepared using powder IAA purchased from Alfa Aesar (A10556) and powder NAA
159 purchased from Sigma-Aldrich (317918) and stored at -20°C. IAA and NAA were then
160 diluted into the NGM agar (cooled to approximately 50°C) at the time of pouring plates.
161 Fresh OP50 was used to seed plates. For control experiments, 0.25% ethanol was used
162 as described previously (Zhang *et al.* 2015). For K-NAA experiments on plates, a 250 mM
163 stock solution in deionized water was prepared using powder K-NAA purchased from
164 PhytoTechnology Laboratories (N610) and stored at 4°C. For control experiments, OP50-
165 seeded NGM agar plates were used. Prior to each NAA experiment in M9 buffer, a fresh
166 1 mM (pH of 7.22) or 4 mM solution (pH of 8.14) in M9 buffer was prepared using 5.4 mM
167 NAA purchased in liquid form from Sigma-Aldrich (N1641). These pH levels are well
168 within the tolerance range of *C. elegans* for pH (Khanna *et al.* 1997). M9 buffer alone (pH
169 of 7.13) was used as a control. A detailed protocol for liquid-based NAA-mediated
170 degradation can be found in File S1. For experiments conducted in the microfluidic
171 platform, a 4 mM NAA or K-NAA solution in M9 buffer containing *E. coli* NA22 was
172 prepared and stored at 4°C for up to 2 weeks. M9 buffer containing NA22 was used as a

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173 control. See File S2 for a detailed protocol describing the preparation of media for the
174 microfluidic device.

175

176 **Brood size and viability assays**

177 Brood size and viability assays were performed as described (Zhang *et al.* 2015). Briefly,
178 L4 hermaphrodites were picked onto individual MYOB plates containing 0% ethanol (K-
179 NAA control), 0.25% ethanol (IAA control), 4 mM K-NAA, or 4 mM IAA. Animals were then
180 transferred to new plates daily over 4 days. The eggs laid on each plate were counted
181 after removing the parent and viable progeny were quantified when the F1 reached L4 or
182 adult stages (2-3 days post egg-laying). At this point, we also scored for dead eggs. Brood
183 size is the sum of live progeny and dead eggs. Percent embryonic lethality was
184 determined by dividing dead eggs by total eggs laid.

185

186 **RNAi experiments**

187 RNAi targeting *cul-1* was constructed by cloning 997 bp of synthetic DNA based on its
188 cDNA sequence available on WormBase (wormbase.org) into the highly efficient T444T
189 RNAi vector (Sturm *et al.* 2018). The synthetic DNA was generated by Integrated DNA
190 Technologies (IDT) as a gBlock gene fragment and cloned into the BgIII/Sall restriction
191 digested T444T vector using the NEBuilder HiFi DNA Assembly Master Mix (E2621).
192 RNAi feeding strains silencing *skr-1/2*, *skr-7*, *skr-10*, and *rbx-1* were obtained from the
193 Vidal RNAi library (Rual *et al.* 2004).

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194 ***Scoring defects in anchor cell (AC) specification***

195 Synchronized L1 stage *nhr-25::GFP::AID; eft-3>TIR1::mRuby* animals were plated onto
196 NGM agar plates containing either control or 4 mM NAA and grown for 24 hours at 25°C
197 until the early L3 stage (P6.p 1-cell stage), after the normal time of AC specification.
198 Images were acquired as specified below to score for the presence or absence of an AC,
199 visualized by characteristic morphology using DIC optics.

200

201 ***Scoring vulva precursor cell (VPC) arrest***

202 Synchronized L1 stage *nhr-25::GFP::AID; eft-3>TIR1::mRuby* animals were plated onto
203 OP50 NGM agar plates and allowed to grow until the P6.p 1-cell stage. Animals were
204 then washed off plates with M9 and transferred onto NGM agar plates containing either
205 control or 4 mM NAA and grown at 25°C until the mid-L3 stage, after the normal time of
206 P6.p cell division. Images were acquired as specified below to score for P6.p divisions
207 using DIC optics. Remaining animals were scored for plate level adult phenotypes
208 approximately 24 hours later.

209

210 ***Image acquisition***

211 Images were acquired using a Hamamatsu Orca EMCCD camera and a Borealis-
212 modified Yokagawa CSU-10 spinning disk confocal microscope (Nobska Imaging, Inc.)
213 with a Plan-APOCHROMAT x 100/1.4 oil DIC objective controlled by MetaMorph software
214 (version: 7.8.12.0). Animals were anesthetized on 5% agarose pads containing 10 mM

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215 sodium azide and secured with a coverslip. Imaging on the microfluidic device was
216 performed on a Zeiss AXIO Observer.Z7 inverted microscope using a 40X glycerol
217 immersion objective and DIC and GFP filters controlled by ZEN software (version 2.5).
218 Images were captured using a Hamamatsu C11440 digital camera. For scoring plate level
219 phenotypes, images were acquired using a Moticam CMOS (Motic) camera attached to
220 a Zeiss dissecting microscope.

221

222 ***Image processing and analyses***

223 All acquired images were processed using Fiji software (version: 2.0.0-rc-69/1.52p)
224 (Schindelin *et al.* 2012). To quantify AC- or VPC-specific degradation of AID::GFP,
225 images were captured at the P6.p 2-cell stage and 4-cell stage (mid-L3 stage) at time
226 points 0, 30, 60, 90, and 120 minutes in the absence or presence of auxin. Expression of
227 *eft-3>AID::GFP* was quantified by measuring the mean fluorescence intensity (MFI) of
228 ACs and VPCs subtracted by the MFI of a background region in the image to account for
229 camera noise. Cells were outlined using the freehand selection tool in Fiji. Data were
230 normalized by dividing the MFI in treated or untreated animals at time points 30, 60, 90,
231 and 120 minutes by the average MFI in untreated animals at 0 minutes. For experiments
232 utilizing RNAi, only ACs were measured due to the variable sensitivity of VPCs to RNAi
233 (Bourdages *et al.* 2014; Matus *et al.* 2014). To quantify AC-specific degradation of
234 AID::GFP in animals fed RNAi overnight, images were captured at the P6.p 2-cell stage
235 before auxin treatment and 60 minutes post-treatment. *eft-3>AID::GFP* expression in the
236 AC was quantified as described above. Data were normalized by dividing the MFI in auxin
237 treated animals by the average MFI in untreated animals. To analyze *nhr-25::GFP::AID*

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238 degradation, GFP levels were quantified by measuring the MFI in individual GFP-
239 expressing nuclei in the AC/VU, AC, or VPCs subtracted by the MFI of a background
240 region in the image to account for background noise. Nuclei were outlined using the
241 threshold tool in Fiji or for animals with no detectable GFP signal, the corresponding DIC
242 image was utilized to identify the nucleus. Images of L3 larvae were captured in a *C.*
243 *elegans* larvae-specific microfluidic device (Keil *et al.* 2017). To quantify AID::GFP
244 degradation, animals were loaded into the microfluidic chamber and fed NA22 bacteria.
245 Images were captured at time points 0, 30, 60, 90, and 120 minutes with or without auxin.
246 Here, *eft-3>AID::GFP* expression was quantified by measuring the MFI in whole animals
247 subtracted by the MFI of a background region in the image to account for background
248 noise. Whole animals were outlined using the freehand selection tool in Fiji. Data were
249 normalized by dividing the MFI in treated or untreated animals at time points 30, 60, 90,
250 and 120 minutes by the average MFI in untreated animals at timepoints 30, 60, 90, and
251 120 minutes respectively to account for photobleaching from imaging the same animal.
252 Cartoons were created with BioRender (biorender.com) and ChemDraw software
253 (version: 18.0). Graphs were generated using Prism software (version: 8.1.2). Figures
254 were compiled using Adobe Photoshop (version: 20.0.6) and Illustrator (version: 23.0.26).

255

256 ***Statistical analyses***

257 A power analysis was performed to determine the sample size (*n*) needed per experiment
258 to achieve a power level of 0.80 or greater (Cohen 1992; Pollard *et al.* 2019). Statistical
259 significance was determined using either a two-tailed unpaired Student's t-test or Mann
260 Whitney U test. $P < 0.05$ was considered statistically significant. The figure legends

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261 specify when error bars represent the standard deviation (SD) or interquartile range
262 (IQR).

263

264 ***Data availability***

265 Supplemental data and key reagents can be found at Figshare. Worm strains CA1202,
266 CA1204, and PS3239 are available to order from the *Caenorhabditis* Genetics Center. All
267 other strains are available upon request. The data that support the findings of this study
268 are available upon reasonable request.

269

270 **RESULTS AND DISCUSSION**

271

272 ***NAA is a synthetic alternative to the natural auxin IAA***

273 Given the recent advances in CRISPR/Cas9-genome editing technology (Dickinson and
274 Goldstein 2016; Dokshin *et al.* 2018), the auxin-inducible degron (AID) with or without a
275 fluorescent reporter (e.g., GFP or its derivatives) can be inserted into a genomic locus of
276 interest (Röth *et al.* 2019). Though this technology can be applied with ease, there are
277 certain limitations that exist with the use of the natural auxin indole-3-acetic acid (IAA),
278 including its limited solubility in water. While levels of the ethanol solvent used to dissolve
279 IAA (0.25–1.52%) are well below the threshold for causing a physiologic response
280 (Morgan and Sedensky 1995; Kwon *et al.* 2004), higher percentages of ethanol (7%) have
281 been shown to cause rapid changes in *C. elegans* gene expression (Kwon *et al.* 2004).

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282 A potentially more problematic limitation of IAA for live-cell imaging-based applications is
283 cytotoxicity related to excitation with UV and blue light. Specifically, IAA has been shown
284 in yeast (Papagiannakis *et al.* 2017) and mouse oocytes (Camlin and Evans 2019) to
285 cause cytotoxicity, likely due to acceleration of the oxidative decarboxylation of IAA to
286 methylene-oxindole (Srivastava 2002). In yeast, IAA exposure during live-cell imaging
287 suppressed cell proliferation (Papagiannakis *et al.* 2017) and mammalian oocytes failed
288 to complete meiotic maturation (Camlin and Evans 2019). In both systems, the use of a
289 synthetic auxin, 1-naphthaleneacetic acid (NAA), rescued these cytotoxic responses. For
290 these reasons, we chose to examine whether NAA would also function in *C. elegans* to
291 degrade AID-tagged proteins in the presence of TIR1. Ultimately, we wished to evaluate
292 AID-mediated degradation (**Figure 1A**) in single cells and tissues using live-cell imaging.
293 Thus, we determined the kinetics of protein degradation using spinning disk confocal
294 microscopy, rather than using low-magnification microscopy and Western blot analysis to
295 measure protein loss in whole animals, as performed previously (Zhang *et al.* 2015). We
296 chose to focus primarily on the L3 stage of post-embryonic development due to many of
297 the dynamic cellular behaviors occurring over relatively short time scales (within minutes
298 to hours) in this developmental window, including uterine-vulval attachment and vulval
299 morphogenesis (**Figure 1B**) (Gupta *et al.* 2012).

300

301 To analyze the kinetics of AID-mediated degradation in the uterine anchor cell (AC)
302 and underlying vulva precursor cells (VPCs) (**Figure 1B**), we utilized a previously
303 published strain expressing AID::GFP and TIR1::mRuby under the same ubiquitously
304 expressed *eft-3* promoter (Zhang *et al.* 2015). The single-cell abundance of AID::GFP

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305 was measured over time in mid-L3 stage animals exposed to different concentrations of
306 auxin incorporated into standard *C. elegans* solid culture media (**Figure 2A**). In addition
307 to testing the natural auxin IAA, also tested whether it was possible to perform auxin-
308 inducible degradation in the AC and VPCs using the synthetic auxin analog NAA (**Figure**
309 **2B**). In the presence of ≥ 1 mM IAA or NAA, AID::GFP abundance in the AC and VPCs
310 was reduced by approximately 80% of its initial level within 30 minutes (**Figure 2C-D**).
311 Within 60 minutes, AID::GFP was virtually undetectable (**Figure 2C-D**). These results
312 indicate that NAA can serve as a viable substitute to IAA. These results indicate that NAA
313 can serve as a viable substitute for IAA. Similar to (Zhang *et al.* 2015), we observed that
314 growth on IAA resulted in similar brood sizes compared to control (**Table S1**). Growth on
315 the potassium salt of NAA (K-NAA) resulted in similar brood sizes compared to control,
316 and also produced a modest but significant ($P = 0.02$) reduction in embryonic lethality
317 compared to IAA treatment (**Table S1**). Consistent with this result, higher levels of toxicity
318 was observed when using IAA over NAA in studies investigating circadian rhythm biology
319 in *Drosophila* (Chen *et al.* 2018). Zhang *et al.* (2015) reported inhibited bacterial growth
320 at high concentrations of auxin. Compared to bacterial growth on IAA and NAA, we
321 observed more robust OP50 growth on K-NAA with no trade-off in degradation rate
322 (**Figure S1A**). Together, these results demonstrate that NAA is a viable alternative to IAA
323 for targeted protein degradation in *C. elegans* larvae.

324

325 ***NAA is soluble in physiological buffer***

326 To compare degradation kinetics on plate-based growth to depletion in liquid culture
327 during imaging, we measured AID::GFP abundance in the AC and VPCs in mid-L3 stage

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328 animals exposed to different concentrations of NAA solubilized in M9 buffer (**Figure 3A-**
329 **D**). In the presence of ≥ 1 mM liquid NAA, AID::GFP in the AC and VPCs was reduced by
330 80%, as compared to initial levels, within 30 minutes and was nearly undetectable within
331 60 minutes (**Figure 3B-D**). These results show that NAA can induce auxin-dependent
332 degradation in liquid culture in *C. elegans*, reducing the need to rear animals on auxin
333 plates and transfer to slides for imaging. This finding raises the possibility of depleting
334 proteins and imaging the resulting developmental consequences at single-cell resolution.

335

336 ***K-NAA is an option for C. elegans researchers employing microfluidics***

337 The ability to easily solubilize NAA in physiological buffer raises the possibility of
338 performing protein degradation experiments paired with microfluidics where individual
339 animals can be imaged over long periods at cellular resolution. *C. elegans* lifespan and
340 behavioral assays can involve, subtle phenotypes sensitive to environmental
341 perturbations. Accordingly, the low level of ethanol present in IAA plates is not optimal for
342 these types of assays. The reduced bacterial growth on ethanol and IAA could also affect
343 nutrition (**Figure S1A**) (Cabreiro *et al.* 2013), making water soluble K-NAA an attractive
344 alternative. To compare degradation kinetics between NAA and K-NAA, we time-lapsed
345 L3 stage animals using a microfluidic device optimized for long-term imaging of *C.*
346 *elegans* larvae (Keil *et al.* 2017), assessing depletion of ubiquitously expressed AID::GFP
347 in trapped animals (**Figure 3A**). At the L3 stage, animals were loaded into the microfluidic
348 chamber in M9 and flushed with a mixture of M9, 4 mM NAA or K-NAA, and NA22 *E. coli*
349 as a bacterial food source (Keil *et al.* 2017). The animals were imaged every 30 minutes
350 for 2 hours. During image acquisition, animals were temporarily immobilized by manually

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351 increasing the negative pressure on the compression layer of the device (Keil *et al.* 2017).
352 Between timepoints, animals were allowed to move and feed freely in 4 mM NAA or K-
353 NAA combined with NA22 in M9. Although degradation kinetics were slower than those
354 observed in NAA solubilized in M9 alone (**Figure 3B-D**) or NGM plates containing NAA
355 or K-NAA (**Figure S1B-C**), we still observed approximately 60–70% reduction of
356 AID::GFP expression within the first 30 minutes of NAA or K-NAA exposure, and nearly
357 80% depletion of whole animal AID::GFP within 1 hour (**Figure 3E-F**). Our results may
358 be under-representing the overall loss of AID::GFP as we did not account for gut
359 autofluorescence in our quantification of fluorescence intensity in whole animals
360 (Teuscher and Ewald 2018). Nonetheless, our results demonstrate that AID-tagged
361 proteins can be depleted in a microfluidic platform which, when combined with long-term
362 high-resolution imaging, provides a powerful tool for studying post-embryonic *C. elegans*
363 development at cellular resolution.

364

365 ***The AID system functions through specific components of the C. elegans SCF***
366 ***complex to degrade target proteins***

367 Our work demonstrates that the AID system functions rapidly to degrade target proteins
368 in *C. elegans* as described previously (Zhang *et al.* 2015). As a heterologous system,
369 researchers have shown in yeast that *Arabidopsis* TIR1 interacts with the *S. cerevisiae*
370 Cul1 homolog, Cdc53 (Nishimura *et al.* 2009). Whether *Arabidopsis* TIR1 also functions
371 through *C. elegans* proteins homologous to yeast SCF proteins is unknown. Thus, to
372 examine interactions between TIR1 and SCF complex proteins in *C. elegans*, we used
373 RNAi technology directed against components of the SCF complex and quantified

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374 AID::GFP in the presence and absence of NAA (**Figure 4 and Figures S2-S3**). This
375 experiment was designed to provide insight into the mechanism through which the AID
376 system depletes target proteins in *C. elegans* and as an intersectional proof-of-concept
377 test of combining auxin-based depletion with a RNAi feeding approach. Briefly, the SCF
378 complex consists of three components: SKP1, CUL1, and RBX1. In contrast to yeast and
379 humans, which contain only one functional SKP1 protein, the scaffold protein CUL-1 is
380 known to interact with eight of the Skp1-related adaptor proteins in *C. elegans*, including
381 SKR-1, -2, -3, -4, -7, -8, -9 and -10 (Nayak *et al.* 2002; Yamanaka *et al.* 2002). We first
382 perturbed *cul-1* expression. To deplete CUL-1, we generated a new RNAi construct
383 targeting *cul-1* in the upgraded T444T RNAi targeting vector (Sturm *et al.* 2018). Notably,
384 this vector contains T7 terminator sequences, which prevents non-specific RNA
385 fragments from being synthesized from the vector backbone (Sturm *et al.* 2018). This
386 vector modification increases the efficiency of mRNA silencing over the original L4440
387 vector (Sturm *et al.* 2018).

388

389 We hypothesized that depleting CUL-1 would strongly interfere with the
390 proteasomal machinery and thus protein turnover. To assess the abundance of AID::GFP
391 in the *C. elegans* AC, we treated animals with either *control(RNAi)* or *cul-1(RNAi)*. As
392 before, we made use of a strain expressing AID::GFP and TIR1::mRuby from an *eft-3*
393 driver (Zhang *et al.* 2015), and we examined animals at the P6.p 2-cell stage (**Figure 4A**).
394 RNAi knockdown of *cul-1* resulted in a modest but statistically significant increase in the
395 abundance of AID::GFP in the AC compared to *control(RNAi)* treatment (+19%, $n = 31$
396 and 33, respectively, $P < 0.0001$) (**Figure 4C**). This result suggests that there is TIR-

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397 dependent, auxin-independent depletion of AID::GFP, similar to reports in other systems
398 (Morawska and Ulrich 2013; Nishimura and Fukagawa 2017; Zasadzińska *et al.* 2018).
399 To further test this notion, we assessed GFP abundance in the AC in animals lacking
400 TIR1::mRuby. AID::GFP and GFP were driven by *eft-3* and *zmp-1* promoters,
401 respectively, and we performed the experiment on animals at the P6.p 2-cell stage. We
402 did not assess AID::GFP abundance in the VPCs due to the variable sensitivity of this
403 tissue to RNAi (compare **Figure 4A and S2**) (Bourdages *et al.* 2014; Matus *et al.* 2014).
404 Depletion of CUL-1 in animals expressing AID::GFP (-1.8%, $n = 29$ for both treatments,
405 $P = 0.1168$; **Figure S3A-B**) or GFP (-2.4%, $n = 20$ for both RNAi treatments, $P = 0.7682$;
406 **Figure S3C-D**) resulted in a slight decrease in GFP abundance, though it was not
407 statistically significant in either case compared to treatment with *control(RNAi)*. The
408 modest decrease in protein abundance in animals lacking TIR1::mRuby suggests that
409 knockdown of *cul-1* might mildly perturb protein homeostasis, but TIR1-mediated
410 proteosomal degradation of AID-tagged proteins independent of auxin exposure
411 requires endogenous levels of CUL-1 to function robustly.

412

413 Next, we tested whether depletion of *cul-1* would inactivate AID-mediated protein
414 degradation. We fed synchronized L1 stage animals with RNAi targeting *cul-1*, and then
415 treated animals at the P6.p 2-cell stage with 1 mM NAA for 60 minutes before quantifying
416 AID::GFP degradation in the AC (**Figure 4A**). For *control(RNAi)*-treated animals, the
417 abundance of AID::GFP in the AC was nearly undetectable within 60 minutes (-94%, $n =$
418 33; **Figure 4D**). Remarkably, the abundance of AID::GFP in the AC was reduced by only

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419 29% within 60 minutes for animals treated with *cul-1(RNAi)* ($n = 31$, $P < 0.0001$; **Figure**
420 **4D**).

421

422 We next wanted to determine if any of the Skp1-related proteins in *C. elegans*
423 function as adaptors that link CUL-1 to the F-box protein TIR1 to mediate degradation of
424 AID-tagged target proteins. Based on the availability of RNAi clones, we fed synchronized
425 L1 stage animals with RNAi targeting four of the eight Skp1-related adaptors known to
426 interact with CUL-1; *skr-1*, *skr-2*, *skr-7*, and *skr-10* (Nayak *et al.* 2002; Yamanaka *et al.*
427 2002). Owing to the 83% sequence homology between *skr-1* and *skr-2* likely stemming
428 from a gene duplication event and predicted cross-RNAi effects, their gene names are
429 unified in this report as *skr-1/2* similar to (Nayak *et al.* 2002). Of all the Skp1 homologs,
430 *C. elegans skr-1* and human Skp1 share the greatest sequence homology (Yamanaka *et*
431 *al.* 2002). We also fed animals RNAi targeting *rbx-1*, which encodes the RING finger
432 protein in the SCF E3 ubiquitin ligase (Yamanaka *et al.* 2002). To assess AID::GFP
433 abundance in the AC, we again used animals expressing *eft-3>AID::GFP* and *eft-*
434 *3>TIR1::mRuby* and examined animals at the P6.p 2-cell stage (**Figure 4B**). RNAi
435 knockdown of *skr-1/2* compared to *control(RNAi)* led to differences in AID::GFP
436 abundance that were not statistically significant ($n = 23$ and 20, respectively, $P = 0.3522$).
437 However, similar to *cul-1(RNAi)*, RNAi silencing of *rbx-1* ($n = 22$) resulted in a statistically
438 significant increase in the abundance of AID::GFP in the AC compared to *control(RNAi)*
439 treatment ($P = 0.0283$) (**Figure 4E**). Interestingly, both *skr-7(RNAi)* ($n = 23$) and *skr-*
440 *10(RNAi)* ($n = 21$) resulted in a statistically significant decrease in AID::GFP abundance
441 compared to control ($P < 0.0001$).

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442 We also wanted to determine whether depletion of *skr-1/2*, *skr-7*, *skr-10*, and *rbx-*
443 *1* could inactivate AID-mediated protein degradation. We fed synchronized L1 stage
444 animals with RNAi targeting these SCF complex components. We treated animals at the
445 P6.p 2-cell stage with 1 mM NAA for 60 minutes and quantified AID::GFP degradation in
446 the AC (**Figure 4F**). For *control(RNAi)*-treated animals, the abundance of AID::GFP in the
447 AC was once again nearly undetectable within 60 minutes (-96%, $n = 21$) (**Figure 4F**).
448 Similarly, the AID::GFP abundance in animals treated with *skr-7(RNAi)* ($n = 21$) and *skr-*
449 *10(RNAi)* ($n = 21$) was undetectable within 60 minutes of NAA exposure (-94% and -93%,
450 respectively). For animals treated with *skr-1/2(RNAi)* ($n = 21$, $P < 0.0001$), the abundance
451 of AID::GFP in the AC was reduced by 57% within 60 minutes (**Figure 4F**). For animals
452 treated with *rbx-1(RNAi)* ($n = 23$, $P < 0.0001$), the abundance of AID::GFP in the AC was
453 reduced by 65% within 60 minutes (**Figure 4F**). These results suggest that: 1)
454 suppression of *cul-1*, *skr-1/2*, or *rbx-1* is sufficient to block TIR1-mediated degradation,
455 while suppression of *skr-7* or *skr-10* is not; 2) TIR1 functions as a substrate recognition
456 component of the *C. elegans* CUL-1-based SCF complex, which was also previously
457 shown in yeast (Nishimura *et al.* 2009); and 3) it is possible to deplete multiple targets
458 simultaneously using both AID and RNAi technology.

459
460 Inhibiting the expression of *cul-1*, *skr-1/2*, or *rbx-1* is a valid approach for reversing
461 AID-mediated degradation in *C. elegans*. We suggest using *cul-1*, *skr-1/2* or *rbx-1(RNAi)*
462 for this purpose with caution, as they have known cell cycle-dependent functions and
463 therefore silencing them may conflate the recovery of AID-tagged proteins with a cell
464 cycle phenotype (Kipreos *et al.* 1996). As an alternative approach to achieving recovery

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465 of AID-tagged proteins, we propose the use of RNAi targeting TIR1 or simply using
466 auxinole, a commercially available inhibitor of TIR1 (Hayashi *et al.* 2012; Yesbolatova *et*
467 *al.* 2019). One caveat to this approach is that auxinole is expensive and thus it may be
468 difficult to obtain stoichiometrically equivalent amounts of auxin and auxinole to truly
469 achieve recovery of one's protein of interest. However, for *C. elegans* researchers
470 requiring tighter temporal control, these may be avenues worth exploring. Presently,
471 recovery from degradation with 1 mM auxin takes up to 24 hours to fully recover
472 expression of the target protein (Zhang *et al.* 2015). Such protein recovery kinetics are
473 insufficient for studying events in the nematode that occur within minutes to hours such
474 as uterine-vulval attachment, vulval morphogenesis, or many other developmental events
475 occurring post-embryonically.

476

477 ***NAA as a tool for exploring phenotypes during development and beyond***

478 As our previous results demonstrate that we could effectively deplete a non-functional
479 AID::GFP reporter expressed in the uterine AC and VPCs, we next tested whether NAA-
480 mediated depletion of target proteins could be utilized to study post-embryonic
481 developmental events occurring over a tight temporal window. We focused on a well-
482 studied system of organogenesis, *C. elegans* uterine-vulval cell specification and
483 morphogenesis (**Figure 1B**) (Schindler and Sherwood 2013). As a proof-of-principle, we
484 chose to deplete the nuclear hormone receptor, *nhr-25*, a homolog of arthropod Ftz-F1
485 and vertebrate SF-1/NR5A1 and LRH-1/NR5A2, which has been shown to function
486 pleiotropically in a wide array of developmental events, from larval molting (Asahina *et al.*
487 2000; Gissendanner and Sluder 2000; Frand *et al.* 2005), heterochrony (Hada *et al.*

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488 2010), and uterine-vulval morphogenesis (Chen *et al.* 2004; Hwang and Sternberg 2004;
489 Asahina *et al.* 2006; Hwang *et al.* 2007; Ward *et al.* 2013). It was this pleiotropy that made
490 targeting NHR-25 an attractive target, as RNAi and mutant analyses have shown
491 previously that it is initially required in the AC during the AC/VU decision for proper
492 specification of AC fate (Hwang and Sternberg 2004; Asahina *et al.* 2006) and
493 approximately 7 hours later it is required in the underlying VPCs for cell division (Chen *et*
494 *al.* 2004; Hwang *et al.* 2007; Ward *et al.* 2013).

495
496 First, we examined the *nhr-25::GFP::AID* expression pattern, and observed GFP
497 localization to the nuclei of the AC/VU cells during the mid-L2 stage, enrichment in the
498 AC following specification, and nuclear localization in the 1° and 2° VPCs during all stages
499 of vulval division and morphogenesis (**Figure 5A**). We quantified GFP fluorescence over
500 developmental time. Consistent with previous reports based on transgene analyses
501 (Gissendanner and Sluder 2000; Ward *et al.* 2013), endogenous *nhr-25::GFP::AID* AC
502 expression peaks after AC specification in the early L3 at the P6.p 1-cell stage and is
503 undetectable above background by the P6.p 4-cell stage at the time of AC invasion.
504 Conversely, *nhr-25::GFP::AID* increases in intensity in the VPCs at the P6.p 4-cell stage,
505 peaking during the morphogenetic events following AC invasion (**Figure 5B**). Given this
506 temporally-driven expression pattern and based on previous experimental results from
507 RNAi and mutant analyses (Chen *et al.* 2004; Hwang and Sternberg 2004; Asahina *et al.*
508 2006; Hwang *et al.* 2007; Ward *et al.* 2013), we hypothesized that depleting AID-tagged
509 NHR-25 prior to AC specification should interfere with the AC/VU decision. To test this
510 hypothesis, we used synchronized L1 stage animals expressing *eft-3>TIR1::mRuby* and

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511 endogenously tagged NHR-25::GFP::AID. We exposed these larvae to 4 mM NAA or a
512 buffer control and examined animals in the early L3 stage, after the normal time of AC
513 specification. Strikingly, all 36 animals examined showed a failure to specify the AC fate,
514 with the presence of either one (10/36) or two (26/36) small AC/VU-like cells in the central
515 gonad as compared to control animals (**Figure 5C**). Next, we repeated the experiment
516 but waited until after AC specification, in the early L3 stage, to expose animals to buffer
517 control or 4 mM NAA. Here, in all animals, we detected the presence of an AC situated
518 over P6.p, but in 34 of the 36 animals, P6.p failed to divide as compared to controls at
519 the mid-L3 stage (**Figure 5E**). Quantification of *nhr-25::GFP::AID* in AC/VU cells (**Figure**
520 **5D**) and VPCs (**Figure 5F**) demonstrated the 4 mM NAA treatment robustly depleted
521 endogenous protein by 95% in the AC/VU and 81% in the VPCs, respectively. Finally, we
522 waited until treated animals (early L3 stage) became adults (approximately 24 hours later)
523 and examined them for plate level phenotypes. We saw a 100% Egg-laying defect (Egl)
524 in 4 mM NAA treated animals as compared to control treated plates (**Figure 5G**).
525 Together, these results indicate that the synthetic auxin, NAA, can robustly deplete target
526 endogenous proteins in a facile, high-throughput fashion during uterine-vulval
527 development. This method should prove valuable in dissecting pleiotropic gene function
528 in the future.

529

530 ***Important caveats of the C. elegans AID system***

531 Our results reported here with a synthetic auxin, NAA, as well as the initial report
532 developing the *C. elegans* AID system (Zhang *et al.* 2015) clearly demonstrate the
533 effectiveness of auxin-induced targeted protein depletion. Several other recent reports

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534 have also effectively used the AID system in *C. elegans* to control protein function,
535 including controlling spermatogenesis by manipulating *spe-44* levels (Kasimatis *et al.*
536 2018), depleting a mediator component to modulate longevity (Lee *et al.* 2019), examining
537 chromosome segregation during oogenesis (Ferrandiz *et al.* 2018), meiotic crossover
538 (Zhang *et al.* 2018), and revealing novel roles of neuronal gene function through
539 conditional depletion (Serrano-Saiz *et al.* 2018). Despite the increasing frequency of AID
540 system usage in the *C. elegans* community, there are only a handful of TIR1 driver lines
541 published, and the importance of copy number and promoter strength has not been
542 systematically assessed.

543

544 While we are optimistic that the use of the synthetic analog of auxin presented
545 here will allow for even more widespread utility of the AID system in the *C. elegans*
546 research community, there are still are some areas open to improvement for the
547 technology. A recent report in mammalian cell culture identified that AID-tagged proteins
548 are depleted in an auxin-independent fashion in the presence of TIR1, relative to wild-
549 type levels (Li *et al.* 2019; Sathyan *et al.* 2019). We examined if this was also occurring
550 in *C. elegans* strains in our laboratory expressing AID-tagged proteins and TIR1. We were
551 able to detect statistically significant auxin-independent depletion of both a ubiquitously
552 expressed AID::GFP transgene under the *eft-3* promoter in ACs (-22%, $n = 24$, $P <$
553 0.0001) and VPCs (-24%, $n = 24$, $P < 0.0001$; **Figure S4A-B**) and an endogenously
554 tagged *nhr-25::GFP::AID* allele in ACs (-22%, $n = 26$, $P = 0.0022$) and VPCs (-35%, $n =$
555 26, $P < 0.0001$; **Figure S4C-D**). As partial loss of an endogenous protein could generate
556 a hypomorphic condition, both from the placement of the AID tag and apparent triggering

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557 of the degradation machinery, we urge caution in carefully evaluating AID-tagged alleles
558 paired with TIR1, independent of auxin delivery. Further optimization of the AID system
559 in *C. elegans* will hopefully ameliorate this concern, as researchers recently used the
560 heterologous co-expression of an auxin response factor (ARF) with TIR1 to rescue auxin-
561 independent degradation in mammalian cell culture (Sathyan *et al.* 2019).

562

563 **Conclusion**

564 The ease of editing the *C. elegans* genome using CRISPR/Cas9-based approaches
565 (Calarco and Friedland 2015) and heterologous gene manipulation tools is ushering in a
566 new era of cellular and developmental biology. Several new tools available to *C. elegans*
567 researchers require the insertion of small amino acid tags into target loci, including ZF1
568 tagging (Armenti *et al.* 2014), sortase A (Wu *et al.* 2017), and the AID system (Zhang *et*
569 *al.* 2015). Alternatively, any GFP fusion can be targeted via a GFP nanobody tethered to
570 ZIF1 (Wang *et al.* 2017). These genomic edits are then paired with single transgene
571 expression to allow for targeted spatial and temporal loss-of-function approaches through
572 manipulation of endogenous loci. Prior to their advent, spatial and temporal control of
573 protein function was largely missing from the *C. elegans* genomic toolkit. With an ever-
574 increasing set of these tools being optimized for *C. elegans*, it is clear that different tools
575 will have strengths and weaknesses depending on multiple variables, including
576 subcellular localization of target protein, availability of tissue- and cell-type specific
577 drivers, and inducibility of depletion. Here, we optimize a powerful heterologous system,
578 the auxin-inducible degradation system. We demonstrate that a synthetic auxin analog,
579 NAA, and its water-soluble, potassium salt, K-NAA, can function equivalently to natural

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580 auxin. The water solubility permits easier preparation of media and allows researchers to
581 perform experiments in liquid culture and microfluidics. Importantly, the use of ethanol
582 free K-NAA may be beneficial to *C. elegans* researchers studying behavior and aging,
583 where introduction of ethanol may lead to confounding results. We also demonstrate the
584 strength of the AID system for studying developmental cell biology by examining multiple
585 spatial and temporal roles of the Ftz-F1 homolog *nhr-25* during uterine and vulval
586 morphogenesis. It is our hope that the use of the synthetic auxin NAA will complement
587 the AID system in *C. elegans* when examining targeted protein depletion phenotypes in
588 tissues and developmental stages of interest. As the library of tissue-specific TIR1 drivers
589 continues to grow, we envision researchers being able to rapidly degrade proteins of
590 interest in specific tissues and visualize the outcome at single-cell resolution.

591

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611

612 **AUTHOR CONTRIBUTIONS**

613 M.A.Q.M. and D.Q.M conceived and designed the experiments. G.A. and J.D.W.
614 designed the constructs. J.M.R. performed the microinjections. G.A. performed the
615 crosses and characterized strains. M.A.Q.M, B.A.K., T.N.M., L.J. and J.A. performed the
616 experiments. M.A.Q.M. and D.Q.M. analyzed and quantified the data. M.A.Q.M. and
617 D.Q.M. wrote the manuscript with contributions from the other authors. The authors
618 declare no competing interests.

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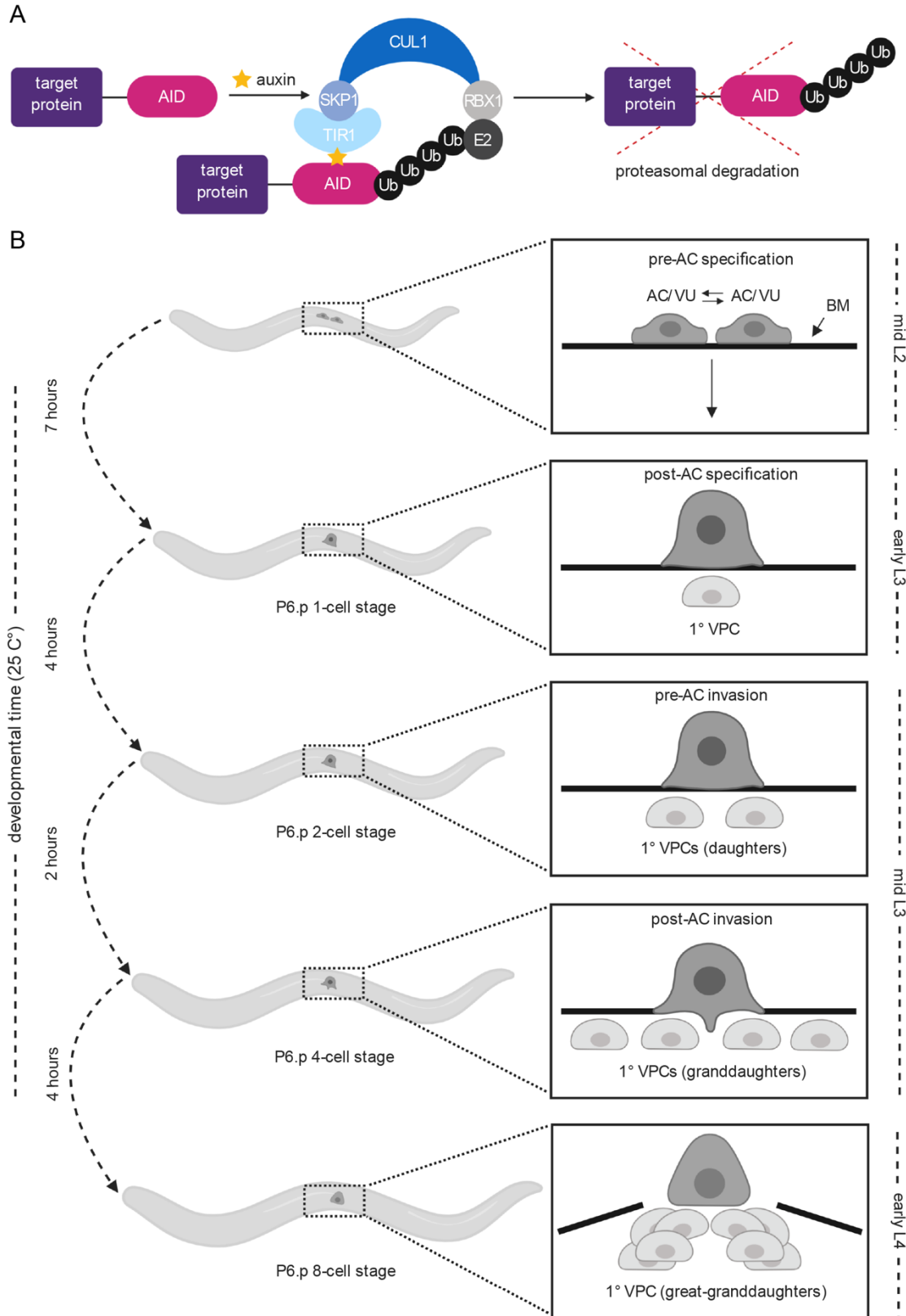
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Figure 1. Overview of the auxin-inducible degradation system and *C. elegans* uterine-vulval development. (A) In this system, a target protein is fused to an auxin-inducible degron (AID). Heterologous expression of *Arabidopsis* TIR1 mediates robust auxin-dependent proteasomal degradation of AID-tagged proteins through the SKP1-CUL1-F-box (SCF) E3 ubiquitin ligase complex. (B) Schematic of uterine-vulval morphogenesis during *C. elegans* larval development. In *C. elegans*, AC specification and morphogenesis of uterine-vulval attachment occurs from the mid-L2 through the early L4 stage (Schindler and Sherwood 2013). The AC is specified in a stochastic reciprocal Notch-Delta signaling event in the mid-L2 stage (top panel). Following AC specification, the AC specifies the 1° fate of the underlying vulval precursor cell, P6.p in the early L3 (second panel), which then divides three times to ultimately give rise to eight of the 22 cells of the adult vulva (bottom three panels).

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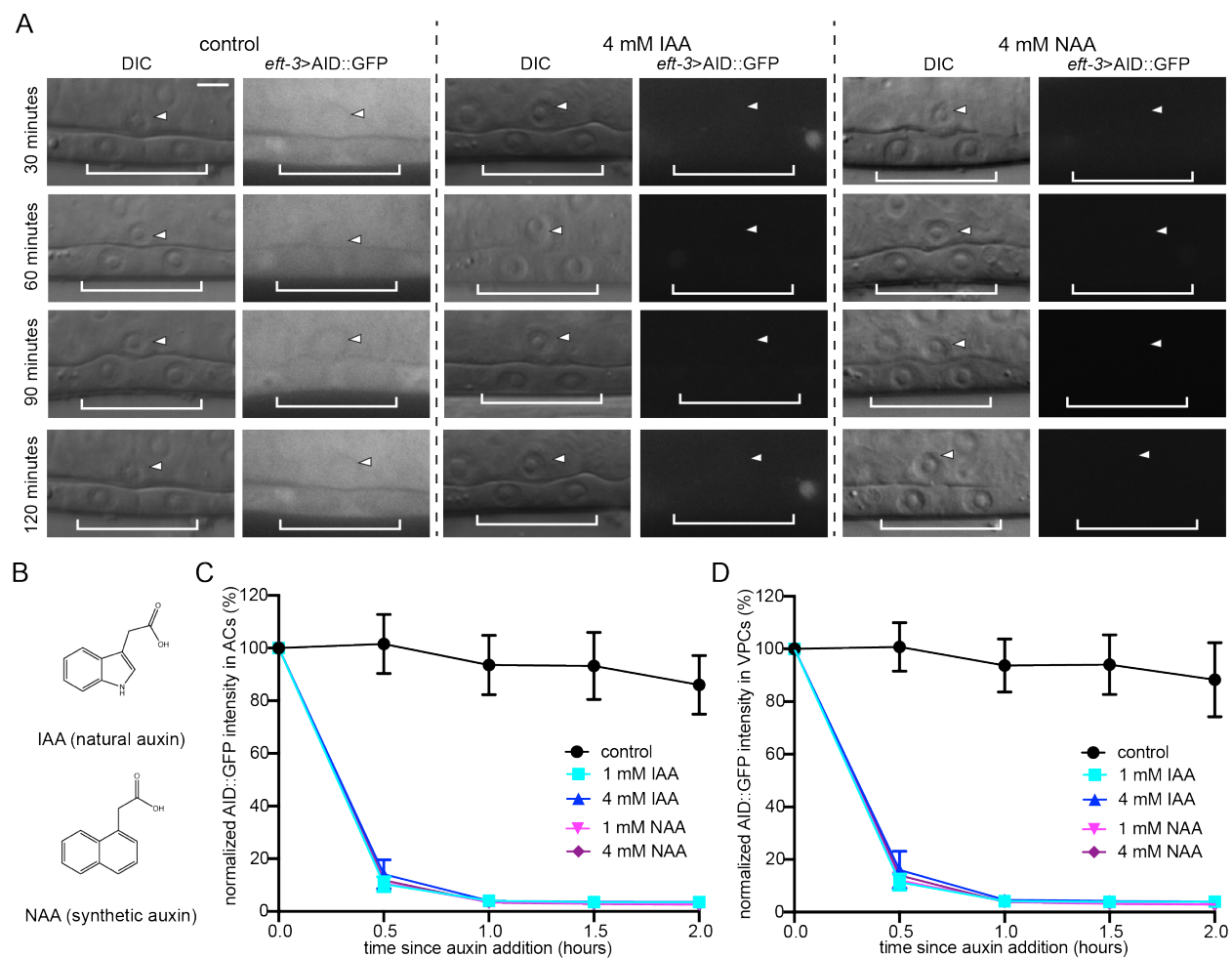
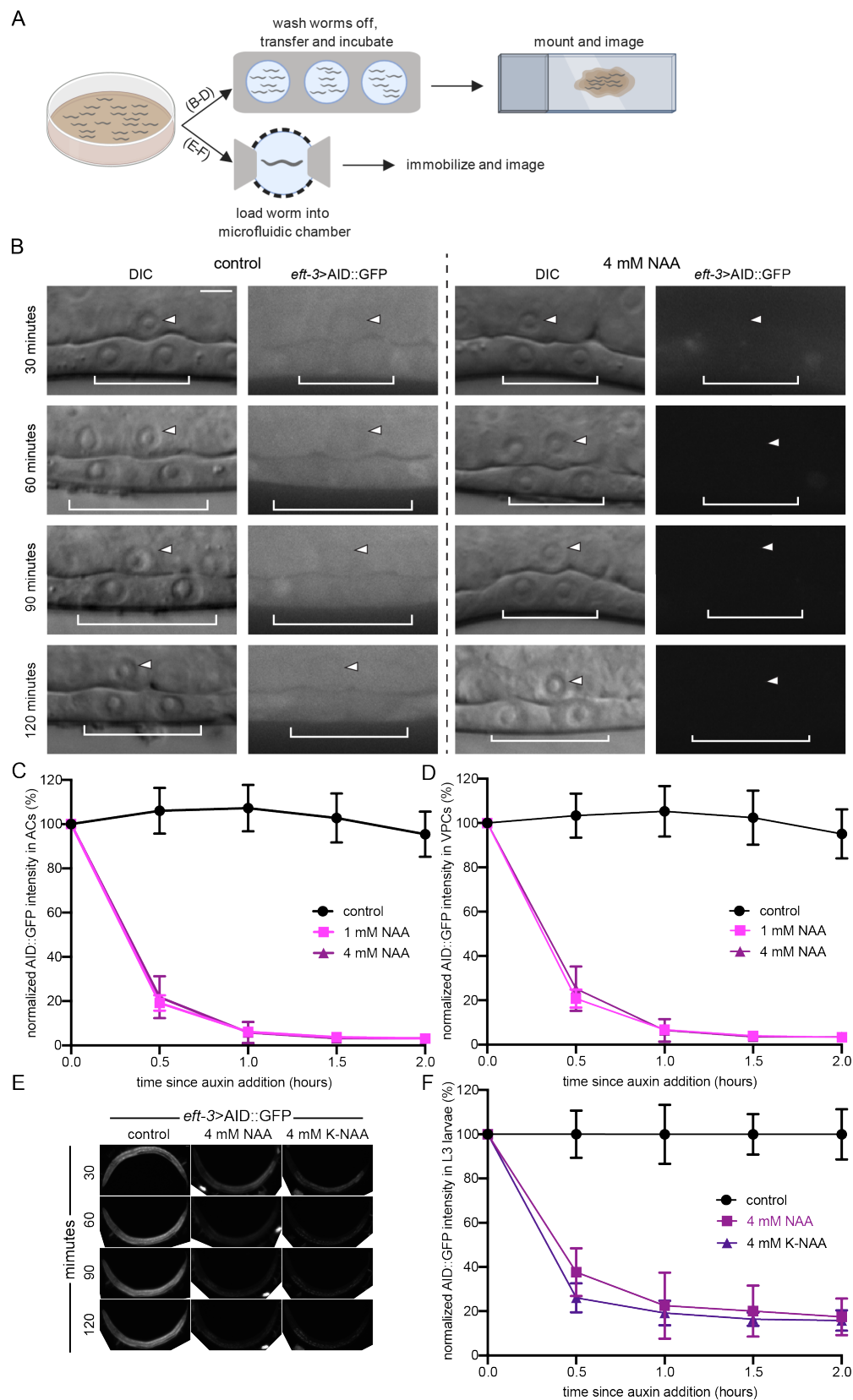


Figure 2. Comparison of IAA- and NAA-mediated degradation in the *C. elegans* AC and VPCs. (A) DIC and corresponding GFP images of ACs (arrowheads) and underlying 1° fated VPCs (brackets) from mid-L3 stage animals at the P6.p 2-cell stage. Animals expressing AID::GFP and TIR1::mRuby under the same *eft-3* promoter were treated with natural auxin indole-3-acetic acid (IAA) and synthetic auxin 1-naphthaleneacetic acid (NAA) in NGM agar containing OP50. (B) Chemical structure of IAA and NAA. (C, D) Rates of degradation determined by quantifying AID::GFP in (C) ACs and (D) VPCs

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following auxin treatment. Data presented as the mean \pm SD ($n \geq 30$ animals examined for each time point).

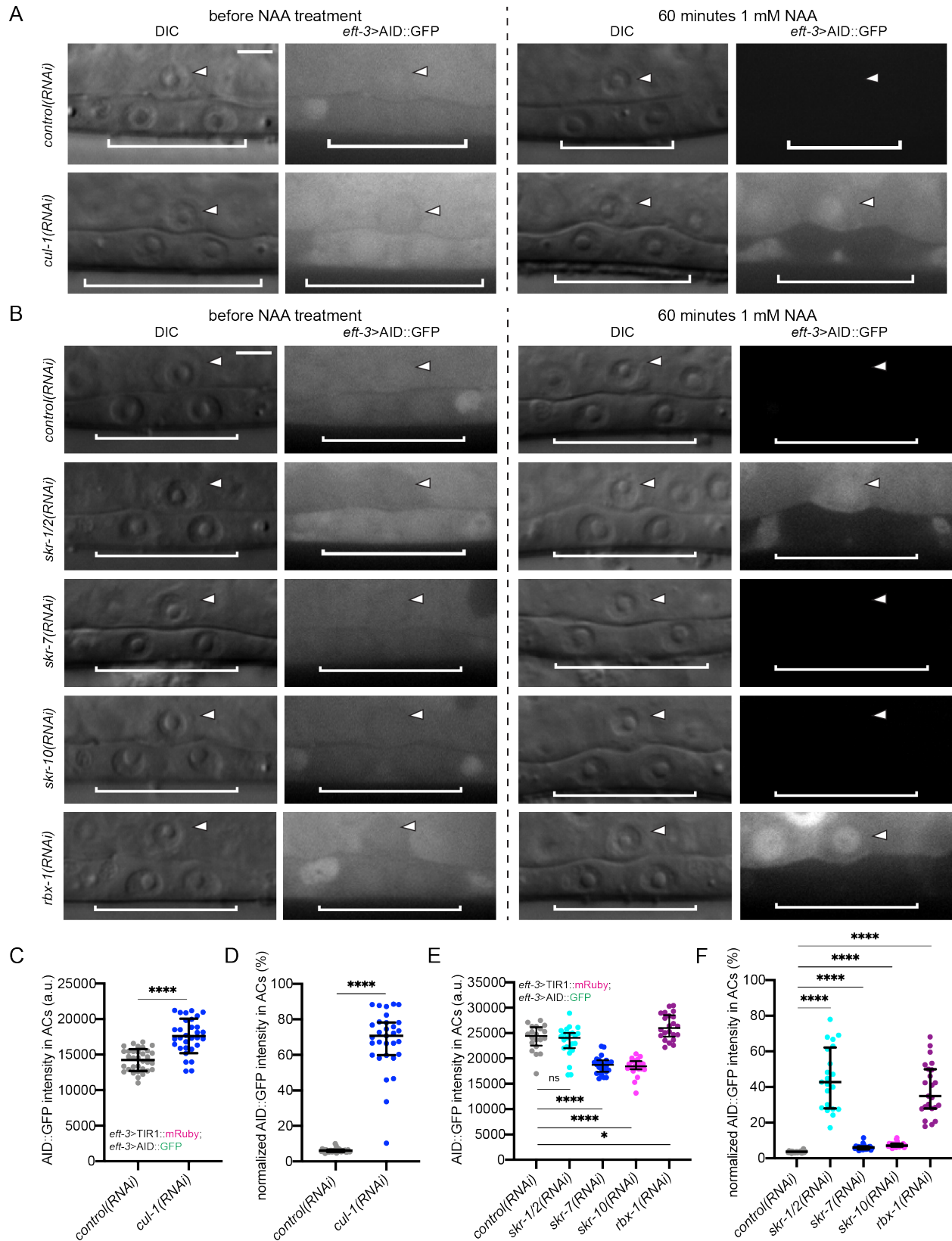
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Figure 3. Solubility of NAA/K-NAA in physiological buffer enhances utility. (A) Schematic representation of the liquid NAA-based degradation protocol for use in high-resolution microscopy or microfluidics-based approaches. (B) DIC and corresponding GFP images of ACs (arrowheads) and underlying VPCs (brackets) from mid-L3 stage animals at the P6.p 2-cell stage. Animals expressing AID::GFP and TIR1::mRuby under the same *eft-3* promoter were treated with NAA in M9. (C, D) Rates of degradation were determined by quantifying AID::GFP in (C) ACs and (D) VPCs following auxin treatment. Data presented as the mean \pm SD ($n \geq 30$ animals examined for each time point). (E) Images of AID::GFP expression from mid-L3 stage animals in control conditions (M9 buffer containing NA22 only, left) or conditions where a 4 mM NAA (middle) or K-NAA (right) solution in M9 buffer containing NA22 was perfused through the microfluidic chamber for the time indicated (Keil et al. 2017). Anterior is left and ventral is down. (F) Rates of degradation were determined by quantifying AID::GFP in whole animals following auxin treatment. Data presented as the mean \pm SD ($n \geq 4$ animals examined for each time point).

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Figure 4. Suppression of SCF complex member expression inhibits TIR1-dependent degradation in the *C. elegans* AC. (A-B) DIC and corresponding GFP images of ACs (arrowheads) and underlying VPCs (brackets) from mid-L3 stage animals at the P6.p 2-cell stage. Animals expressing AID::GFP and TIR1::mRuby under the same *eft-3* promoter were treated with (A) *cul-1(RNAi)* and (B) *skr-1/2*, *skr-7*, *skr-10* and *rbx-1(RNAi)*. (C) Quantification of AID::GFP in ACs following *cul-1(RNAi)* treatment. Data presented as the mean±SD ($n \geq 30$ animals examined for each, and $P < 0.0001$ by a Student's t-test). (D) Quantification of AID::GFP in ACs following treatment with NAA. Data presented as the median+IQR ($n \geq 30$ animals examined for each, and $P < 0.0001$ by a Mann Whitney U test). (E) Quantification of AID::GFP in ACs following RNAi knockdown of Skp1-related (*skr*) genes and *rbx-1*. Data presented as the median+IQR ($n \geq 20$ animals examined for each, and P values examined by a Mann Whitney U test). (F) Quantification of AID::GFP in ACs following treatment with NAA. Data presented as the median+IQR ($n \geq 21$ animals examined for each, and P values examined by a Mann Whitney U test).

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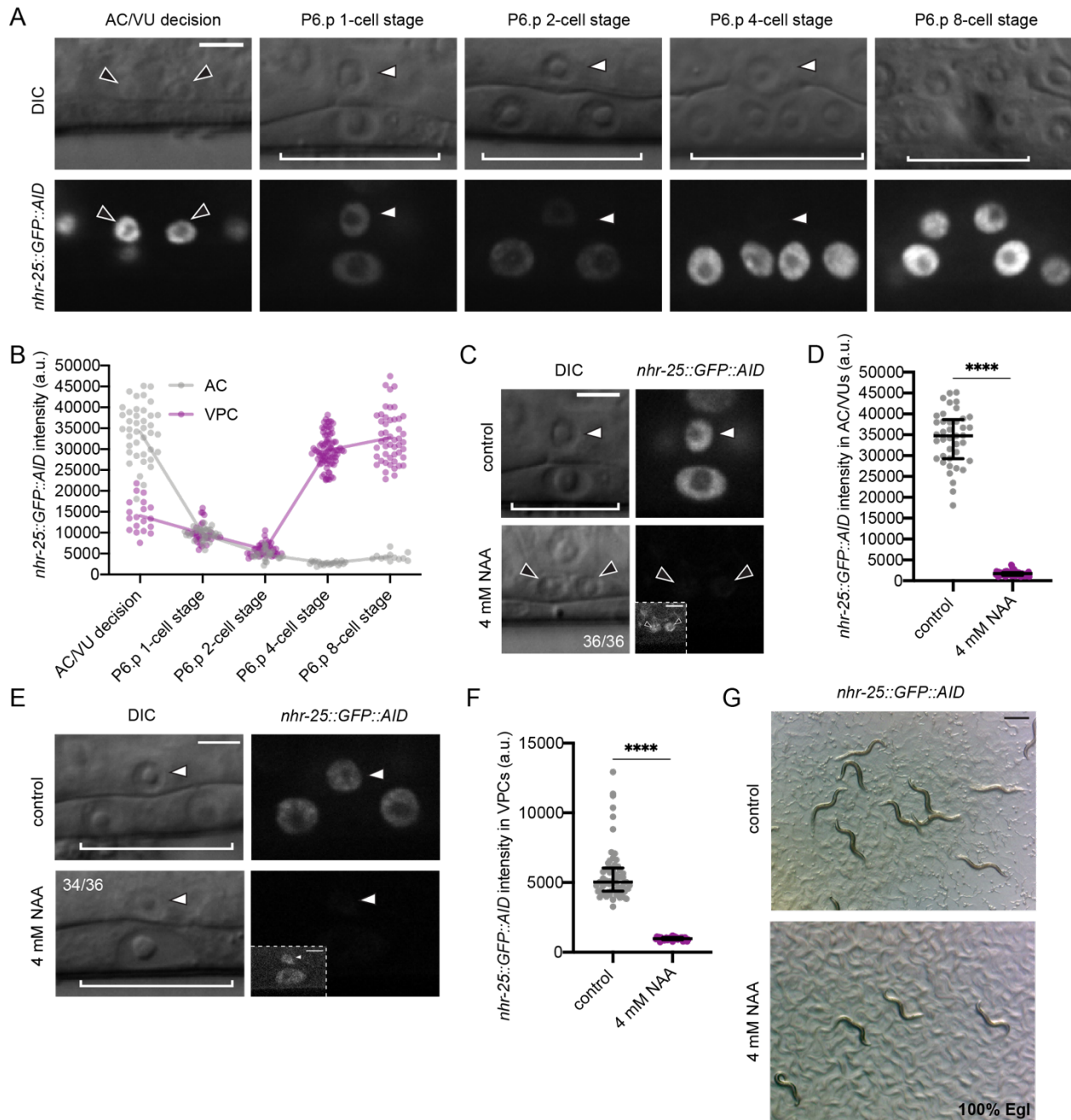


Figure 5. NAA-mediated degradation of NHR-25 causes AC specification and VPC division defects. (A) *nhr-25::GFP::AID* localizes to the nuclei of the AC/VU (black arrowheads), the AC (white arrowheads) and VPCs (brackets). At P6.p 8-cell stage (far right) the AC is not in the same focal plane as the 1° VPCs. (B) Quantification of *nhr-*

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25::*GFP*::*AID* over developmental time, from the AC/VU decision to the P6.p 8-cell stage. The curve is connected by the mean at each developmental stage ($n = 20, 31, 20, 21,$ and 12 animals quantified, respectively). (C) DIC and corresponding GFP images of ACs (arrowheads) and underlying VPCs (brackets) from early L3 stage animals. Animals expressing *nhr-25*::*AID*::*GFP* and *eft-3*>*TIR1*::mRuby were treated with control and 4 mM NAA. (D) Quantification of *nhr-25*::*GFP*::*AID* in AC/VUs following NAA treatment. Data presented as the median+IQR ($n \geq 20$ animals examined for each, and $P < 0.0001$ by a Mann Whitney U test). (E) DIC and corresponding GFP images of ACs (arrowheads) and underlying VPCs (brackets) from mid-L3 stage animals. Animals expressing *nhr-25*::*AID*::*GFP* and *eft-3*>*TIR1*::mRuby were treated with control and 4 mM NAA. (F) Quantification of *nhr-25*::*GFP*::*AID* in VPCs following NAA treatment. Data presented as the median+IQR ($n \geq 30$ animals examined for each, and $P < 0.0001$ by a Mann Whitney U test). (G) Representative images of adult plate level phenotypes following control and 4 mM NAA treatments added at the L3 stage ($n \geq 30$ animals examined). Scale bar in (G), 500 μm .