

1                   **Integrity of the short arm of nuclear pore Y-complex**  
2                   **is required for mouse embryonic stem cell growth**  
3                   **and differentiation**

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6   **Running title:**      Roles of Y-complex Nups in mESCs

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22

23   **Keywords:**

24   Nucleoporin, Seh1, Nup43, Nup85, Mios, mouse embryonic stem cells.

25 **Summary statement**

26 Seh1 and Nup43, although dispensable in pluripotent mouse embryonic stem cells ,  
27 are required for normal cell growth, viability upon differentiation, and maintenance  
28 of proper NPC density.

29

30 **Abstract**

31 Many cellular processes, ranging from cell division to differentiation, are controlled  
32 by nuclear pore complexes (NPCs). However studying contributions of individual NPC  
33 subunits to these processes in vertebrates has long been impeded by their  
34 complexity and the lack of efficient genetic tools. Here we use genome editing in  
35 mouse embryonic stem cells (mESCs) to characterize the role of NPC structural  
36 components, focusing on the short arm of the Y-complex that comprises Nup85,  
37 Seh1 and Nup43. We show that Seh1 and Nup43, although dispensable in  
38 pluripotent mESCs, are required for their normal cell growth rates, their viability  
39 upon differentiation, and for the maintenance of proper NPC density. mESCs with an  
40 N-terminally truncated Nup85 mutation (in which interaction with Seh1 is greatly  
41 impaired) feature a similar reduction of NPC density. However, their proliferation  
42 and differentiation are unaltered, indicating that it is the integrity of the Y-complex,  
43 rather than the number of NPCs, that is critical to ensure these processes.

44

45 **List of Symbols and Abbreviations used.**

46 mAID: mini Auxin Inducible Degron; mESC: mouse embryonic stem cell ; NE: nuclear  
47 envelope; NPC: Nuclear pore complex; Nups: nucleoporins ; n.s.: not significant; SD:  
48 standard deviation; *Tigre* locus: *Tightly regulated* locus ; WT: Wild type.

49

49 **INTRODUCTION**

50 Nuclear pore complexes (NPCs) are huge structures embedded in the nuclear  
51 envelope (NE). They provide the sole gateways for bidirectional nucleocytoplasmic  
52 transport, but also participate in a wide variety of other cellular processes including  
53 cell division and gene regulation (reviewed in Buchwalter et al., 2019; Hezwani and  
54 Fahrenkrog, 2017). NPCs are composed of ~ 30 distinct proteins (called nucleoporins  
55 or Nups), each present in multiple copies and forming a ring with an eightfold  
56 rotational symmetry. Among them, structural Nups assemble to form a scaffold that  
57 anchors Nups with unfolded domains, cytoplasmic filaments and the nuclear basket  
58 (reviewed in Hampoelz et al., 2019; Lin and Haelz, 2019).

59 The three-dimensional organization of the NPC scaffold has been determined at  
60 atomic resolution (reviewed in (Hampoelz et al., 2019; Lin and Haelz, 2019)). It is  
61 formed by an inner rim sandwiched by two outer (cytoplasmic and nuclear) rims,  
62 whose main component is the evolutionarily-conserved Y-complex. In metazoans,  
63 this complex (also named Nup107-160 complex) comprises Nup133, Nup107, Nup96  
64 and Sec13 (forming the stem of the Y); Nup160, Nup37 and Elys (building the long  
65 arm); and Nup85, Seh1 (also named Seh1l) and Nup43 (forming the short arm)(**Fig.**  
66 **1A**) (Loiodice et al., 2004; Rasala et al., 2006; von Appen et al., 2015).

67 Functional studies in vertebrates have shown that the Y-complex is critical for NPC  
68 assembly, both after mitosis and during interphase (Doucet et al., 2010; Harel et al.,  
69 2003; Walther et al., 2003). Studies in mammalian cells also showed that in mitosis a  
70 fraction of the Y-complex localizes at kinetochores (Loiodice et al., 2004; Rasala et  
71 al., 2006) where it is required for proper chromosome congression and segregation  
72 (Platani et al., 2009; Zuccolo et al., 2007). Because the members of the Y-complex (Y-  
73 Nups) are tightly associated throughout the cell cycle (Loiodice et al., 2004; Rabut et  
74 al., 2004), they were long anticipated to work as an entity. However, one of its  
75 components, Sec13, is also part of the COPII coat complex involved in vesicle  
76 budding (Salama et al., 1993). In addition, Sec13 and Seh1 also belong to the  
77 unrelated GATOR2 complex, an indirect regulator of the mTORC1 pathway that  
78 controls cell growth and proliferation (Bar-Peled et al., 2013), further complicating  
79 the study of their function in the context of the Y-complex.

80 In mice, inactivation of most Y-Nups genes (namely *Elys*, *Nup96*, *Nup133*, *Nup85*,  
81 *Sec13* and *Seh1*, but not *Nup37*) lead to embryonic lethality (Faria et al., 2006; Liu et  
82 al., 2019; Lupu et al., 2008; Moreira et al., 2015; Okita et al., 2004; Terashima et al.,  
83 2020; <https://www.mousephenotype.org/data/genes/MGI:1919964>). In particular,  
84 *Nup133* was found to be essential for mouse development beyond gastrulation  
85 (Lupu et al., 2008). Studies performed in mouse embryonic stem cell (mESCs)  
86 showed that *Nup133* is dispensable for cell growth at the pluripotent stage, but is  
87 required for mESC differentiation (Lupu et al., 2008). In mESCs, *Nup133* is  
88 dispensable for NPC scaffold assembly but required for the proper assembly of the  
89 nuclear pore basket (Souquet et al., 2018). However, it is not clear if the role of  
90 *Nup133* in NPC basket assembly underlies its functions in cell differentiation.

91 More recently, *Seh1*, which is critical for proper mitotic progression in cancer cell  
92 lines (Platani et al., 2018; Platani et al., 2009; Zuccolo et al., 2007), was found to be  
93 required for differentiation of oligodendrocyte progenitors (Liu et al., 2019).  
94 However, the potential contribution of *Seh1* to cell cycle progression in non-  
95 transformed cells and at other stages of cell differentiation needed to be addressed.

96 Here we assessed the requirements for *Seh1* in pluripotent mESCs and upon their  
97 differentiation towards neuroectodermal lineage, determined whether these  
98 requirements reflect its role in the GATOR2-complex or in the short arm of the Y-  
99 complex, and further addressed the specific function of these proteins in NPC  
100 integrity. This systematic analysis enabled us to disentangle the processes underlying  
101 the contribution of these of these Y-Nups in NPC assembly, nuclear size, cell growth  
102 and differentiation.

103

103 **Results**

104 **Seh1 is required for mESC growth and survival upon differentiation**

105 Using CRISPR/Cas9 gene-editing technology we obtained several independent *Seh1*<sup>-/-</sup>  
106 mESC clones (of which three were further examined in this study; see Materials and  
107 Methods and Table S2) (**Fig. 1B**). This indicates that *Seh1* is dispensable for mESC  
108 viability at the pluripotent stage. We noticed however that *Seh1*<sup>-/-</sup> mESCs formed  
109 smaller colonies than did *WT* mESCs (**Fig. 1C**). Consistently, automated cell growth  
110 analyses of *Seh1*<sup>-/-</sup> mESCs showed a clear reduction of cell confluence compared to  
111 *WT* (**Fig. 1D**). More strikingly, *Seh1*<sup>-/-</sup> cells showed a strong impairment in viability  
112 from the very early stages of monolayer differentiation towards neuroectodermal  
113 lineage and almost no cells were recovered after 5 days (**Fig. 1E and Movies S1-S2**).

114 To verify the specificity of these phenotypes, we next integrated at the permissive  
115 *Tigre* locus (Tightly regulated; Zeng et al., 2008) of *Seh1*<sup>-/-</sup> mESCs a GFP-tagged *Seh1*  
116 cDNA expressed under the control of the pCAG promoter. The resulting cell lines  
117 (subsequently named “*Rescue*” #1 and #2) expressed GFP-*Seh1* at a level comparable  
118 to that of the endogenous untagged protein (**Fig. 1B**). We observed a specific  
119 enrichment of GFP-*Seh1* at nuclear pores in interphase and at kinetochores  
120 throughout mitosis (**Fig. 1F**). Most importantly, the growth rate of the *Rescue* cell  
121 lines was comparable to that of *WT* cells both at the pluripotent stage (**Fig. 1D**) and  
122 upon neuroectodermal differentiation (**Fig. 1E and Movie S3**).

123 To exclude the possibility that phenotypes observed in *Seh1*<sup>-/-</sup> mESCs at the  
124 pluripotent stage could be due to cell adaptation, we also generated cell lines in  
125 which endogenous *Seh1* was N-terminally tagged with the 7 kDa mini Auxin Inducible  
126 Degron (mAID) sequence, to induce its acute degradation upon auxin addition  
127 (Natsume et al., 2016). A GFP tag was also introduced to allow visualization of both  
128 the localization and degradation of the resulting GFP-mAID-*Seh1* fusion. Upon  
129 addition of auxin to *GFP-mAID-Seh1* mESCs, the GFP signal rapidly declined in mitotic  
130 cells whereas, as also previously observed in HCT116 cells (Platani et al., 2018), the  
131 decay was more progressive in interphasic cells (**Fig. S1 A-D**). While the *GFP-mAID-*  
132 *Seh1* clones showed normal cell growth and differentiation properties in control

133 conditions, addition of auxin recapitulated both the cell growth and differentiation  
134 defects observed in *Seh1*<sup>-/-</sup> mESCs (**Fig. S1 E, F**).

135

136 Together these data reveal that the lack of *Seh1* specifically causes an impaired cell  
137 growth of mESCs at the pluripotent stage and drastically reduced viability upon  
138 induction of neuroectodermal differentiation.

139

140 **The altered growth rate of pluripotent *Seh1*<sup>-/-</sup> mESCs is mainly caused by extended  
141 interphases.**

142 *Seh1* is known to play a role in mitosis in cancer cell lines, in which its depletion  
143 causes a delay in mitotic progression associated with chromosome congression and  
144 segregation defects (Platani et al., 2018; Platani et al., 2009; Zuccolo et al., 2007).  
145 Whether these defects are caused by the mislocalization of the entire Y-complex  
146 from kinetochores, as observed in HeLa cells (Platani et al., 2009; Zuccolo et al.,  
147 2007) or by the removal of *Seh1* alone has recently been questioned (Platani et al.,  
148 2018). To study the mitotic role of *Seh1* in mESCs, *WT* and *Seh1*<sup>-/-</sup> cells were  
149 transfected with GFP-H2B and imaged for 4-6 hours. Quantification of progression  
150 time from prometaphase to anaphase onset showed a ~10 min delay in *Seh1*<sup>-/-</sup> as  
151 compared to *WT* mESCs (from  $23.7 \pm 10.1$  min in *WT* to  $32.8 \pm 14.5$  min in *Seh1*<sup>-/-</sup>  
152 cells; mean $\pm$ SD) (Fig. S2 A). This delay is clearly milder than the one initially reported  
153 upon RNAi-induced depletion of *Seh1* in HeLa cells (~45 to 60 min; Platani et al.,  
154 2009; Zuccolo et al., 2007) but comparable to the delay recently measured upon  
155 conditionally-induced degradation of *Seh1* in a HCT116-derived cell line (~12 min;  
156 Platani et al., 2018). In *Seh1*<sup>-/-</sup> or auxin-treated *GFP-mAID-Seh1* mitotic mESCs, the Y-  
157 complex (visualized by Nup133 and Nup85) was still properly localized at  
158 kinetochores despite the complete lack of *Seh1* (**Figs. S1C** and **S2 B,C**). This indicates  
159 that the mitotic delay observed in *Seh1*-deficient mESCs is not merely caused by the  
160 mislocalization of the Y-complex from kinetochores.

161 The 10-minute prolongation of mitosis was however unlikely to explain the cell  
162 growth defect of *Seh1*<sup>-/-</sup> mESCs (**Fig. 1C,D**). We therefore also measured the length of  
163 interphase by imaging mCherry-H2B-expressing mESCs during 24-30 hours.

164 Quantification of progression time from the end of one mitosis (set at anaphase  
165 onset) to the beginning of the next (set at prometaphase) showed that interphase  
166 length is significantly longer in *Seh1*<sup>-/-</sup> as compared to *WT* mESCs (9.4 ± 2.2 hours in  
167 *WT* versus 14.0 ± 4.5 and 14.1 ± 2.1 hours in *Seh1*<sup>-/-</sup> #1 and #2, respectively; means ±  
168 SD) (**Fig. 2A**).

169 To determine if the lengthening of interphase in *Seh1*<sup>-/-</sup> mESCs was caused by  
170 retention in a specific phase of the cell cycle, we analyzed EdU-labelled and DAPI-  
171 stained *WT* and *Seh1*<sup>-/-</sup> mESCs by imaging flow cytometry (**Figs. 2B** and **S2D**). Except  
172 for a mild increase in the percentage of the mitotic fraction, this analysis revealed a  
173 comparable distribution of the G1, S and G2 phases of the cell cycle between *Seh1*<sup>-/-</sup>  
174 and *WT* mESCs (**Fig. 1H**). Therefore, the altered growth rate of *Seh1*<sup>-/-</sup> mESCs reflects  
175 a lengthening distributed over all phases of the cell cycle.

176

### 177 **Lack of Seh1 leads to a decrease of both NPC density and nuclear size**

178 The viability of *Seh1*<sup>-/-</sup> mESCs at the pluripotent stage and their impaired survival  
179 upon differentiation was reminiscent of the phenotype observed upon inactivation  
180 of Nup133, another member of the Y-complex (Lupu et al., 2008). Because Nup133  
181 loss was recently demonstrated to affect NPC basket assembly (as revealed by lack of  
182 TPR staining in about 50% of the NPCs) (Souquet et al., 2018), we decided to  
183 examine the impact of *Seh1* inactivation on NPC assembly. We therefore quantified  
184 the average fluorescence intensity at the nuclear envelope (NE) of TPR, Nup98 and  
185 Nup133 in *WT* and *Seh1*<sup>-/-</sup> mESCs, using a GFP-tagged cell line for internal reference,  
186 as previously reported (Souquet et al., 2018; see Materials and Methods). This  
187 analysis revealed a mild but significant reduction (in the range of 20-35%) of the  
188 signal of these three nucleoporins in *Seh1*<sup>-/-</sup> relative to *WT* mESCs (**Fig. 3 A-C**). The  
189 fact that the reduced intensity at the NE is not restricted to TPR indicates that, unlike  
190 what was previously observed in *Nup133*<sup>-/-</sup> mESCs, the lack of Seh1 leads to a  
191 decrease in the total number of NPCs rather than alteration of a specific  
192 substructure. This defect in NPC density was also observed upon auxin-induced  
193 depletion of Seh1 (**Fig. 3D**) and was largely rescued by stable expression of GFP-Seh1  
194 (**Fig. 3, A-C**).

195 It was recently proposed that nuclear size is sensitive to NPC density and nuclear  
196 import capacity in cultured mammalian cells (Jevtić et al., 2019). The decreased NPC  
197 density observed upon *Seh1* inactivation thus prompted us to measure nuclear size  
198 in these mutant mESCs. This analysis revealed a ~10% reduction of the nuclear  
199 surface in *Seh1*<sup>-/-</sup> mESCs, a phenotype that was rescued by the GFP-*Seh1* transgene  
200 (**Fig. 4A**). A significant reduction in nuclear size could also be observed in auxin-  
201 treated *GFP-mAID-Seh1* mESCs (**Fig. 4B**).

202 *Seh1*-deficient mESCs thus exhibit several distinct phenotypes: altered cell growth,  
203 lethality upon differentiation, reduced NPC density and nuclear size. We next aimed  
204 to determine whether these defects were linked to each other and whether they  
205 reflected functions of *Seh1* as part of the Y-complex or the GATOR2 complex, or  
206 both.

207

#### 208 **Mios is not required for proper cell growth and cell differentiation in mESCs.**

209 Within the GATOR2 complex, *Seh1* directly interacts with Mios (also known as Mio,  
210 missing oocyte in *Drosophila* and *Sea4* in budding yeast) (Senger et al., 2011; Bar-  
211 Peled et al., 2013; Algret et al., 2014). Our western blot analyses revealed decreased  
212 protein levels of Mios in *Seh1*<sup>-/-</sup> compared to *WT* mESCs (**Fig. 5A**), a result consistent  
213 with studies in other species and cell types (Platani et al., 2018; Platani et al., 2015;  
214 Senger et al., 2011). To assess if this reduction in Mios could cause the cell growth  
215 and differentiation phenotypes observed in *Seh1*<sup>-/-</sup> mESCs, we inactivated *Mios* in  
216 mESCs via CRISPR/Cas9 (**Fig. 5A** and **Table S2**). Immunoprecipitation experiments  
217 performed using anti-*Seh1* antibodies revealed that lack of Mios prevents *Seh1*  
218 interaction with *Wdr24*, another GATOR2 complex component (Bar-Peled et al.,  
219 2013) (**Fig. 5B**). This points to Mios as being the main direct partner linking *Seh1* to  
220 the rest of the GATOR2 complex, a result complementing data previously obtained in  
221 budding yeast and *drosophila* (Algret et al., 2014; Dokudovskaya and Rout, 2015; Cai  
222 et al., 2016). Analyses of independent *Mios*<sup>-/-</sup> clones revealed only a minor reduction  
223 of cell growth at the pluripotent stage as compared to *WT* mESCs (14±15% decrease  
224 in confluence after 48h of growth, while the reduction was 44±14% for *Seh1*<sup>-/-</sup>  
225 mESCs; **Fig. 5C**). In addition, *Mios*<sup>-/-</sup> cells underwent differentiation towards the

226 neuroectodermal lineage with a comparable cell density (**Fig. 5D**) and morphology  
227 (our unpublished data) as *WT* cells. Finally, quantitative analyses did not reveal any  
228 significant alteration in either NPC density or nuclear size in *Mios*<sup>-/-</sup> as compared to  
229 *WT* mESCs (**Fig. 5E, F**).

230 Together these experiments indicate that neither the growth and differentiation  
231 defects, nor the altered nuclear pore density and nuclear sizes observed in *Seh1*<sup>-/-</sup>  
232 mESCs can be merely attributed to the decreased levels of Mios.

233

234 **Mutations affecting the short arm of the Y-complex impair NPC assembly, but with**  
235 **distinct impacts on cell proliferation and differentiation**

236 Having excluded Mios destabilization as the main cause of the defects of *Seh1*<sup>-/-</sup>  
237 mESCs, we next focused our attention on Seh1's partners localized on the short arm  
238 of the Y-complex (**Figs. 1A and 6A**).

239 We first inactivated *Nup43*, another small β-propeller-folded nucleoporin that is  
240 specific to metazoan Y-complexes (Neumann et al., 2010). We obtained viable clones  
241 upon CRISPR/Cas9-mediated *Nup43* knockout in mESCs that however displayed  
242 impaired proliferation at the pluripotent stage (**Fig. 6B, C**). Although this growth  
243 defect was milder than the one observed in pluripotent *Seh1*<sup>-/-</sup> mESCs, *Nup43*<sup>-/-</sup>  
244 nevertheless underwent drastic cell death upon neuroectodermal differentiation,  
245 comparable to that of differentiating *Seh1*<sup>-/-</sup> cells (**Fig. 6D**). *Nup43*<sup>-/-</sup> mESCs also  
246 displayed a reduced NPC density comparable to that observed in the various *Seh1*<sup>-/-</sup>  
247 mESC lines, as revealed by the reduced intensity of Nup133, Nup98 and TPR labelling  
248 at nuclear pores (**Fig. 7A-C**, see also **Fig. 2A-C**). Finally, these cells showed no  
249 significant reduction in nuclear size (**Fig. 7D**). Together these data indicate that  
250 inactivation of *Nup43* mimics, albeit with a slightly milder impact, most of the  
251 phenotypes caused by *Seh1* inactivation.

252 To determine whether these shared phenotypes reflect a function of these Nups  
253 within the short arm of the Y-complex, we next aimed to impair Seh1 recruitment to  
254 the NPCs. Structural studies have shown that budding yeast Seh1 binds to Nup85  
255 through its N-terminal domain forming the 7<sup>th</sup> blade to complete the β-propeller

256 structure of Seh1 (Brohawn et al., 2008; Debler et al., 2008). Homology modelling  
257 predicted that the human Nup85-Seh1 interface similarly involves  $\beta$ -sheets within  
258 Nup85 N-terminal domain that complete Seh1  $\beta$ -propeller (von Appen et al., 2015);  
259 see **Fig. 6A**). We thus attempted to prevent Seh1 binding to Nup85 by deleting most  
260 of this blade (two  $\beta$ -sheets encoded by exon 2 of mouse Nup85; colored in yellow in  
261 **Fig. 6A**) and inserting instead the sequence of the bulky GFP. We obtained viable  
262 mESC lines in which the resulting  $\Delta$ E2-GFP-Nup85 fusion, expressed as the unique  
263 form of Nup85 in the cell (**Figs. 6B and S3C**), was properly localized at both NPCs and  
264 kinetochores (**Fig. S3A**).

265 To determine whether this deletion within Nup85 indeed prevented its interaction  
266 with Seh1, we performed immunoprecipitation on *WT* or  $\Delta$ E2-GFP-Nup85 mESC  
267 lysates using antibodies directed against either Nup85 itself, or Nup107, another Y-  
268 complex constituent (**Fig. 1A**). Mass-spectrometry analysis showed that  $\Delta$ E2-GFP-  
269 Nup85 interacts with all the members of the Y-complex except Seh1 (**Fig. S3B**).  
270 Because none of the available Seh1 antibodies we tested properly recognized  
271 endogenous mouse Seh1 in immunofluorescence experiments, we also introduced  
272 within *GFP-Seh1* cells the same N-terminal deletion of Nup85, this time tagged with  
273 mCherry (**Fig. S3C** and **Table S2**). Although  $\Delta$ E2-mCherry-Nup85 was properly  
274 localized at NPCs and kinetochores (**Fig. 8A,B**), GFP-Seh1 was at most only barely  
275 detectable at kinetochores in these cells (**Fig. 8B,D**; quantifications revealed  $7-8 \pm 5-$   
276 7% residual signal in  $\Delta$ E2-mCherry-Nup85- compared to wt-Nup85-expressing cells).  
277 The mislocalization of Seh1 from kinetochores is consistent with its impaired  
278 interaction with  $\Delta$ E2-GFP-Nup85 seen at the biochemical level and with previous  
279 studies indicating that the Y-complex is recruited as an entity to kinetochores  
280 (Loiodice et al., 2004). In contrast, we could still detect some punctate GFP-Seh1  
281 signal at the nuclear envelope (26-27% residual signal; **Fig. 8A,C**). The relative  
282 persistence of GFP-Seh1 at NPCs as compared to kinetochores in  $\Delta$ E2-mCherry-  
283 Nup85 cells likely reflects the existence of additional minor binding sites for Seh1 at  
284 NPCs, that either involve Nups not belonging to the Y-complex, or imply interfaces  
285 generated by the 3D-organization of Y-complex within the assembled mammalian  
286 NPC (Huang et al., 2020; Kosinski et al., 2016; von Appen et al., 2015).

287 Unexpectedly, analysis of cell differentiation did not reveal any significant  
288 differences between  $\Delta E2$ -GFP-*Nup85* and *WT* mESCs (Fig. 6D). In addition  $\Delta E2$ -GFP-  
289 *Nup85* mESCs only displayed a minor cell growth defect in the pluripotent state  
290 (9±7% decrease in confluence after 48h of growth compared to *WT* mESCs, while the  
291 reduction was 60±7% and 38±11% for *Seh1*<sup>-/-</sup> and *Nup43*<sup>-/-</sup> cells, respectively; Fig. 6C).  
292 Nevertheless, pluripotent  $\Delta E2$ -GFP-*Nup85* cells display a significant reduction in the  
293 intensity of Nup133, Nup98 and TPR at the nuclear envelope (Fig. 7A-C). These  
294 decreased signals were comparable to those observed in *Seh1*<sup>-/-</sup> and *Nup43*<sup>-/-</sup> mESCs,  
295 and yet, as in the case of *Nup43*<sup>-/-</sup>, they were not accompanied by a significant  
296 reduction in nuclear size (Fig. 7D). Analysis of the  $\Delta E2$ -GFP-*Nup85* cells lines thus  
297 showed that perturbed recruitment of Seh1 at NPCs leads to a reduction in NPC  
298 number, but does not impact cell growth and differentiation.

299

300 **Discussion**

301 This study has revealed that Seh1 and Nup43 are dispensable for mESC viability in  
302 the pluripotent state but become critical upon their differentiation. In view of the  
303 reported embryonic lethality of the *Seh1* knockout in mouse, an impaired  
304 differentiation of *Seh1*<sup>-/-</sup> mESCs could have been anticipated (Liu et al., 2019). In  
305 contrast, no role in development had been previously described for Nup43, which is  
306 specific to metazoans (Neumann et al., 2010). Although the requirement for  
307 differentiation was reminiscent of the phenotype observed upon inactivation of  
308 *Nup133*, we observed that Seh1 and Nup43, are, unlike Nup133 (Lupu et al., 2008),  
309 also required for proper growth of mESCs in the pluripotent state. Importantly, this  
310 altered growth is not simply caused by a mitotic defect, as might have been assumed  
311 given the mitotic roles of Seh1 in cancer cells (Platani et al., 2018; Platani et al.,  
312 2009; Zuccolo et al., 2007), but rather reflects a lengthening of all phases of the cell  
313 cycle.

314 We initially anticipated that the phenotypes of *Seh1*<sup>-/-</sup> mESCs could be caused by a  
315 combination of its functions within the Y- and the GATOR2-complexes. However, the  
316 fact that the *Mios*<sup>-/-</sup> cells do not feature any NPC assembly, nuclear size, or cell

317 differentiation defects rather suggests that *Seh1*<sup>-/-</sup> phenotypes (except perhaps a  
318 mild contribution to cell growth rates) are unlikely to result from a combination of  
319 defects in NPC and GATOR function. Moreover, *Nup43*<sup>-/-</sup> mESC phenotypes are very  
320 similar to those of *Seh1*<sup>-/-</sup> mESCs, despite the fact that Nup43 does not interact with  
321 Mios.

322 Our data also showed that integrity of the short arm of the Y-complex is important  
323 for proper NPC density, further distinguishing its function from that of Nup133,  
324 which specifically affects NPC basket assembly in mESCs (Souquet et al., 2018). The  
325 observed reduction in NPC density in our *Seh1*<sup>-/-</sup>, *Nup43*<sup>-/-</sup> and  $\Delta$ E2-GFP-*Nup85* clones  
326 likely reflects an absolute reduction in total NPC number, since there was no  
327 corresponding increase in nuclear surface in these cells (instead, nuclear size was  
328 mildly reduced in *Seh1*<sup>-/-</sup> mESCs). Different mechanisms may explain the requirement  
329 for an intact short arm of the Y-complex to ensure proper NPC numbers. Considering  
330 the critical roles of the Y-complex in both NPC re-assembly after mitosis and *de novo*  
331 NPC assembly in interphase (Doucet et al., 2010; Harel et al., 2003; Walther et al.,  
332 2003), the short arm of the Y-complex might be required for the efficient  
333 recruitment of the Y-complex either to the mitotic chromatin (an hypothesis  
334 consistent with the minor reduction of Y-complex levels on chromatin reported upon  
335 *Seh1* depletion in HCT116 cells - Platani et al., 2018), or to the nuclear envelope in  
336 interphase (as process involving Nup153 Vollmer et al., 2015). Alternatively, Nup43  
337 and *Seh1* may contribute to the stabilization of the NPC scaffold, by virtue of their  
338 direct interactions with neighbouring subunits from either Y-complexes or inner ring  
339 complexes (Huang et al., 2020; Kosinski et al., 2016; von Appen et al., 2015). NPCs  
340 lacking these stabilizing interactions might then be recognized by one of the recently  
341 described quality-control mechanisms that mediate the removal of some  
342 misassembled NPCs from the nuclear envelope (reviewed in Webster and Lusk,  
343 2016).

344 Finally, our analysis of the  $\Delta$ E2-GFP-*Nup85* cell lines indicates that the reduction in  
345 NPC density observed in *Seh1*<sup>-/-</sup> and *Nup43*<sup>-/-</sup> mESCs is not sufficient to impact cell  
346 growth and differentiation. The lack of major growth and differentiation defects in  
347  $\Delta$ E2-GFP-*Nup85* cells, in which *Seh1* is largely mislocalized from NPCs, could reflect

348 an “off-pore” function of Seh1, or a function of Seh1 that does not require its normal  
349 stoichiometry within NPCs (for instance, a localization restricted to the cytoplasmic  
350 or nuclear side of the NPCs). At NPCs, Seh1 and Nup43 might be required for the  
351 proper recruitment and positioning of the mRNA export and remodelling machinery,  
352 an established function of the short arm of the Y-complex in budding yeast  
353 (Fernandez-Martinez et al., 2016). Alternatively, whether at pores or elsewhere in  
354 the nucleus, Seh1 and Nup43 may impact cell growth and differentiation by directly  
355 contributing to gene regulation, as now reported for a few Nups in mammalian cells  
356 (reviewed in Buchwalter et al., 2019; Pascual-Garcia and Capelson, 2019; see also  
357 Scholz et al., 2019). In particular, Seh1 was recently found to participate in  
358 oligodendrocyte differentiation, acting as a platform to recruit transcription and  
359 chromatin remodelling factors (Olig2 and Brd7) (Liu et al., 2019). We may  
360 hypothesize that both Seh1 and Nup43 may specifically interact with factors  
361 required for gene regulation and chromatin organization in mESCs, hence  
362 contributing to early stages of pluripotent cell growth and differentiation.

363

363 **Materials and Methods**

364 **Plasmids** used in this study are listed in **Table S1**. They were either previously  
365 published or generated using standard molecular cloning techniques including  
366 restriction digestions (Fastdigest, Thermo Fisher Scientific, Waltham, MA), PCR  
367 amplification using proofreading DNA polymerases (Phusion HF, New England Biolab,  
368 Ipswich, MA) and In-Fusion HD Cloning Kit (Clontech, Mountain View, CA) or  
369 NEBuilder HiFi DNA Assembly Cloning Kits (New England Biolab). The *Mios* and  
370 *Nup43* gRNAs were integrated in a linear plasmid (GeneArt™ CRISPR Nuclease Vector  
371 – OFP-Cas9) following manufacturer's instructions. The other Cas9 vectors (pX-280,  
372 pX-672, pX-853 and pX-864) were assembled by golden gate cloning (Engler et al.,  
373 2009). For all constructs, PCR-amplified fragments and junctions were checked by  
374 sequencing. Plasmid maps are available upon request.

375

376 **Cell lines, growth condition, transfection, and CRISPR/CAs9-based genome editing**

377 Cell lines used in this study are listed in **Table S2**. All cells were grown at 37°C and 5%  
378 CO<sub>2</sub>.

379 DR4-mouse embryonic fibroblast feeder cells, DR4-MEFs (Applied StemCells), were  
380 grown in Dulbecco's Modified Eagle's Medium (DMEM) (Gibco/Thermofisher)  
381 supplemented with 15% heat-inactivated foetal bovine serum (FBS, Gibco), 100U/ml  
382 Penicillin-100µg/ml Streptomycin (P/S) (Gibco) and 2mM L-Glutamine (Gibco). DR4-  
383 MEFs were inactivated using 8.5 µg/ml Mitomycin-C (Sigma-Aldrich, St Louis, MO) for  
384 3 hours.

385 HM1 (Selfridge et al., 1992) and derivative mESCs clones were grown in  
386 serum/leukemia inhibitory factor (LIF)-containing stem cell medium: DMEM  
387 (EmbryoMax, Millipore, Burlington, MA), P/S (Gibco), 2mM L-Glutamine (Gibco), 15%  
388 heat-inactivated ESC-Qualified FBS (Gibco), non-essential amino acids (Gibco),  
389 nucleosides (Millipore), 2-mercaptoethanol (Gibco) and 10<sup>3</sup> units/ml LIF (ESGRO,  
390 Millipore). mESCs were grown on inactivated DR4-MEFs (MEF-derived feeders)  
391 plated on 0.1% gelatin (Sigma-Aldrich) and were passaged every 2 or 3 days using  
392 0.05% Trypsin (Gibco). mESCs were used at passages below 30. Lack of  
393 contamination in-between the mutant cell lines was assessed by PCR on genomic

394 DNA, proper GFP-or mCherry expression when pertinent, and western blots  
395 analyses. Frequent DAPI staining ensured lack of major contamination by  
396 mycoplasm. When required, cells were counted using a Countess automated cell  
397 counter (Invitrogen, Carlsbad, CA).

398 For transfections, mESCs were plated onto DR4-MEFs in medium without P/S.  
399 Plasmid DNA and Lipofectamine 2000 (Invitrogen) were mixed in OptiMEM  
400 (Invitrogen) and added to the cells according to the manufacturer's instructions.

401 For CRISPR/Cas9 editing,  $5 \cdot 10^5$  mESCs were transfected as indicated in **Table S2** with  
402 one or two plasmids (3 $\mu$ g each) directing the expression of one or two gRNAs along  
403 with Cas9 (WT or high fidelity, HF) fused to GFP, mCherry or OFP. gRNAs were  
404 designed using the Benchling website (<https://benchling.com>) and are listed in **Table**  
405 **S3**. When indicated, DNA sequences of interest (PCR product 1-4  $\mu$ g, or 3 $\mu$ g of  
406 linearized plasmid) flanked by homology directed repair arms were co-transfected  
407 (**Fig. S4**). Following selection (as detailed below), individual clones were picked,  
408 amplified, and further characterized. For each clone, chromosome spreads were also  
409 performed (chromosome counts are indicated in **Table S2**).

410 To establish *Seh1*<sup>-/-</sup>, *Mios*<sup>-/-</sup> and *Nup43*<sup>-/-</sup> cell lines, cells were collected by  
411 trypsinization two days after transfection, resuspended in 1 mL Fluorescence-  
412 activated cell sorting (FACS) buffer (PBS +10% FBS, Gibco + P/S), and sorted based on  
413 Cas9 expression (EGFP or OFP signal). 2000 FACS-sorted cells were plated in 100mm  
414 culture dishes. Individual clones picked 6-12 days after sorting were then  
415 characterized using Western blot and PCR on genomic DNA followed by sequencing  
416 (the identified Indels are listed in **Table S2**).

417 To establish the *OsTir* cell line, 200 $\mu$ g/mL Geneticin (Geneticin® Gibco, Life  
418 technologies 10131-019) was added to the medium two days after transfection.  
419 Geneticin-resistant clones (expected to have integrated the pCAG-OsTir-T2A-NeoR  
420 sequence at the *Tigre* locus) were picked after five days and characterized by  
421 Western-blot with antibodies directed against the OsTir receptor. PCR on genomic  
422 DNA was also performed to determine the number of *Tigre* alleles bearing the  
423 transgene.

424 To generate the *Seh1 rescue* (expressing GFP-Seh1 under the pCAG promoter at the  
425 *Tigre* locus), *GFP-Seh1*, *GFP-mAID-Seh1*, *ΔE2-GFP-Nup85* and *[ΔE2-mCherry-Nup85]*  
426 cell lines, GFP [mCherry] positive cells were FACS-sorted 3 days after transfection to  
427 select for cells expressing the tagged nucleoporin. Individual clones were picked 6-7  
428 days after sorting and characterized using immunofluorescence (to confirm the  
429 localization of the tagged protein at the nuclear periphery) and western blot (to  
430 identify clones lacking the endogenous protein). The selected clones were then  
431 further validated by PCR on genomic DNA and sequencing.

432 To achieve inducible degradation of GFP-mAID-Seh1, Auxin (Sigma-Aldrich) was  
433 added to the medium at 500µM (from a stock at 280 mM in EtOH). For control  
434 experiments, the same amount of EtOH was added.

435

#### 436 **Cell growth and differentiation assays**

437 To evaluate cell growth at pluripotent stage, cells were plated at 1 - 2·10<sup>5</sup> cells per  
438 well in TPP 12-well plate. Photomicrographs were taken every two hours using an  
439 IncuCyte® live cell imager (Essen Biosciences, Ann Arbor, MI) and confluence of the  
440 cultures was measured using IncuCyte® software (Essen Biosciences, Ann Arbor,  
441 MI). To improve comparison in-between experiments or cell lines, the same mask  
442 was always used and time was set at t=0 when confluence reached 1% (**Figs. 1D and**  
443 **6C**), 2% (**Fig. 5C**) or 3% (**Fig. S1E**). Graphs were generated using Excel. Error bars  
444 correspond to standard deviations from the indicated [n=] independent  
445 experiments.

446

447 Neuroectodermal differentiation of mESCs grown as monolayers was adapted from  
448 (Ying and Smith, 2003). Following trypsinization, feeders were removed by plating  
449 the resuspended cells in gelatin-free wells for 20 minutes. Feeder-free mESCs were  
450 collected and resuspended in N2B27 medium [DMEM F-12, DMEM Neurobasal, BSA,  
451 L-glutamin, 2-mercaptoethanol, N2 (Gibco) and B27 (Gibco)]. Cells were plated in  
452 gelatin-coated wells at 1·10<sup>5</sup> or 3·10<sup>4</sup> cells per well in TPP 12 well plate. At day 2,  
453 N2B27 medium containing 1µM RA (all-trans-Retinoic acid, Sigma) was added for 24  
454 hours. From day 3 to 7 medium was changed every day with fresh N2B27 without

455 RA. Confluence analyses, used as a proxy to evaluate cell growth and viability, was  
456 performed as described above except that time was set at t=0 the beginning of the  
457 differentiation process (i.e., upon plating in N2B27 medium).

458

#### 459 **Fluorescence Videomicroscopy**

460 mESCs were transiently transfected using plasmids expressing H2B-GFP or H2B-  
461 mCherry on microscopy-adapted 35-mm dishes ( $\mu$ -dish, 35 mm, high; Ibidi,  
462 Germany) coated with 0.1% gelatin and DR4-MEFs. Acquisitions were performed  
463 about 36 hours after transfection at 37°C and 5% of CO<sub>2</sub> using an AxioObserverZ1  
464 inverted microscope (Zeiss, Germany) equipped with a 63 oil objective, a CSU-X1  
465 spinning-disk head (Yokogawa, Japan), and a sCMOSPRIME 95B (Photometrics)  
466 camera.

467 The whole setup was driven with MetaMorph software (Molecular Devices,  
468 Sunnyvale, CA). Eleven Z sections with a step of 1 $\mu$ m were acquired at intervals of 5  
469 minutes for the mitotic progression experiments (4-6 hours) and of 15 minutes for  
470 the cell cycle length experiments (24-30 hours). Laser intensity was set between 10-  
471 20% power and acquisition time was 500ms. The raw data were processed using  
472 ImageJ software (National Institutes of Health, Bethesda, MD). Images stacks were  
473 processed as max projections. Cells were tracked manually setting prometaphase at  
474 the moment at which chromatin starts to be seen condensed and anaphase at the  
475 first time point at which chromosome segregation is observed.

476

#### 477 **FACS analyses**

478 To perform a bi-parametric analysis of cell cycle based on DNA content (DAPI) and  
479 DNA synthesys (EdU) we used the Click-it-EdU Imaging kits (Invitrogen). 0.5 10<sup>6</sup> mESC  
480 were plated on MEF-derived feeders plated on 0.1% gelatin 2 days before the  
481 experiment. Cells were incubated with EdU 50  $\mu$ m for 15 minutes, then collected by  
482 trypsinization and plated on gelatin dishes for 20 minutes to remove feeders.

483 Cells were then collected and centrifuged, washed in PBS and then fixed in PFA 3%  
484 for 15 minutes. Cells were then permeabilized with PBS+ 0.2% Triton-X100 and  
485 washed with PBS + 2% SVF. Click reaction (30 minutes) was performed following  
486 manufacturer's protocol. DNA was stained for 20 minutes (DAPI 5  $\mu$ M, RNaseA 0.1

487 mg/mL, 1% SVF in PBS), then samples were centrifuged and resuspended in 60 $\mu$ L PBS  
488 +2% SVF.

489 Sample acquisition was achieved with the ImageStream® X (Amnis, Austin, TX)  
490 imaging flow cytometer and captured using the ISX INSPIRE™ data acquisition  
491 software. Images of 5.000–20.000 cells were acquired at 40x magnification using the  
492 following channels: Ch1= 430-470 nm, BF (bright field); Ch6= 720-800nm, SS (side  
493 scatter); Ch7= 430-505 nm, DAPI; Ch9= 570-595, BF; Ch11= 660-720 nm, EdU-AF647.  
494 A compensation matrix was generated using fluorescence controls and applied to all  
495 samples. Analysis was then performed with the IDEAS software as follows: 1)  
496 definition of cells in focus, based on the gradient RMS; 2) definition of singlets,  
497 according to area and aspect ratio; 3) definition of cells using contrast and gradient  
498 RMS; 4) definition of nucleated cells using DNA content; 5) cell cycle phases were  
499 then identified using DAPI and EdU intensity; 6) mitotic cells were finally defined  
500 according to DAPI bright detail intensity and DAPI area threshold (**Fig. S2 D**).

501

## 502 **Immunostaining and quantitative image analyses**

503 mESCs grown on coverslips were washed with PBS, then fixed using 3% para-  
504 formaldehyde (VWR, Radnor, PA) for 20 min and washed again with PBS. For all  
505 conditions, cells were then permeabilized in PBS + 0.2% Triton X-100, 0.02% SDS  
506 (Euromedex, Souffelweyersheim, FR), and 10 mg/mL BSA (Sigma). Antibody  
507 hybridizations and washes were also performed in this buffer. Primary and  
508 secondary antibodies (listed in **Table S4**) were incubated for 1 hr at room  
509 temperature. Cells were then incubated 5 min with 280nM DAPI (Sigma) in PBS and  
510 mounted with Vectashield (Vector, Maravai Life Sciences, San Diego, CA). Images  
511 were acquired using 100x/1.4 oil objectives on inverted and motorized microscopes,  
512 either a DMI8 (Leica), equipped with a CSU-W1 spinning-diskhead (Yokogawa, Japan)  
513 and 2 Orca-Flash 4 V2+ sCMOS cameras (Hamamatsu), or an Axio Observer.Z1  
514 (Zeiss), equipped with CSU-X1 spinning-diskhead (Yokogawa, Japan) and 2 sCMOS  
515 PRIME 95 cameras (Photometrics).

516 Quantification of NPC density at the nuclear envelope (NE) was performed  
517 essentially as described (Souquet et al., 2018). Briefly, mESCs of interest were mixed

518 with a GFP cell line of reference (*Rescue-Seh1* or  $\Delta E2$ -GFP-*Nup85* cells, used for  
519 normalization) and grown on coverslips for 24h prior to fixation and immunostaining.  
520 For each acquired image, one z section was selected; 8-pixel-thick regions of interest  
521 (ROIs) were drawn freehand on the NE of both GFP-negative and -positive  
522 (reference) mESCs. Following subtraction of background, the signal intensity at the  
523 NE for each cell was normalized to the average NE intensity measured for the GFP-  
524 positive mESCs acquired in the same field. All values were then divided by the mean  
525 normalized intensity of *WT* mESCs acquired in the same experiment. Box plots were  
526 generated using GraphPad Software: each box encloses 50% of the normalized  
527 values obtained, centred on the median value. The bars extend from the 5<sup>th</sup> to 95<sup>th</sup>  
528 percentiles. Values falling outside of this range are displayed as individual points.

529 For kinetochore quantifications, mixed *GFP-Seh1* and  $\Delta E2$ -*mCherry-Nup85/GFP-Seh1*  
530 mESCs grown on coverslips were fixed, permeabilized, and stained with DAPI. Fields  
531 containing both *GFP-Seh1* and  $\Delta E2$ -*mCherry-Nup85/GFP-Seh1* mitotic cells were  
532 selected. For each mitotic cell, the mean intensities of five distinct kinetochores  
533 (regions of 10-pixels in diameter) and of two "background" regions in the mitotic  
534 cytoplasm (40-pixels in diameter) were measured on a unique z-section. Following  
535 background subtraction, the average intensity of *GFP-Seh1* at kinetochores in each  
536 mitotic cell was normalized to the intensity measured for the control (*wtNup85/*  
537 *GFP-Seh1*) mitotic cells acquired in the same experiment. Box plots were generated  
538 as described above.

539  
540 To quantify nuclear surfaces, mESCs of interest were mixed with a GFP cell line of  
541 reference and grown on coverslips for 24 hr prior to fixation and immunostaining as  
542 described above. For each field, 33 to 45 optical sections (0.5  $\mu$ m apart) were  
543 acquired and nuclei were segmented based on TPR immunostaining with the Fiji  
544 plugin Lime-Seg (Machado et al., 2019). A circular ROI was drawn within the nucleus  
545 of each cell in the field and the LimeSeg Plugin "Sphere Seg advanced" was run with  
546 the following parameters: D0: 4; Zscale: 7.143; range in D0 units: 2; real xy pixel size:  
547 0.07; F pressure: 0.025 for TPR-Cy3 staining and 0.019 for TPR-Cy5. Segmented  
548 structures for which the "free edges" values were above 0 (segmentation could not  
549 close the structure), and those for which the Euler characteristic was not comprised

550 between -4/+4 (aberrant structures very far from a spherical shape) were discarded.  
551 For each cell, the segmentation perimeter and TPR staining along the z-axis were  
552 compared to further validate proper segmentation (less than 8% of the identified  
553 structures, frequently corresponding to the merge of two closely apposed nuclei,  
554 were manually discarded at that stage). Nuclear surfaces and volumes were then  
555 exported. To compensate for variability occurring during fixation or IF processing,  
556 nuclear surface values were first normalized to the average of the GFP-reference  
557 cells acquired within the same coverslip, and then to the mean of *WT* mESCs  
558 acquired in the same experiment. Nuclear surface graphs were generated using  
559 GraphPad Software: average and standard deviation (boxes and bars) of nuclear  
560 surface are displayed, along with values for each experiment (dots).

561

## 562 **Western blot analyses**

563 To prepare whole cell lysates, mESCs were lysed in 2× Laemmli lysis buffer (150-mM  
564 Tris-HCL (pH 6.8), 5 % (wt/vol) SDS, 25 % (vol/vol) glycerol, and 0.01 % (wt/vol)  
565 bromophenol blue). Lysates were incubated for 3 min at 95 °C, clarified by sonication  
566 (Bioruptordiagenode: 4 cycles of 30 s on/off, high power), and denatured again for 3  
567 min at 95°C. Protein concentration was then determined using a BCA assay kit  
568 (Thermo Fisher Scientific). Total protein extracts supplemented with  $\beta$ -  
569 mercaptoethanol (750 mM final, Sigma-Aldrich) were analysed by western blot. 10  
570  $\mu$ g of mESC lysate were separated on 4–12% or 10% SDS-PAGE gels (pre-cast  
571 NuPage® GE healthcare or Mini-Protean TGX Stain free precast gels, Biorad,  
572 Hercules, CA) and transferred to nitrocellulose (GE healthcare). The resulting blots  
573 were stained using Ponceau, saturated with TBS buffer + 0.1% Tween and 5% dried  
574 milk, and incubated in TBS + 0.1% Tween and 5% dried milk with primary antibodies,  
575 followed by either HRP-conjugated secondary antibodies of interest or HRP-  
576 conjugated anti-rabbit TrueBlot® secondary antibody (used in **Fig. 5B** to prevent  
577 interference from the denatured/reduced heavy and light chains of the anti-Seh1  
578 antibody used for immunoprecipitation (primary and secondary antibodies are listed  
579 in **Table S4**). Signals were detected by enhanced chemiluminescence (SuperSignal®  
580 Pico or Femto, ThermoScientific) using ChemiDoc (Biorad).

581

582 **Immunoprecipitation experiments and mass spectrometry analyses**

583 Immunoprecipitation experiments: Protein G beads (GE Helathcare) were washed  
584 three times with Wash buffer (100mM NaCl, 1mM EDTA, 25mM TRIS pH 7.5, 1mM  
585 DTT + protease inhibitor [Pi] solution); (bead centrifugations were performed at 500g  
586 4°C). 30µL of beads were then incubated for 2 hours at 4°C in 250µL wash buffer  
587 containing 5µL of rabbit anti-Seh1 antibody (for **Fig. 5B**), or 25µL of rabbit polyclonal  
588 anti-Nup107 or anti-Nup85 serum or a pre-immune rabbit serum as control (for **Fig.**  
589 **S3B**). Antibodies used are listed in **Table S4**. After incubation, beads were washed 4  
590 times with Wash buffer.

591 In the meantime, lysates were prepared from *WT*, *Seh1*<sup>-/-</sup>, *Mios*<sup>-/-</sup> or *ΔE2-GFP-Nup85*  
592 mESCs by resuspending frozen pellets of 4·10<sup>6</sup> mESCs (~500µg total proteins) in  
593 200µL Lysis buffer (100mM NaCl, 1mM EDTA, 25mM TRIS pH 7.5, 1mM DTT, Tween-  
594 20 0.5%, Triton-100 1.2% + PI solution). Samples were vortexed and incubated 15  
595 min on ice. 600µL of Dilution buffer (100mM NaCl, 1mM EDTA, 25mM TRIS pH 7.5,  
596 1mM DTT, Tween-20 0.5%, + PI solution) was then added and samples were  
597 centrifuged at 16.000 g for 30 min at 4°C. The resulting supernatant was pre-cleared  
598 by a 1 hour incubation at 4°C with 30µL Protein G beads equilibrated with wash  
599 buffer.

600 The cleared supernatants (inputs) were then incubated at 4°C with 30µL of the anti-  
601 Seh1, control, anti-Nup107- or anti-Nup85-coated Protein G beads. After overnight  
602 (for anti-Seh1) or 2 hours incubation (for anti-Nup107 and anti-Nup85), samples  
603 were centrifuged and washed 5 times in wash buffer. The proteins were either  
604 eluted in 40µL of Laemmli and boiled 10 minutes for subsequent western blot  
605 analysis (**Fig. 5B**), or split in 2 and then either eluted in 20µL of Laemmli and boiled 3  
606 minutes for subsequent western blot analyses, or processed for both mass-  
607 spectrometry and western blot analyses (for experiments presented in **Fig. S3**).

608 Samples preparation prior to LC-MS/MS analysis: Proteins on beads were digested  
609 overnight at 37°C with trypsin (Promega, Madison, WI, USA) in a 25-mM NH<sub>4</sub>HCO<sub>3</sub>  
610 buffer (0.2µg trypsin in 20µL). The resulting peptides were desalted using ZipTip µ-  
611 C18 Pipette Tips (Pierce Biotechnology, Rockford, IL, USA).

612 **LC-MS/MS acquisition:**

613 Samples were analyzed using an Orbitrap Fusion, coupled to a Nano-LC Proxeon  
614 1200, equipped with an easy spray ion source (Thermo Scientific, Waltham, MA,  
615 USA). Peptides were loaded with an online preconcentration method and separated  
616 by chromatography using a Pepmap-RSLC C18 column (0.75 x 750 mm, 2  $\mu$ m, 100  $\text{\AA}$ )  
617 from Thermo Scientific, equilibrated at 50°C and operated at a flow rate of 300  
618 nl/min. Solvents (MS grade H<sub>2</sub>O, formic acid (FA) and Acetonitrile (ACN)) were from  
619 Thermo Chemical (Waltham, MA, USA).

620 Peptides were eluted by a gradient of solvent A (H<sub>2</sub>O, 0.1 % FA) and solvent B  
621 (ACN/H<sub>2</sub>O 80/20, 0.1% FA). The column was first equilibrated 5 min with 95 % of  
622 solvent A, then solvent B was raised to 28 % in 105 min and to 40% in 15 min. Finally,  
623 the column was washed with 95% solvent B during 20 min and re-equilibrated at  
624 95% solvent A during 10 min. On the Orbitrap Fusion instrument, peptides precursor  
625 masses were analyzed in the Orbitrap cell in full ion scan mode, at a resolution of  
626 120,000, a mass range of *m/z* 350-1550 and an AGC target of 4.10<sup>5</sup>. MS/MS were  
627 performed in the top speed 3s mode. Peptides were selected for fragmentation by  
628 Higher-energy C-trap Dissociation (HCD) with a Normalized Collisional Energy of 27%  
629 and a dynamic exclusion of 60 seconds. Fragment masses were measured in an Ion  
630 trap in the rapid mode, with and an AGC target of 1.10<sup>4</sup>. Monocharged peptides and  
631 unassigned charge states were excluded from the MS/MS acquisition. The maximum  
632 ion accumulation times were set to 100 ms for MS and 35 ms for MS/MS acquisitions  
633 respectively.

634 Data analysis: Raw data were processed on Proteome Discoverer 2.2 with the  
635 mascot node (Mascot version 2.5.1) and the Swissprot protein database release  
636 2017\_06. The *Mus musculus* taxonomy was used and a maximum of 2 missed  
637 cleavages was authorized. Precursor and fragment mass tolerances were set to 7  
638 ppm and 0.5 Da. The following Post-translational modifications were included as  
639 variable: Acetyl (Protein N-term), Oxidation (M), Phosphorylation (STY). Spectra were  
640 filtered using a 1% FDR with the percolator node.

641

642 **Statistics**

643 For cell confluence analyses, statistical analyses were performed at the latest time  
644 points (48h for cell growth in the pluripotent state and day 5 for neuroectodermal  
645 differentiation) using paired two-tailed Student's t-test. For each mutant cell line,  
646 the % of confluence was compared to that of *WT* cells measured in the same  
647 experiment. To obtain more robust statistics, the paired two-tailed Student's t-test  
648 was also used to compare all the values obtained with distinct clones bearing the  
649 same mutation to *WT* cells. For studies of interphase and mitosis duration and for  
650 quantifications of fluorescence intensity at the NE, statistical analyses were  
651 performed using unpaired non-parametric Mann-Whitney test. For nuclear surfaces,  
652 statistical analyses were performed using paired two-tailed Student's t-test. P values  
653 and significance: \*\*\*\*: P <0.0001; \*\*\*: P <0.001; \*\*: P <0.01; \* :P <0.05.

654

655 **DATA AND SOFTWARE AVAILABILITY**

656 The mass spectrometry proteomics data reported in this study have been deposited  
657 to the ProteomeXchange Consortium  
658 (<http://proteomecentral.proteomexchange.org>) via the PRIDE partner repository  
659 (Perez-Riverol et al., 2019) with the dataset identifier PXD022190.

660

661 The original 16-bit images and montages of the western blots used in this study  
662 are available as Mendeley dataset under <http://dx.doi.org/10.17632/8g59mp92bs.1>

663

664

665

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678 **Competing interests**

679 No competing interests declared.

680

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695

696 **AUTHOR CONTRIBUTIONS**

697 A.G-E., A.V., C.O., B.R-S-M., and V.D. conceived and designed the experiments.  
698 A.G-E., A.V., and C.O. performed the experiments. A.G-E., A.V., C.O., and V.D.  
699 analyzed the data. A.G-E., A.V., and V.D. wrote the manuscript with contribution  
700 from all co-authors.

701

701 **References**

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840

840

841 **FIGURE AND FIGURE LEGENDS**

842 **Figure 1: Seh1 depletion leads to cell growth delay and impaired cell survival upon**  
843 **differentiation.**

844 **A.** Schematic representation of the Y-complex (adapted from von Appen et al.,  
845 2015), highlighting the components of the short arm, namely Nup43 (shown in  
846 purple), Nup85 (blue) and Seh1 (red).  $\beta$ -propellers are outlined with black strokes. **B.**  
847 Western-blot showing the expression of endogenous or GFP-tagged Seh1, Nup107  
848 and GAPDH (used as loading control) in whole cell extracts from the indicated cell  
849 lines. 1/2 and 1/4 dilutions of *WT* extracts were also loaded. Molecular weights are  
850 indicated (kilodaltons). **C.** Representative phase contrast images of *WT* and *Seh1*<sup>-/-</sup>  
851 (#1) mESCs colonies acquired after 2 days of growth on the IncuCyte® imager. Scale  
852 bar, 300 $\mu$ m. **D-E.** Confluence of *WT* (blue), *Seh1*<sup>-/-</sup> (#1 and #2, red) and *Rescue* (#1  
853 and #2, grey) mESCs was quantified with the IncuCyte® system either at pluripotent  
854 stage (**D**) or upon differentiation towards neuroectodermal lineage (**E**). Error bars  
855 correspond to the standard deviation arising from the indicated number of  
856 independent experiments [n]. Statistical analyses of these confluence curves were  
857 performed at the last time-points. Brackets indicate statistics performed using all  
858 values from cell lines bearing a given mutation, compared to *WT* (see Materials and  
859 Methods). **F.** Representative spinning disk images of *Rescue* (#1) mESCs showing  
860 proper localization of GFP-Seh1 at the NE in interphase (one plane) and at  
861 kinetochores in mitosis (a projection of 3 optical sections is presented). Scale bars,  
862 10  $\mu$ m.

863

864 **Figure 2: The altered growth rate of *Seh1*<sup>-/-</sup> mESCs reflects a lengthening**  
865 **distributed over all phases of the cell cycle**

866 **A.** Quantification of interphase length of *WT* and *Seh1*<sup>-/-</sup> mESCs (2 distinct clones).  
867 The black bars represent the median and each dot represent one individual cell. The  
868 mean duration of interphase, as well as the number of imaged cells (N) and  
869 experiments [n] is indicated **B.** Cell cycle profiles of *WT* and two *Seh1*<sup>-/-</sup> clones  
870 generated by ImageStream® using the workflow analysis presented in **Fig. S2D**. For  
871 each cell line, at least 3000 cells acquired in at least 3 distinct experiments were

872 analyzed.

873

874 **Figure 3: Quantification of NPC density in *Seh1* mutant cell lines.**

875 Normalized signal intensities at the NE of Nup133 (**A**), Nup98 (**B**) and TPR (**C-D**) were  
876 quantified and box plots generated as described in Materials and Methods. The  
877 mean value was set at 1 for *WT* mESCs. For each cell line, the number of cells  
878 quantified (N), the number of distinct experiments [n], the mean value and standard  
879 deviation are indicated. In (**D**), cells were treated with EtOH (control) or Auxin as  
880 indicated.

881

882

883 **Figure 4: Quantification of nuclear sizes in *Seh1* mutant cell lines.**

884 Quantification of nuclear surface was performed as described in Materials and  
885 Methods. The mean value was set at 1 for *WT* mESCs. Graphs show average and  
886 standard deviation of nuclear surface values from [n] independent experiments  
887 (displayed as dots). Unless specified by lines, samples were compared to *WT* for  
888 statistical analyses (see Materials and Methods)

889

890 **Figure 5: *Mios*<sup>-/-</sup> mESCs do not mimic the growth and differentiation defects of  
891 *Seh1*<sup>-/-</sup> mESCs, nor their decreased NPC density and nuclear size.**

892 **A.** Whole cell extracts of *WT*, *Seh1*<sup>-/-</sup> (#1 and #2) and *Mios*<sup>-/-</sup> mESCs (three  
893 independent clones) were analyzed by western-blot using the indicated antibodies.  
894 Molecular masses are indicated on the right (kDa). **B.** Immunoprecipitation  
895 experiment using anti-*Seh1* antibodies and *WT*, *Mios*<sup>-/-</sup> (#1 and #2), or *Seh1*<sup>-/-</sup> (#1)  
896 mESC protein extracts. Inputs and eluates (20x equivalent) were analyzed by western  
897 blot using the indicated antibodies. Molecular markers are indicated on the right  
898 (kDa). **C-D.** Cell growth analyses (using percentage of confluence as proxy) were  
899 performed with the IncuCyte® system for *WT*, *Seh1*<sup>-/-</sup> and three distinct *Mios*<sup>-/-</sup>  
900 clones at pluripotent state (**C**), and upon neuroectodermal differentiation (**D**). Error  
901 bars correspond to the standard deviation arising from the indicated number of

902 independent experiments [n]. Statistical analyses were performed at the last time-  
903 points. Brackets indicate statistics performed using all values from cell lines bearing a  
904 given mutation, compared to *WT* (see Materials and Methods). **E-F.** Quantifications  
905 of TPR signal intensity at the NE (**E**, presented as box plots) and of the nuclear  
906 surface (**F**, graphs presenting the mean values and standard deviations from [n]  
907 distinct experiments, each displayed as a dot) were performed for *WT*, *Seh1*<sup>-/-</sup> and  
908 two *Mios*<sup>-/-</sup> clones as described in Materials and Methods. For each cell line, the total  
909 number of cells (N) acquired in [n] distinct experiments, and the mean and standard  
910 deviation values are indicated. For statistical analyses (see Materials and Methods)  
911 samples were compared to *WT*. Note that the mild (9%) increase in TPR density in  
912 *Mios*<sup>-/-</sup> # 1 mESCs was not observed for *Mios*<sup>-/-</sup> #2 cells and likely reflects a clonal-  
913 related variation not linked to the lack of Mios. Note that data for *WT* and *Seh1*<sup>-/-</sup>  
914 mESCs (used as reference strains) shown in panels C, D and F, include some data  
915 from experiments already presented in Figs. 1D-E and 4A.

916

917 **Figure 6: Impact of Y-complex short arm mutations on mESC proliferation and**  
918 **differentiation.**

919 **A.** Predicted model of human Nup43 (shown in purple), Nup85 (blue and yellow) and  
920 Seh1 (red) interactions (von Appen et al., 2015; PDB code: 5A9Q) visualized using  
921 Pymol. The  $\beta$ -sheets within the N-terminal domain of Nup85 that are deleted in the  
922  $\Delta$ E2-GFP/mCherry-Nup85 fusions are shown in yellow. **B.** Whole cell extracts of the  
923 indicated cell lines were analyzed by western-blot using anti-Seh1, -Nup43, -Nup85,  
924 and  $\gamma$ -tubulin antibodies. Two-and four fold dilution (1/2, 1/4) of the *WT* mESC  
925 extract were also loaded. Molecular markers are indicated on the right (kDa). **C-D.**  
926 Cell growth analyses were performed with the IncuCyte® system for *WT*, *Seh1*<sup>-/-</sup>,  
927  $\Delta$ E2-GFP-Nup85 and *Nup43*<sup>-/-</sup> mESCs at pluripotent state (**C**), and upon  
928 neuroectodermal differentiation (**D**). Error bars correspond to the standard deviation  
929 arising from [n] independent experiments. Statistical analyses were performed at the  
930 last time-points. Brackets indicate statistics performed using all values from cell lines  
931 bearing a given mutation, compared to *WT* (see Materials and Methods). Note that

932 data for *WT* and *Seh1*<sup>-/-</sup> (used as reference strains) shown in panels **C-D** include  
933 experiments already presented in Fig. 1 D-E.

934

935 **Figure 7: Quantification of NPC density and nuclear size in  $\Delta E2$ -GFP-*Nup85* and**

936 *Nup43*<sup>-/-</sup> mESCs lines.

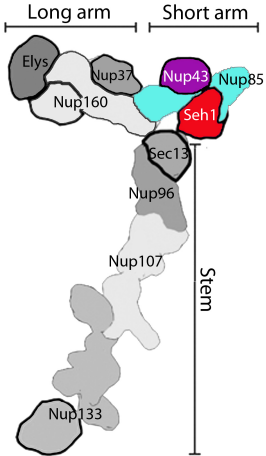
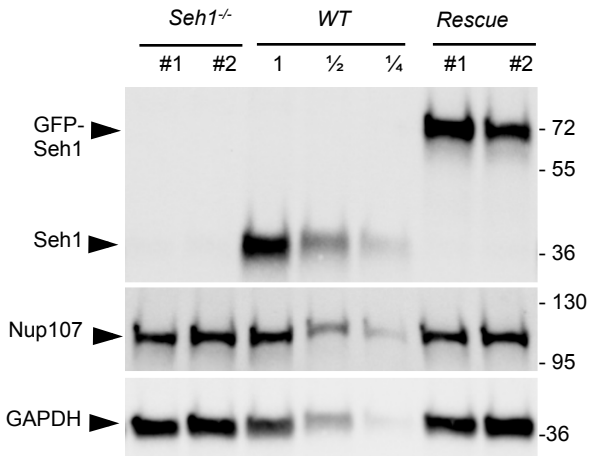
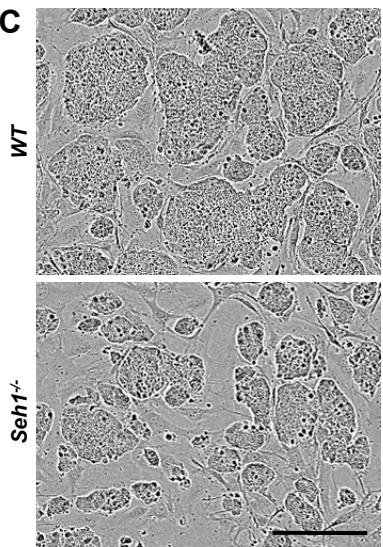
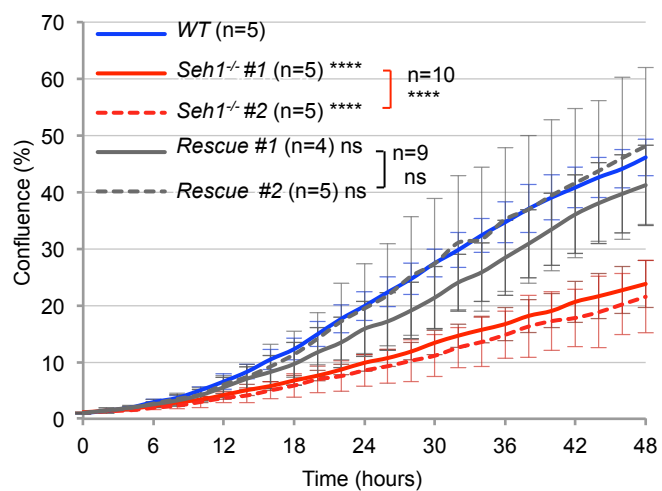
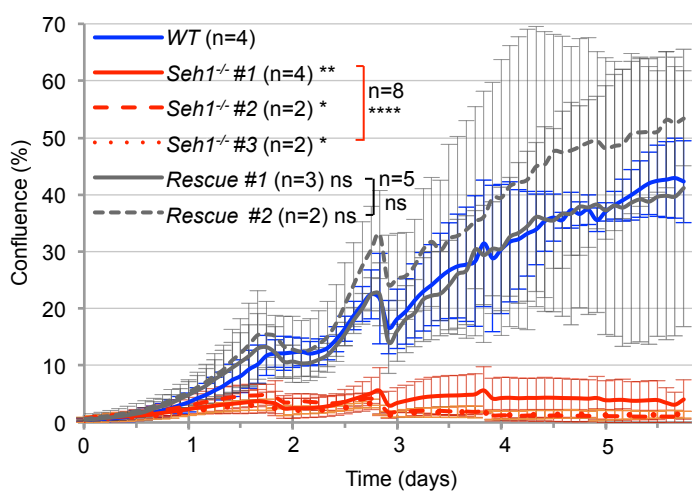
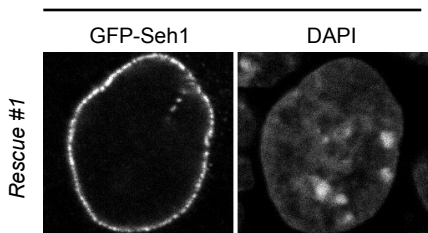
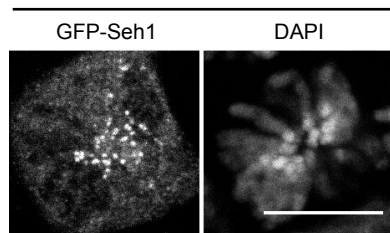
937 **A-C.** Normalized signals intensities at the NE of Nup133 (**A**), Nup98 (**B**) and TPR (**C**),  
938 (presented as box plots) and nuclear surfaces (**D**; graphs presenting the mean values  
939 and standard deviations from [n] distinct experiments, each displayed as a dot) were  
940 quantified as described in Materials and Methods. The number of cells (N), and of  
941 distinct experiments [n], the mean value and standard deviation are indicated. For  
942 statistical analyses (see Materials and Methods) samples were compared to *WT*.  
943 Note that data for *WT* and *Seh1*<sup>-/-</sup> (used as reference strains) include experiments  
944 already presented in Fig. 3.

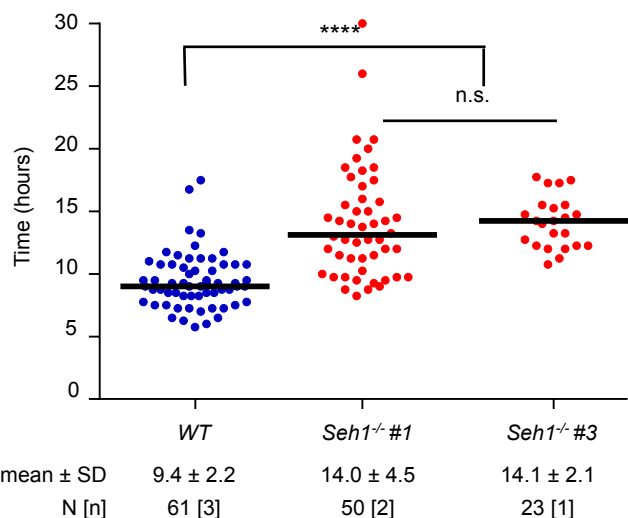
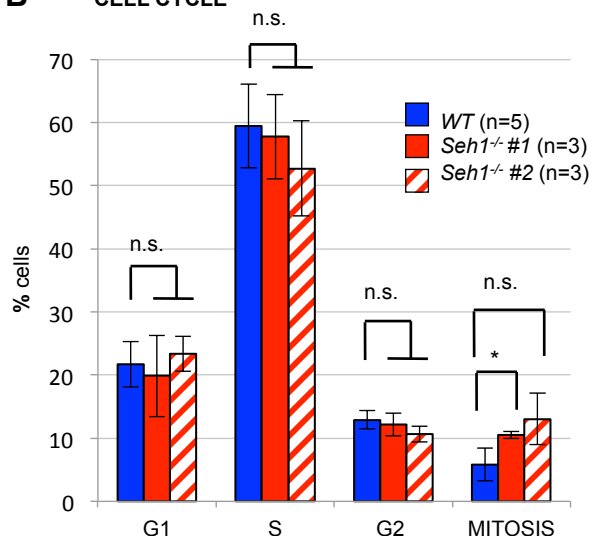
945

946 **Figure 8. Impaired GFP-Seh1 localization at NPCs and Kinetochores in  $\Delta E2$ -mCherry-**

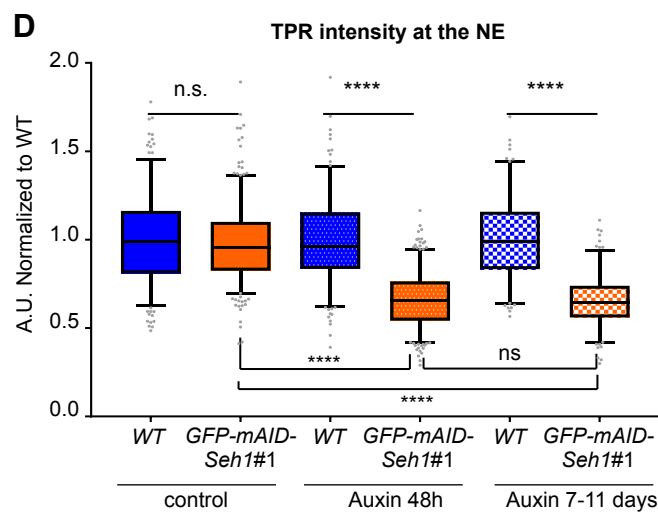
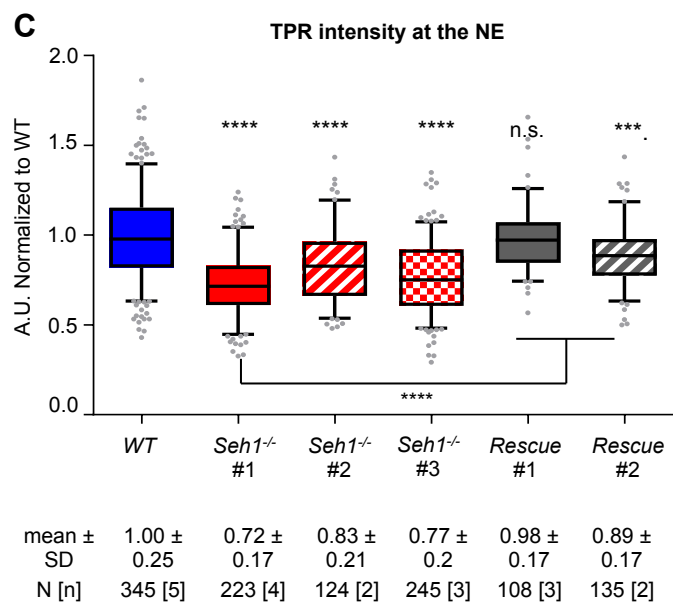
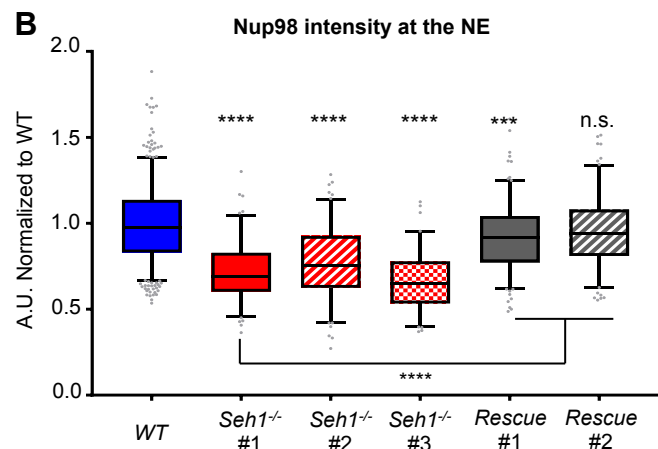
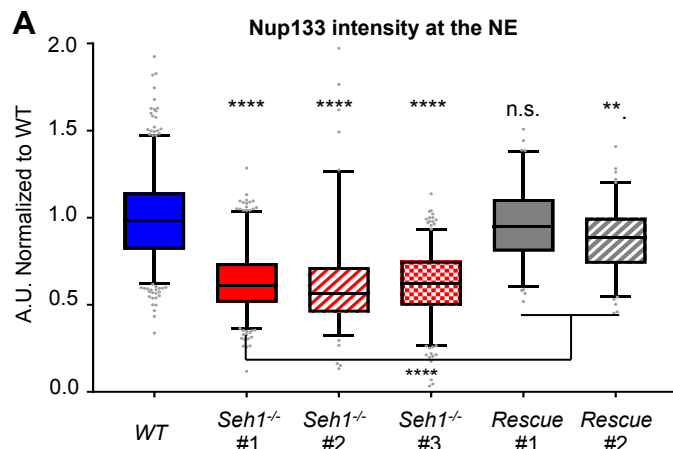
947 *Nup85* mESCs.

948 **A-B.** Representative spinning disk images (single z-section) of interphase (**A**) and  
949 mitotic (**B**) *GFP-Seh1* cells (left) mixed with  $\Delta E2$ -mCherry-*Nup85*/GFP-*Seh1* cells  
950 (right). Scale bars, 10  $\mu$ m. **C-D.** GFP-*Seh1* intensity at NE (**C**) and at kinetochores (**D**)  
951 was quantified in *GFP-Seh1* and  $\Delta E2$ -mCherry-*Nup85*/GFP-*Seh1* for the indicated  
952 number of cells (N) acquired in [n] independent experiments as described in  
953 Materials and Methods. Values were normalized for each field (**C**) or for each  
954 experiment (**D**) to the average intensity of the signal acquired for *GFP-Seh1* cells at  
955 the NE and kinetochores, respectively.

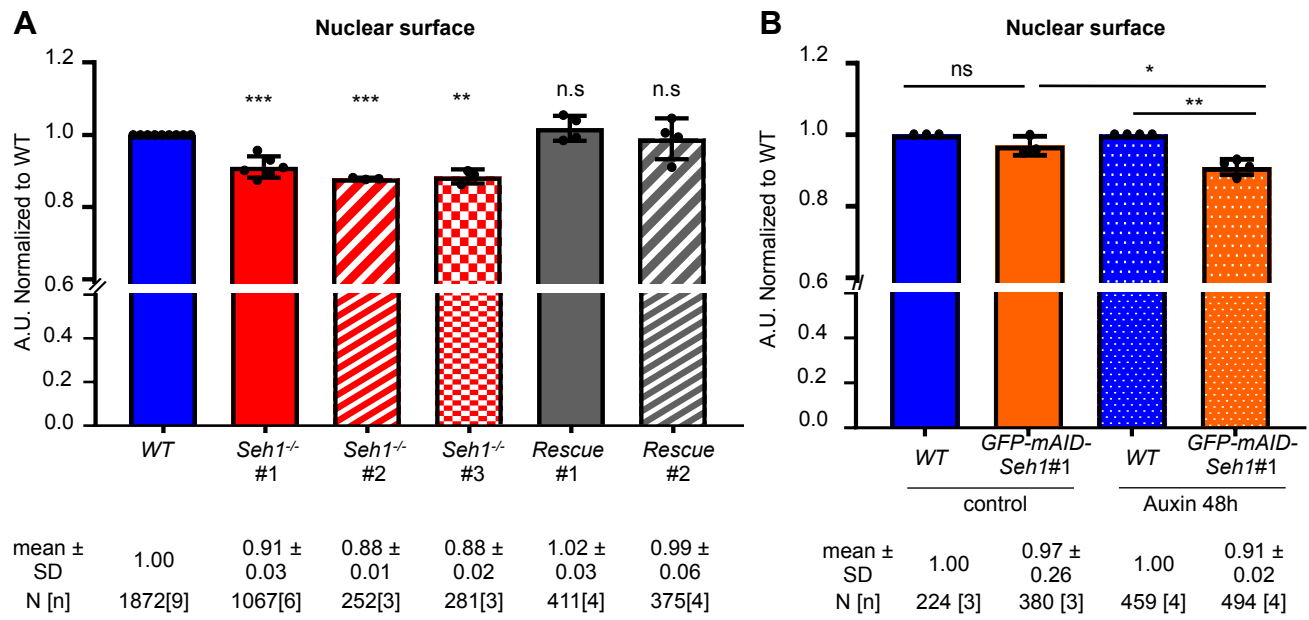
**A****B****C****D****PLURIPOTENT STATE****E****NEUROECTODERMAL DIFFERENTIATION****F****Interphase****Mitosis**

**A****INTERPHASE LENGTH****B****CELL CYCLE**

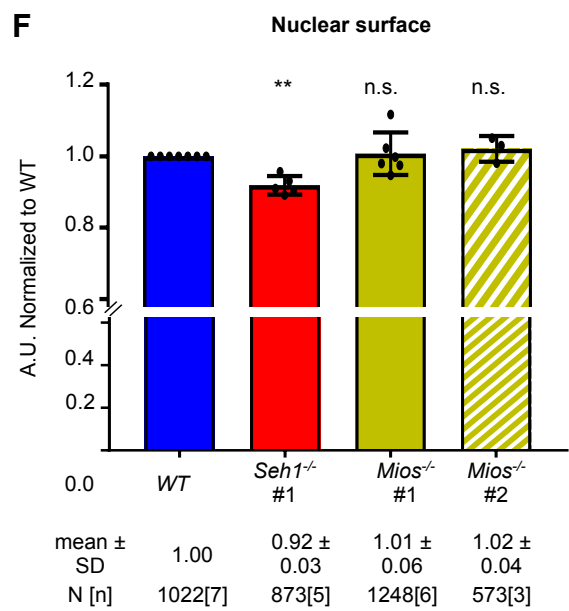
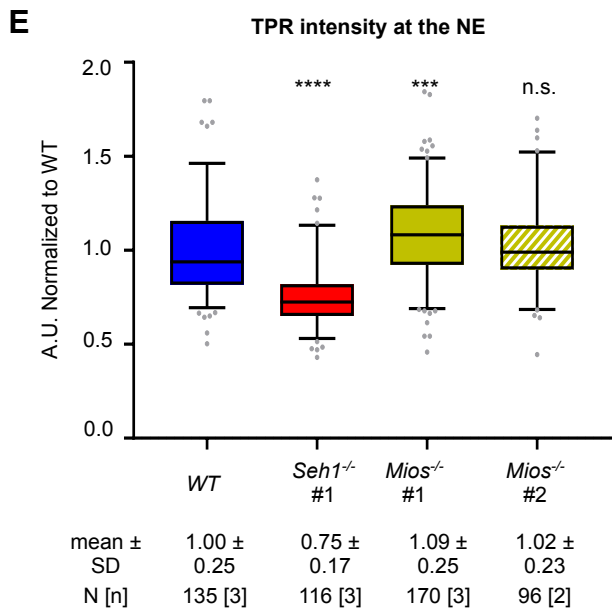
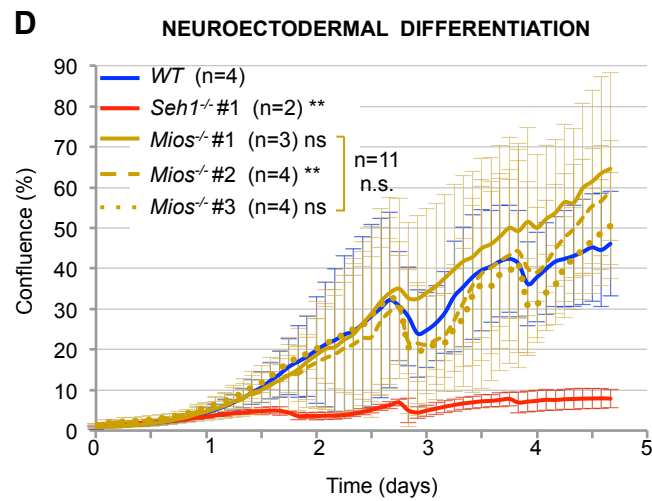
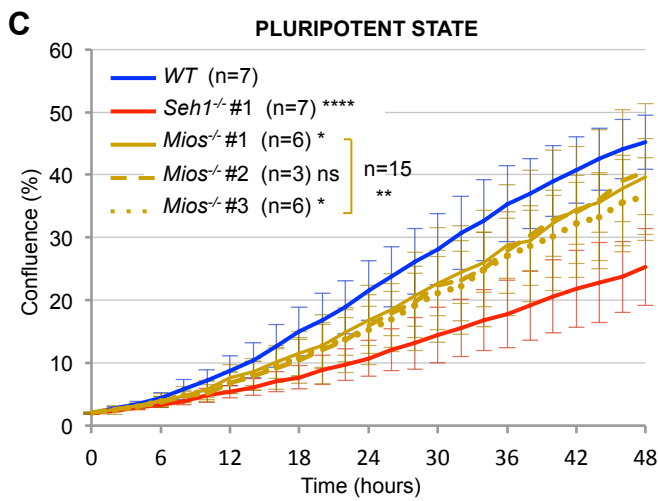
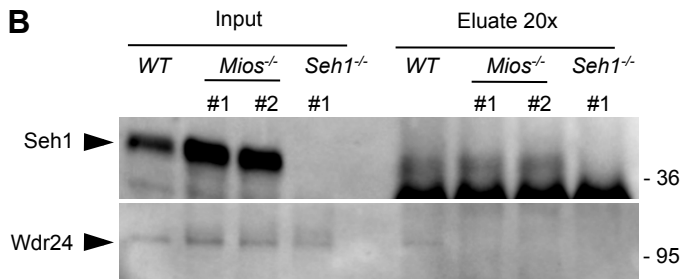
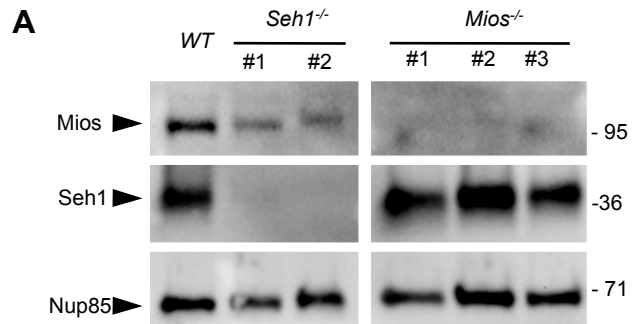
**Gonzalez-Estevez, Verrico, et al., Figure 2**



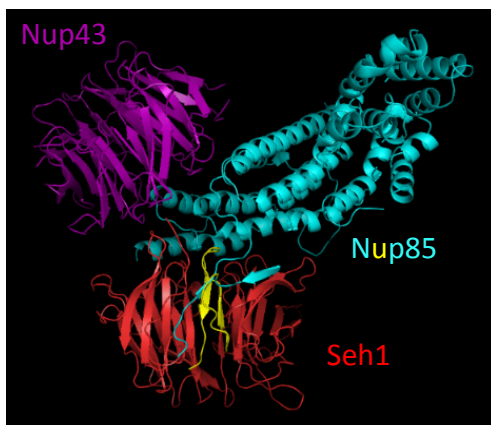
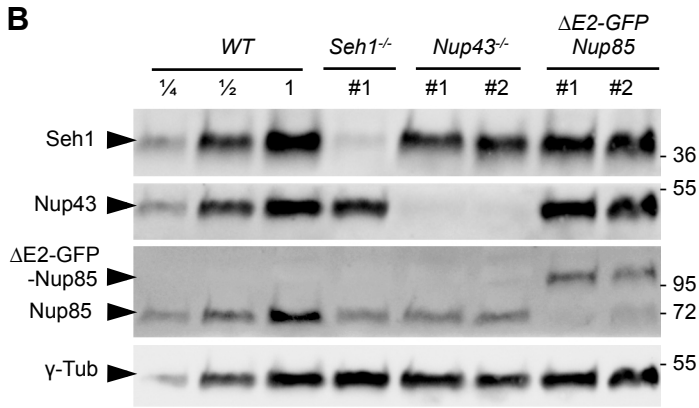
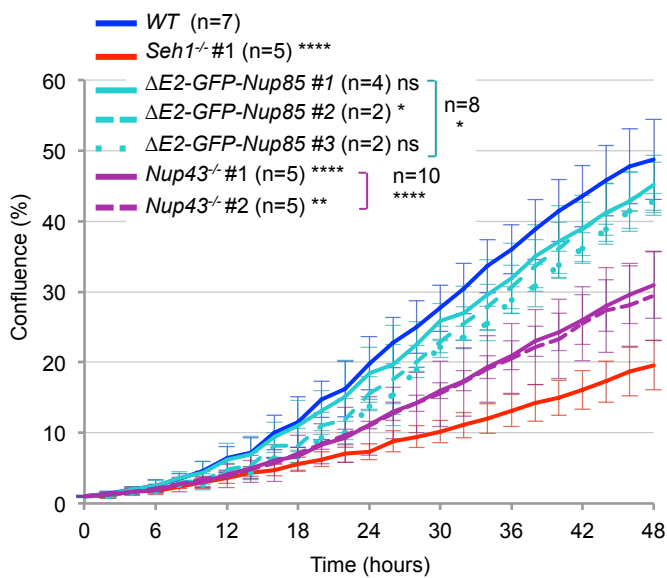
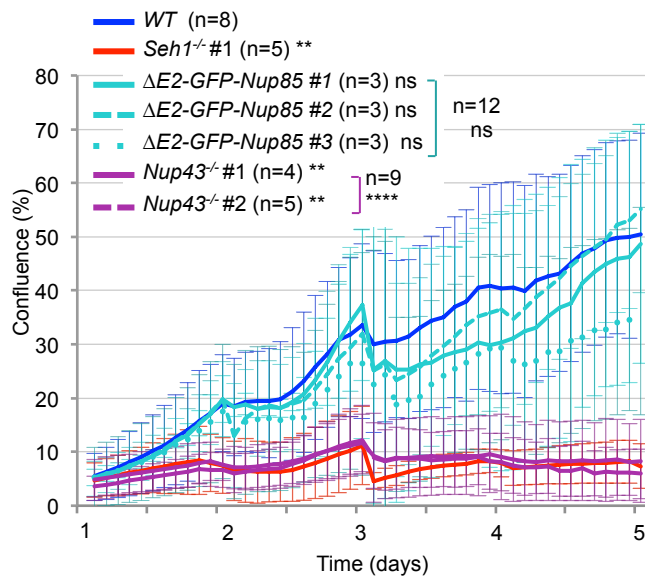
Gonzalez-Estevez, Verrico, et al., Figure 3

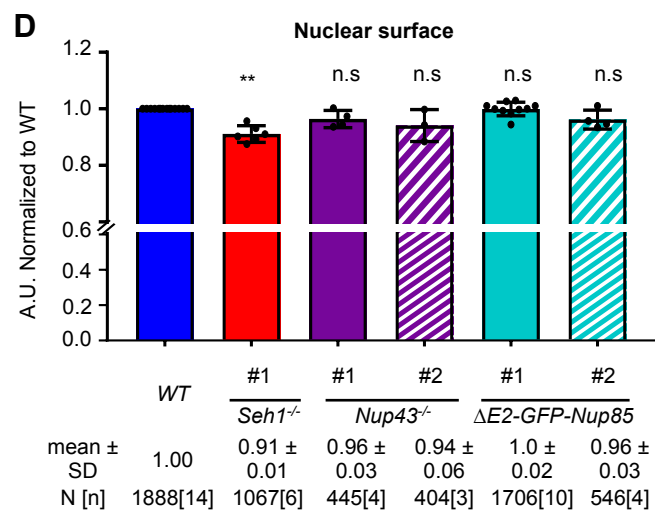
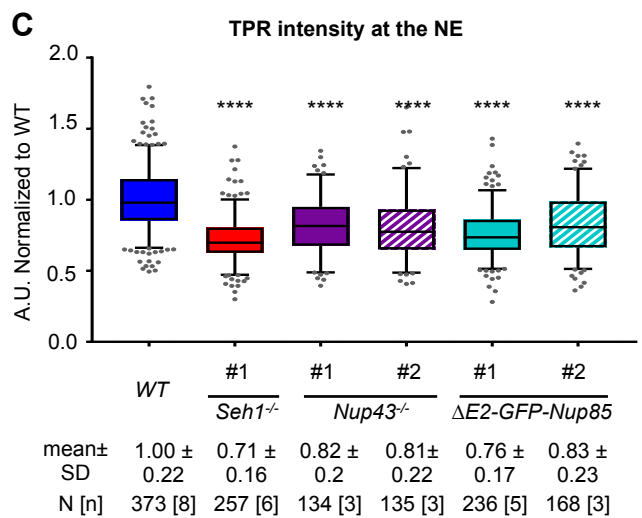
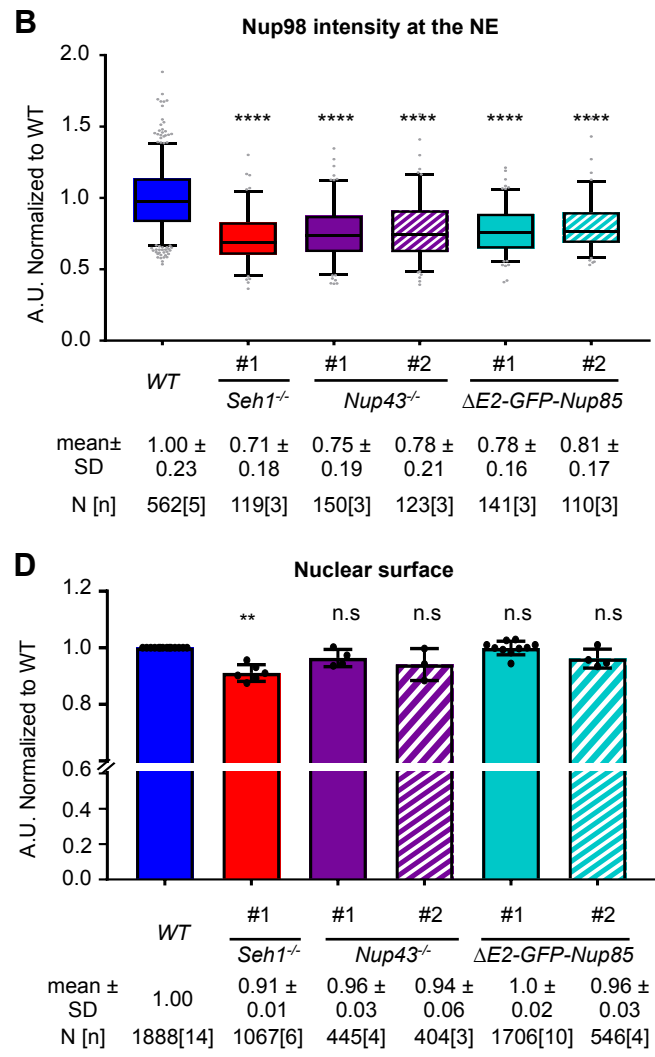
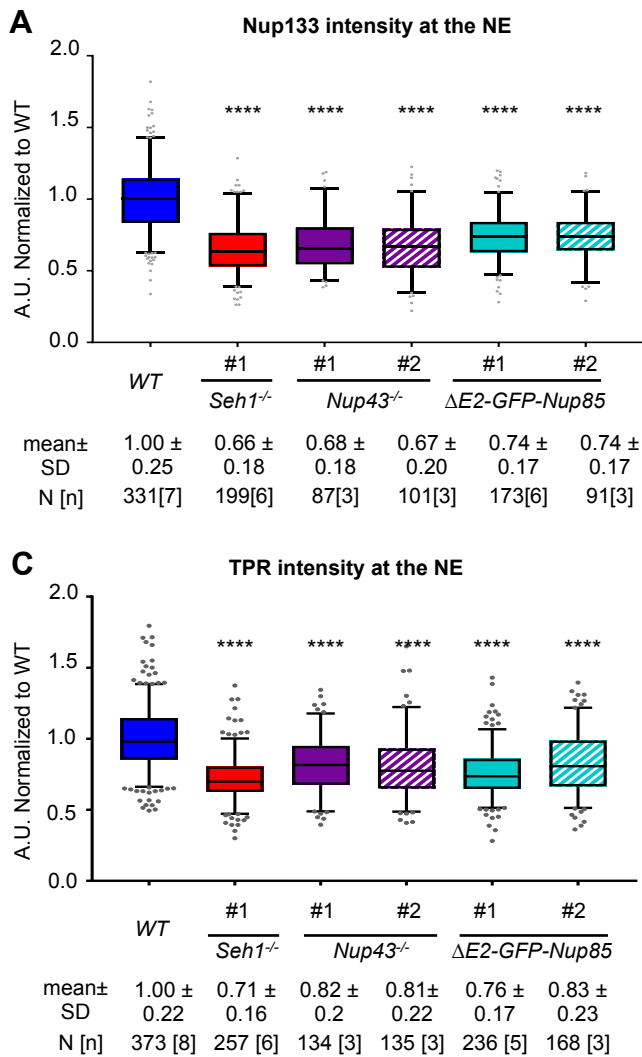


Gonzalez-Estevez, Verrico, et al., Figure 4

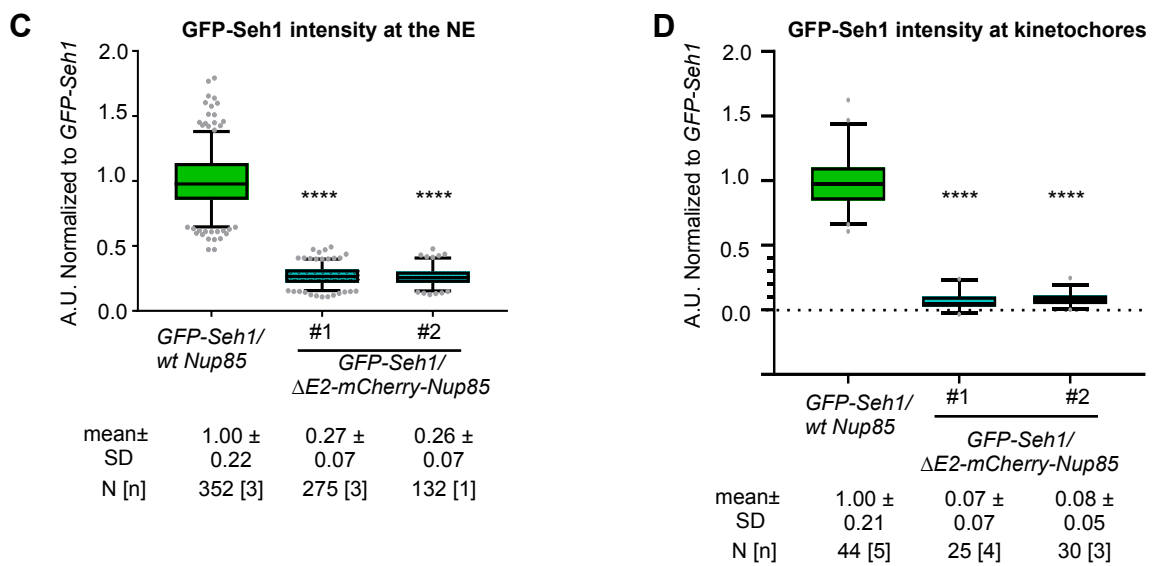
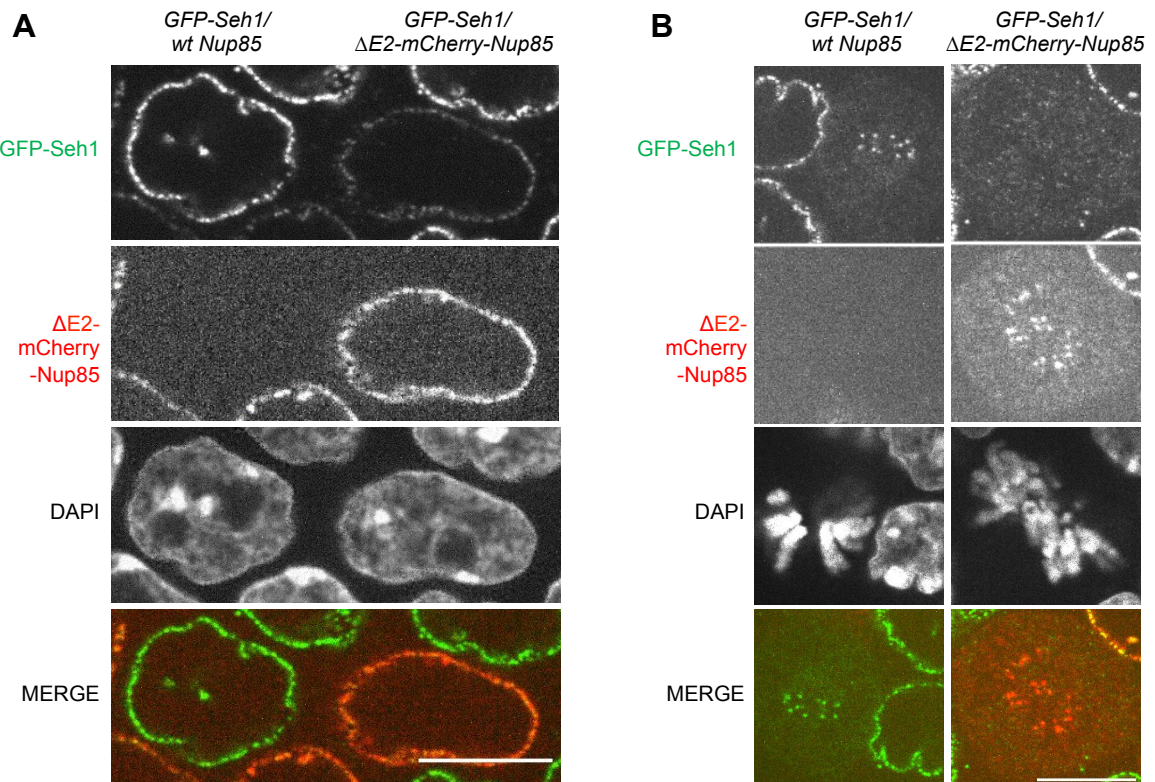


Gonzalez-Estevez, Verrico, et al., Figure 5

**A****B****C****PLURIPOtent STATE****D****NEUROECTODERMAL DIFFERENTIATION****Gonzalez-Estevez, Verrico, et al., Figure 6**



Gonzalez-Estevez, Verrico, et al., Figure 7



Gonzalez-Estevez, Verrico, et al., Figure 8

## Inventory of Supplementary Material

### Supplemental Figures and Figure legend:

**Figure S1, related to Fig. 1:** Auxin-induced GFP-mAID-Seh1 degradation in mESCs recapitulates *Seh1*<sup>-/-</sup> phenotypes

**Figure S2, related to Fig. 2:** *Seh1*<sup>-/-</sup> mESCs have prolonged mitosis despite proper localization of the Y-complex at kinetochores

**Figure S3, related to Figs. 6-8:** Characterization of the  $\Delta E2$ -GFP-*Nup85* and  $\Delta E2$ -*mCherry-Nup85* cell lines

**Figure S4, related to Materials and Methods and Tables S2 and S3:** Strategies for CRISPR-Cas9-mediated cell line establishment via homologous recombination.

### Supplemental Tables:

**Table S1:** Plasmids used in this study

**Table S2:** Cell lines used in this study

**Table S3:** Sequences of gRNAs and Primers used in this study

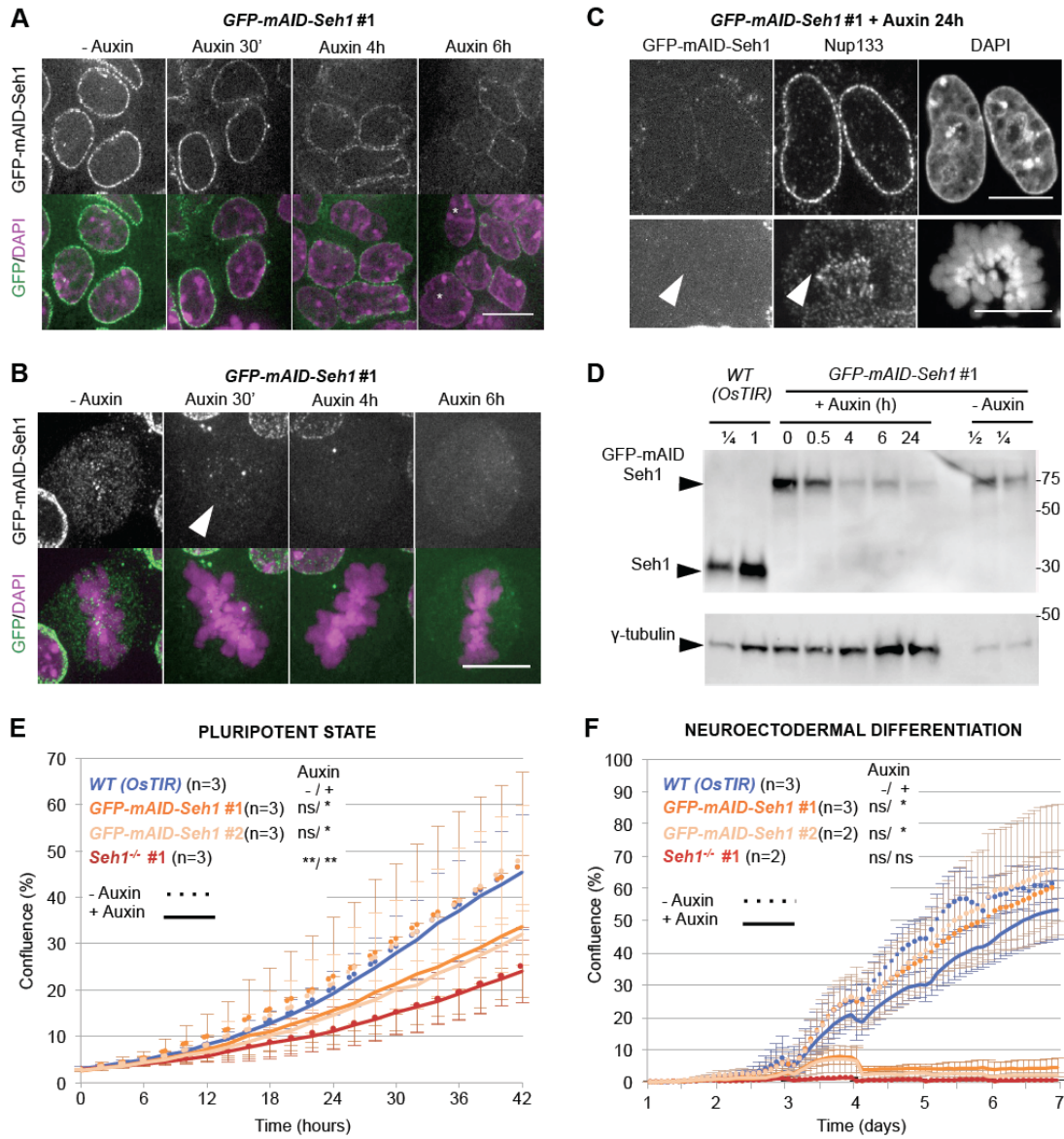
**Table S4:** Antibodies used in this study

### References cited in Supplemental Figures and Tables

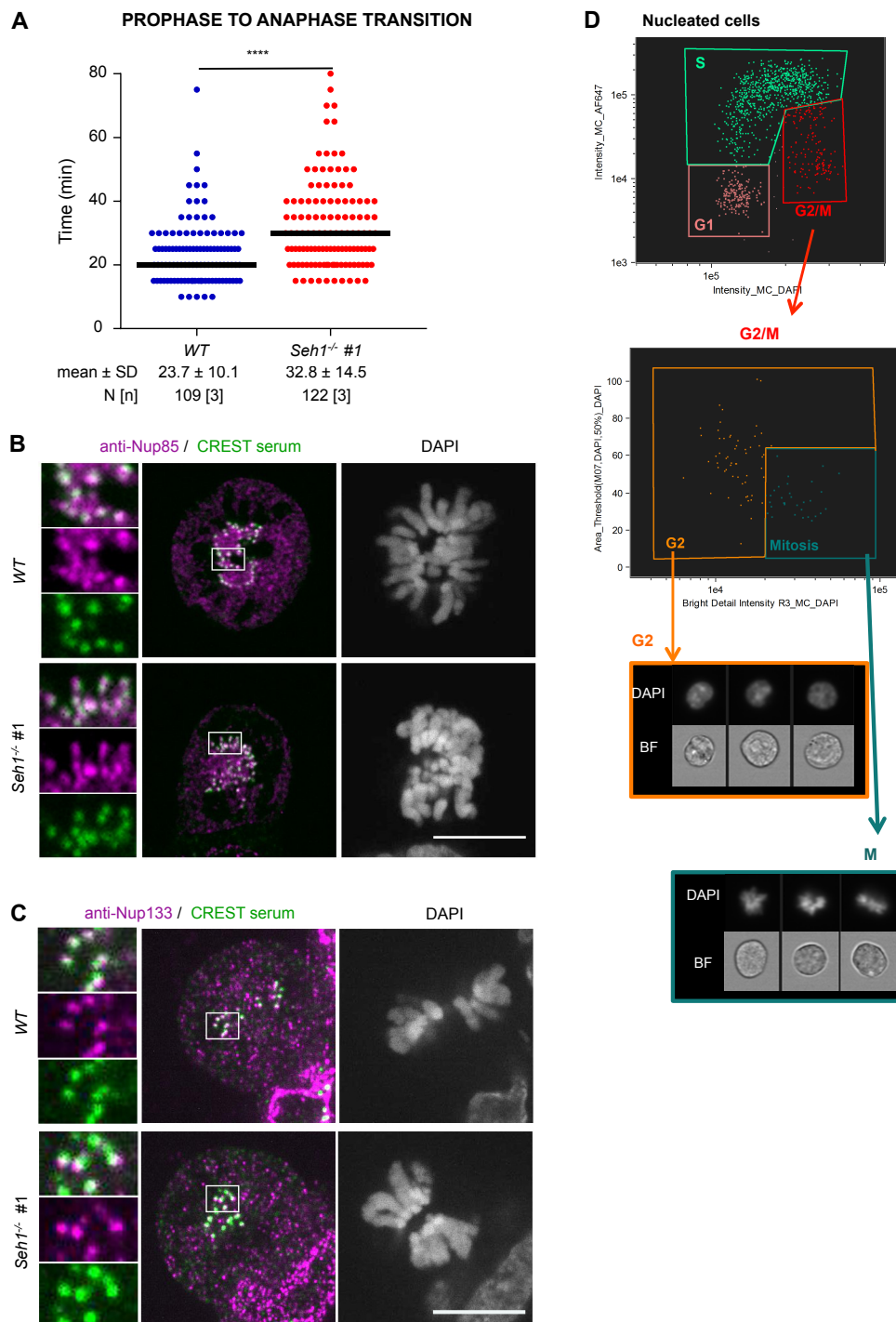
### Supplemental Movies, related to Figure 1:

Examples of neuroectodermal differentiation experiments as exported from IncuCyte® device. Time and scale bars are indicated

- Movie S1: *WT*
- Movie S2: *Seh1*<sup>-/-</sup> #1
- Movie S3: *Rescue* #2

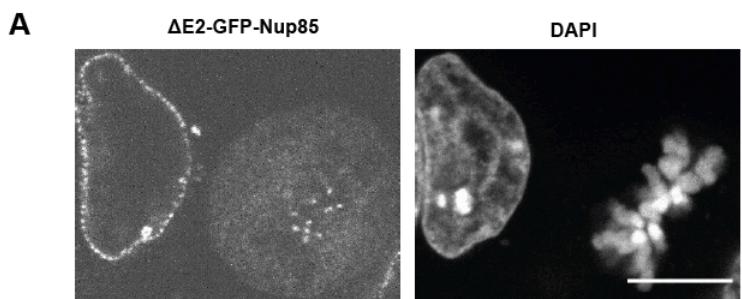


**Figure S1, related to Fig. 1. Auxin-induced GFP-mAID-Seh1 degradation in mESCs recapitulates *Seh1*<sup>-/-</sup> phenotypes. A-B.** GFP-mAID-Seh1 localizes at NE in interphase cells (A; a single confocal plane is presented) and is enriched at kinetochores in mitotic cells (B; a projection of 9 optical sections is presented). After 30 minutes of auxin treatment, the GFP signal is only slightly decreased at the NE in interphase cells but no longer detectable in mitotic cells (arrowhead). Longer treatments (4 to 6 hours) with auxin lead to a progressive decrease of the NE signal, and to the appearance of GFP negative cells (\*, likely corresponding to cells that went through mitosis). Scale bars, 10 μm. **C.** Localization of Nup133 in interphase (upper panels) and mitotic (lower panels) *GFP-mAID-Seh1* #1 mESCs after 24h of auxin treatment. The arrowheads point to the kinetochores that are labelled by Nup133 despite the lack of GFP-mAID-Seh1. Scale bar, 10 μm. **D.** Whole cell extracts from *WT* (*OsTIR*) and *GFP-mAID-Seh1* #1 mESCs, treated or not with auxin for the indicated time, were analyzed by western blot using anti-Seh1 (top) and an anti-gamma-tubulin antibodies (bottom, used as loading control). 1/2 and 1/4 dilutions of *WT* (*OsTIR*) and non-treated *GFP-mAID-Seh1* #1 cell extracts were also loaded. Molecular weights are indicated (kilodaltons). **E.** Cell growth at pluripotent stage of the indicated cell lines was analyzed using cell confluence measurements with the IncuCyte® system. Cells were treated with Auxin for 18 to 26 h prior to time 0. Error bars correspond to standard deviations from 3 independent experiments. **F.** The % of cell confluence upon neuroectodermal differentiation was quantified with the IncuCyte® system. Cells were exposed to Auxin at the beginning of differentiation (day 0). Error bars correspond to standard deviations of 4 (*GFP-mAID-Seh1* #1 ad *OsTIR* WT) or 3 (*GFP-mAID-Seh1* #2 and *Seh1*<sup>-/-</sup> #1) distinct wells acquired in 3 and 2 independent experiments, respectively. Statistical analyses were performed using the two-tailed Paired Student's t-test (See Materials and Methods), by comparing all control (-Auxin) or treated (+Auxin) mutant cell lines to *WT* (*OsTIR*) -Auxin or +Auxin, respectively.



**Figure S2, related to Fig. 2: *Seh1*<sup>-/-</sup> mESCs have prolonged mitosis despite proper localization of the Y-complex at kinetochores**

**A.** Quantification of the time spent from prometaphase to anaphase in *WT* and *Seh1*<sup>-/-</sup> mESCs. Each dot represents one cell and the median time is represented with a black bar. The number (N) of cells, quantified in 3 independent experiments is indicated. **B, C.** Representative spinning-disk images of mitotic *WT* and *Seh1*<sup>-/-</sup> mESCs, immunolabeled with (**C**) anti-Nup85 or (**D**) anti-Nup133, along with CREST serum (centromere marker) and stained with DAPI. A projection of three Z sections is presented. 3-fold enlargements of the boxed areas are also presented. Scale bars, 10  $\mu$ m. **D.** Representative plots of the last steps of our ImageStream® cell cycle analyses (*WT* mESCs). Top: the G1, S and G2/M phases of the cell cycle were defined based on DNA content (DAPI intensity) and EdU incorporation (AF647 channel). Middle: the G2 and M (Mitotic) population were then discriminated based on DAPI intensity and area. Bottom: bright Field (BF) and corresponding DAPI images of representative cells defined by these gating as G2 (orange) or mitotic (blue) are presented.

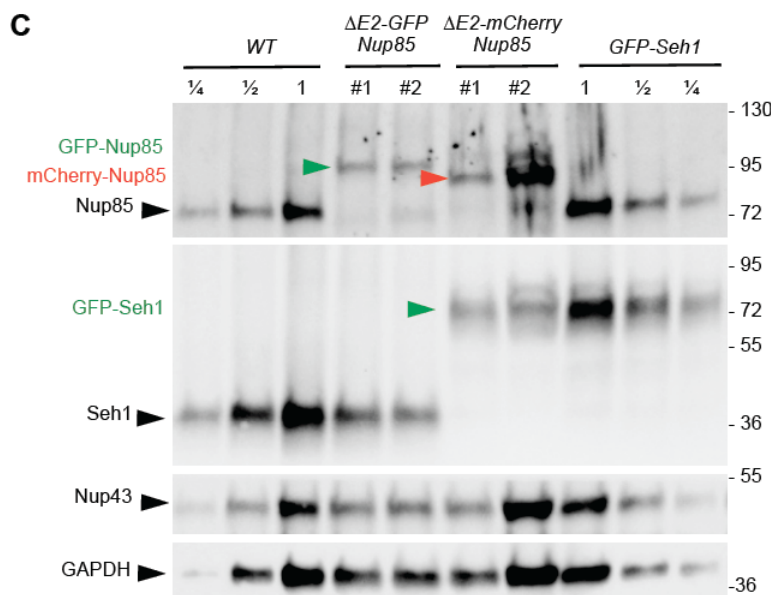


**B**

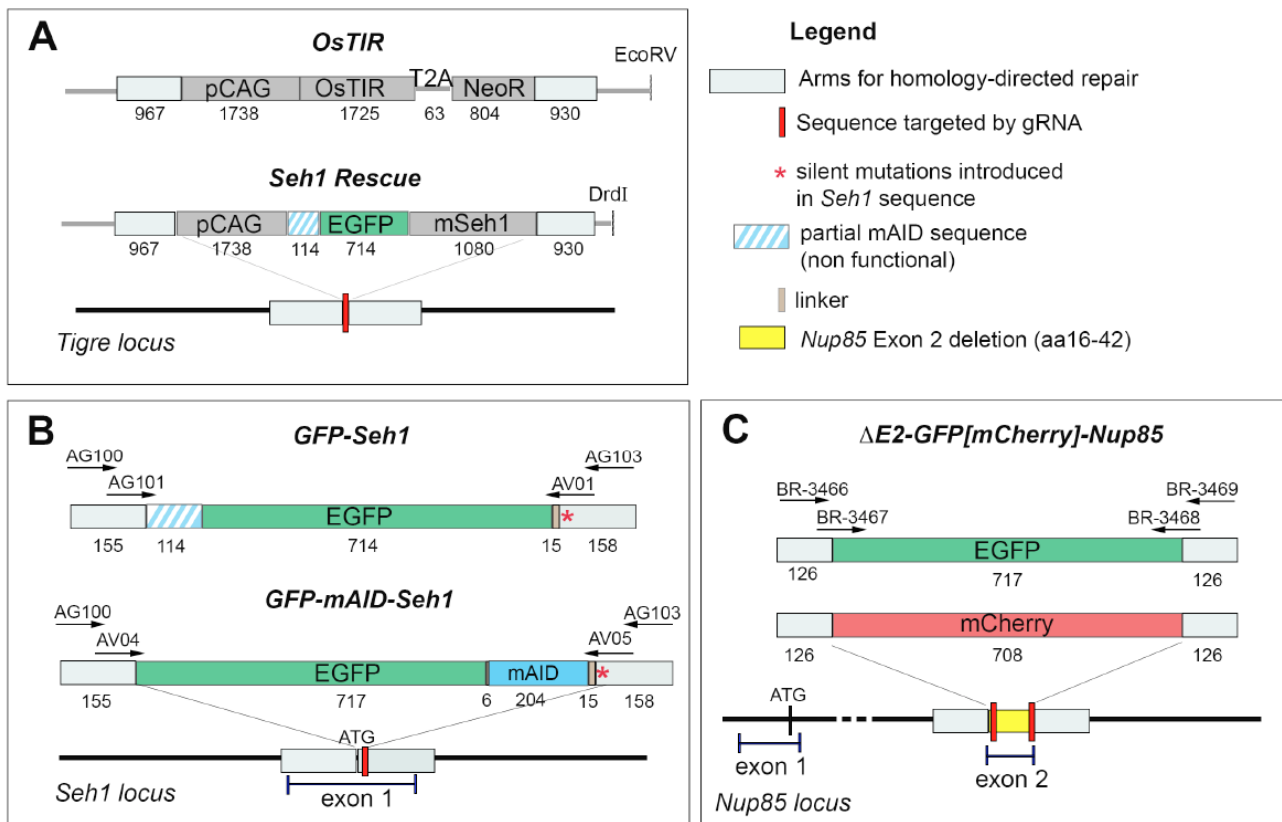
Antibody : control      anti-Nup107      anti-Nup85

Cells : WT      WT      ΔE2- GFP- Nup85      WT      ΔE2- GFP- Nup85

Y-complex Nups <sup>a</sup>	Cells :	control	anti-Nup107	anti-Nup85	
		WT	WT	ΔE2- GFP- Nup85	
Seh1l			106	34	
Nup85			185	215	431
Nup43			55	19	56
Nup160			340	125	255
Nup37			61	34	25
Nup133			536	349	345
Nup107			192	99	95
Nup98-96 <sup>b</sup>	16 <sup>c</sup>		139	200	204
Sec13			27	15	24
					60



**Figure S3, related to Figs 6-8. Characterization of the ΔE2-GFP-Nup85 and ΔE2-mCherry-Nup85 cell lines.** (A) Representative spinning disk images (single plane) of interphase (left) and mitotic (right) ΔE2-GFP-Nup85 mESCs fixed and stained with DAPI. Scale bar, 10 μm. (B) The table provides the Mascot score for an immunoprecipitation experiment (see Materials and Methods) using either a pre-immune serum (control) or anti-Nup107 or -Nup85 antibodies, to pull down the Y-complex from WT or ΔE2-GFP-Nup85 mESC protein extracts. <sup>a</sup>Elys/AHCTF1 was not identified in this immunoprecipitation experiment. <sup>b</sup>The autoproteolytic cleavage of the Nup98-Nup96 precursor generates Nup98 and Nup96 (Fontoura et al., 1999); only the latter is a stable component of the Y-complex. Nearly all (35 out of 36) peptides identified in these immune pellets originates from Nup96. <sup>c</sup> Samples in which the Nup98 peptide was identified. (C) Whole cell extracts of the indicated cell lines were analyzed by western-blot using anti-Nup85, -Seh1, -Nup43, and -GAPDH antibodies. Two-and four fold dilution (1/2, 1/4) of the WT and GFP-Seh1 mESC extracts were also loaded. Molecular markers are indicated on the right (kDa).



**Figure S4, related to Materials and Methods and Tables S2 and S3: Strategies for CRISPR-Cas9-mediated cell line establishment via homologous recombination.**

The constructs used for homologous recombination (A, plasmid digested with the indicated restriction enzyme, and B-C, PCR products: the sequences of the oligonucleotides (arrows) used to amplify the template are listed in **Table S3**) leading to the indicated cell lines are represented above each targeted genomic locus (A: *Tigre*, B: *Seh1*, and C: *Nup85*). The sequences targeted by the gRNAs (red boxes), the sequences used for homology-directed repair (light grey boxes), and when relevant, the position of the exons and the first ATG are displayed. The size of the various segments is indicated in bp. Note that the construct used for GFP-Seh1 expression (A. *Seh1* rescue and B. *GFP-Seh1* lines) further contains a non-functional 38 amino-acid long fragment of mAID. In B., the red star indicates the position of silent CRISPR/Cas9 blocking mutations introduced in one *Seh1* HR-arm.

**Table S1: Plasmids.**

Plasmids used (sequence upon request)	Source	Identifier
pX-280_Cas9WT-EGFP-Seh1gRNA1 - Seh1gRNA 2	This paper	#1808
pU6-sgTigre_CBh-Cas9-T2A-mCherry-3UTR	From P. Navarro Gil and N. Festuccia. (Festuccia et al., 2019)	#2061
<i>Tigre</i> HR-pCAG-EGFP-mSeh1- <i>Tigre</i> HR	This paper	#2077
<i>Tigre</i> HR-pCAG-OsTir-T2A-NeoR- <i>Tigre</i> HR	From P. Navarro Gil and N. Festuccia. (Festuccia et al., 2019)	#2064
pX-853_Cas9WTmCherry Seh1-gRNA1	This paper	#2098
pCMV-EGFP-mAID-hSeh1	This paper	#2100
GeneArt CRISPR Nuclease Vector (OFP_Cas9)	Invitrogen, Carlsbad, CA	A21174
GeneArt CRISPR pOFP-Cas9-Mios gRNA2	This paper	#2110
GeneArt CRISPR pOFP-Cas9-Mios gRNA4	This paper	#2111
GeneArt CRISPR pOFP-Cas9-Nup43 gRNA1	This paper	#2104
pX-672-Cas9HF-mCherry-gRNANup85-1_gRNANup85-2	This paper	#2005
pX-864-Cas9HF-GFP-gRNANup85-1_gRNANup85-2	This paper	#2101
pBOS-H2B-Cherry-IRES-Neo	From Doye lab (Bolhy et al., 2011)	#824
pBOS-H2B-GFP-IRES-Neo	From Doye lab (Bolhy et al., 2011)	#805

**Table S2, related to Experimental Procedures. Cell lines used in this study**

NAME	SOURCE (IF COMMERCIAL) OR DESIGN FOR CELL LINES GENERATED IN THIS STUDY	IDENTIFIER	Plate ref.	MUTATIONS ( genomic sequences set at 1 for the first ATG )	Chromosome count
MEF (DR4)	Applied StemCell	ASF-1001	-	-	-
WT mESCs (HM1)	ThermoFisher Scientific (Selfridge et al., 1992)	MES4303	-	-	40
<i>Seh1</i> <sup>-/-</sup>	<u>Parental cell line:</u> HM1 <u>Cas9 &amp; gRNA plasmid:</u> pX-280_Cas9WT-EGFP-Seh1gRNA1 - Seh1gRNA 2 <u>HR template:</u> (200 bp synthetic oligonucleotide initially designed to facilitate screening)	<i>Seh1</i> <sup>-/-</sup> #1	A18	Only one PCR band observed suggesting that both alleles bear the same mutation. <u>Allele 1 (and 2):</u> Deletion of 351bp from -317 to +33. Insertion of 32bp (from HR template) at +71.	40 Chr (>60%)
		<i>Seh1</i> <sup>-/-</sup> #2	B42	Allele 1: 1bp indel (c>a) at -2 plus Deletion of 77bp from +4 to +77. Insertion of 31 bp (from HR template) at +4. Allele 2: Deletion of 30bp from -2 to +27; one bp deletion at +70.	40 Chr (>60%)
		<i>Seh1</i> <sup>-/-</sup> #3	B37	Allele 1: Deletion of 52 bp from +19 to +71 Allele 2: Deletion of 15bp from +13 to +27; one bp deletion at +70.	40 Chr (>60%)
<i>Seh1-Rescue</i>	<u>Parental cell line:</u> <i>Seh1</i> <sup>-/-</sup> #1 mESCs <u>Cas9 &amp; gRNA plasmid:</u> pU6-sgTIGRE_Cbh-Cas9-T2A-mCherry-3UTR <u>HR template:</u> linearized (with DrdI) TIGRE HR-pCAG-EGFP-mSeh1-TIGRE HR (#2077)(Figure S4)	<i>Seh1-Rescue</i> #1	R11G	Rescue of <i>Seh1</i> <sup>-/-</sup> #1 (A18). Insertion of pCAG-GFP-Seh1 in only one of the <i>TIGRE</i> alleles	40 Chr (55%) 39 Chr (45%)
		<i>Seh1-Rescue</i> #2	R11F	Rescue of <i>Seh1</i> KO A18 (#1). Insertion of pCAG-GFP-Seh1 in only one of the <i>TIGRE</i> alleles	40 Chr (>60%)
<i>OsTIR</i>	<u>Parental cell line:</u> HM1 <u>Cas9 &amp; gRNA plasmid (s):</u> pU6-sgTIGRE_Cbh-Cas9-T2A-mCherry-3UTR <u>HR template:</u> linearized (with EcoRV) TIGRE HR-pCAG-OsTir-T2A-NeoR-TIGRE HR (#2064) (Figure S4)	<i>OsTIR</i>	11B	Insertion of pCAG-OsTir-T2A-NeoR in only one of the <i>TIGRE</i> alleles.	42Chr (>60%)

**Table S2- continued**

NAME	DESIGN OF THE CELL LINES	IDENTIFIER	Plate ref.	MUTATIONS (genomic sequences set at 1 for the first ATG)	Chromosome count
<i>GFP-mAID-Seh1</i>	<u>Parental cell line:</u> <i>OsTIR</i> <u>Cas9 &amp; gRNA plasmid (s):</u> <b>pX-853</b> <u>Cas9WTmCherry Seh1-gRNA1</u> <u>HR template:</u> PCR product: HR-EGFP-mAID-mSeh1 (gRNA resistant)-HR; amplification of plasmid #2100 with primers indicated in <b>Figure S4</b> and <b>Table S3</b>	<i>GFP-mAID-Seh1</i> #1	F9	<u>Alleles 1 and 2:</u> GFP-mAID-Seh1	42Chr (>60%)
		<i>GFP-mAID-Seh1</i> #2	E6	<u>Allele 1:</u> GFP-mAID Seh1 <u>Allele 2:</u> <i>Seh1</i> KO; 1 bp insertion at + 18	42Chr (>60%)
<i>Mios</i> <sup>-/-</sup>	<u>Parental cell line:</u> HM1/ <u>Cas9 &amp; gRNA plasmid (s):</u> GeneArt CRISPR pOFP-Cas9-Mios gRNA2 GeneArt CRISPR pOFP-Cas9-Mios gRNA4 <u>HR template:</u> none	<i>Mios</i> <sup>-/-</sup> #1	5C	Only one PCR band observed suggesting that both alleles bear the same mutation. <u>Allele 1 (and 2):</u> Deletion of 593 bp from -310 to 282.	40 Chr (>60%)
		<i>Mios</i> <sup>-/-</sup> #2	7C	<u>Allele 1:</u> Deletion of 33bp from +74 to 106. <u>Allele 2:</u> Deletion of 3 bp from +84 to +86	40 Chr (>60%)
		<i>Mios</i> <sup>-/-</sup> #3	7D	<u>Allele 1:</u> Deletion of 3bp from +83 to 85 and of 7 bp from 195 to 201. <u>Allele 2:</u> Replacement from bp 85 C by GT (=1bp) and insertion of a T after bp 196.	40 Chr (>60%)
<i>ΔE2-GFP-Nup85</i>	<u>Parental cell line:</u> HM1 <u>Cas9 &amp; gRNA plasmid (s):</u> <b>pX-672</b> -Cas9HF-mCherry-gRNANup85-1_gRNANup85-2 <u>HR template:</u> PCR product: HR-EGFP-HR ( <b>Figure S4</b> ); amplification of EGFP with primers indicated in Figure S4 and Table S3	<i>ΔE2-GFP-Nup85</i> #1	B2	<u>Allele 1:</u> In frame integration of GFP in exon 2 and deletion from bp 639 to 716. <u>Allele 2:</u> <i>Nup85</i> KO; deletion of 2bp at +633-634 leading to premature stop codon.	41 Chr (56%) 40 Chr (25%)
		<i>ΔE2-GFP-Nup85</i> #2	B10	<u>Allele 1:</u> In frame integration of GFP in exon 2 and deletion from bp 639 to 716. <u>Allele 2:</u> <i>Nup85</i> KO; Deletion of 20 bp from 622 to 641.	42 Chr (57%) 41 Chr (29%) 40 Chr (14%)
		<i>ΔE2-GFP-Nup85</i> #3	B3	<u>Allele 1:</u> In frame integration of GFP in exon 2 and deletion from bp 639 to 716. <u>Allele 2:</u> <i>Nup85</i> KO; deletion of 13 bp from 627 to 647	42 Chr (>60%)

**Table S2- continued**

NAME	DESIGN OF THE CELL LINES	IDENTIFIER	Plate ref.	MUTATIONS (genomic sequences set at 1 for the first ATG)	Chromosome count
<b>GFP-Seh1</b>	<u>Parental cell line:</u> HM1 <u>Cas9 &amp; gRNA plasmid (s):</u> pX-853 <u>_Cas9WTmCherry Seh1-gRNA1</u> <u>HR template:</u> PCR product HR_EGFP-mSeh1 (gRNA resistant)-HR. Amplification of plasmid #2077 with primers indicated in <b>Figure S4</b> and <b>Table S3</b>	<i>GFP-Seh1</i>	B10	<u>Allele 1:</u> <i>GFP-Seh1</i> . In frame N-term integration of GFP at +4 in <i>Seh1</i> sequence. <u>Allele 2:</u> <i>Seh1</i> KO. Deletion from -2 to +27.	40 Chr (>60%)
<b>ΔE2-mCherry-Nup85</b>	<u>Parental cell line:</u> <i>GFP-Seh1</i> <u>Cas9 &amp; gRNA plasmid (s):</u> pX-864-Cas9HF-GFP-gRNANup85-1_gRNANup85-2 <u>HR template:</u> PCR product HR-mCherry-HR ( <b>Figure S4</b> ); amplification of mCherry with primers indicated in Figure S4 and Table S3	<i>ΔE2-mCherry-Nup85</i> #1	B6	<u>Allele 1:</u> in frame mCherry integration in exon2 and deletion from bp 639 to 716. <u>Allele 2:</u> <i>Nup85</i> KO. Deletion of 8bp (+630 -- +638) and of 10 bp (+706 to +716) in exon 2.	41 Chr (>60%)
		<i>ΔE2-mCherry-Nup85</i> #3	C5	Same genotype as above (#1 =B6)	41 Chr (>60%)
		<i>ΔE2-mCherry-Nup85</i> #2	G4	<u>Allele 1:</u> in frame mCherry integration in exon2 and deletion from bp +639 to +716. <u>Allele 2:</u> <i>Nup85</i> KO. Deletions of 42 (+635 -- +677; 12 bp also inserted) and 33 bp (+683 -- +716; 9 bp also inserted) in exon 2.	41 Chr (>60%)
<b>Nup43<sup>-/-</sup></b>	<u>Parental cell line:</u> HM1 <u>Cas9 &amp; gRNA plasmid (s):</u> GeneArt CRISPR pOFP-Cas9-Nup43 gRNA1 <u>HR template:</u> none	<i>Nup43<sup>-/-</sup></i> #1	E4	<u>Allele 1:</u> <i>Nup43</i> KO. Deletion of 40 bp from +22 to +62. <u>Allele 2:</u> <i>Nup43</i> KO. Deletion of 32 bp from +38 to +70.	40 Chr (>60%)
		<i>Nup43<sup>-/-</sup></i> #2	B5	<u>Allele 1:</u> <i>Nup43</i> KO. Insertion of 1 bp at +52. <u>Allele 2:</u> <i>Nup43</i> KO. Insertion of 2 bp at +52.	40 Chr (>60%)

**Table S3: Sequences of gRNAs and PCR primers used to generate the template for homology-directed repair (HDR)**

Sequences of gRNAs used in this study		
IDENTIFIER	SEQUENCE	SOURCE
Seh1-gRNA1 [Exon 1 Fw]	TTGTGGCGCGCAGCATCG	This paper
Seh1-gRNA2 [Exon 1 Rv]	TGGCCATCCGGCGCCCG	This paper
sgTigre	ACTGCCATAACACCTTAACCTT	This paper
Mios-gRNA2 [Exon 3 Fw]	CCTATATGAAATGTGTTGCG	This paper
Mios-gRNA4 [Exon 3 Fw]	AGAATTGAGCCTTATCATG	This paper
ΔE2-GFP-Nup85-gRNA1 [Exon 2 Rv]	TTCTTCTTGGAAATTACACCC	This paper
ΔE2-GFP-Nup85-gRNA2 [Exon 2 Fw]	CTTCCTTCAACCAAACAGGT	This paper
Nup43-gRNA1 [Exon 1 Fw]	AAATTAGCAAAACCCGCTGG	This paper

PCR primers used to generate the template for homology-directed repair (HDR)		
IDENTIFIER	SEQUENCE	SOURCE
AG100 [Fw, HDR left arm] <i>GFP-Seh1, GFP-mAID-Seh1</i>	CCCGCGCAGGGCGTGTCCCTCGGGGCGTGGCGCGCCGGGCC GACGCGGGCGGGCAGGCAGGGCCGACGTGCCGTACGTGCCT CCGCGGTGG	This paper
AG101 [Fw, HDR left arm] <i>GFP-Seh1</i>	TACGTGCCTCCGCGGTGGTCCGGGGCTGCGGGCCGCACCGC CGCCCTCTGGAGGCAGGCGGGCCGGCAGACGGCGCGGCCATGG TTTCTGCCAAAAATCAAGC	This paper
AG103 [Rv, HDR right arm] <i>GFP-Seh1, GFP-mAID-Seh1</i>	GCGCCGCCTCCCTCCCTCCGGCCGGCAGCGCGAGTGCCG CGCGCACCTTGACGCTCTGATCGCTGGAGCAGGTGGCCATCCG GCGCCGTGGAAGTCG	This paper
AV01 [Rv, HDR right arm] <i>GFP-Seh1</i>	CGGCGCCGTGGAAGTCGAAAGACACATCGTGGATGAGGTCC TTGTGGTCCGCCCGATAGACCGGGCAGCAAGGATCTGAGT CCGGACTTGTACAGC	This paper
AV04 [Fw, HDR left arm] <i>GFP-mAID-Seh1</i>	CGGCGCCGTGGAAGTCGAAAGACACATCGTGGATGAGGTCC TTGTGGTCCGCCCGATAGACCGGGCAGCAAGGATCTGAGT CCGGATTATA	This paper
AV05 [Rv, HDR right arm] <i>GFP-mAID-Seh1</i>	TACGTGCCTCCGCGGTGGTCCGGGGCTGCGGGCCGCACCGC CGCCCTCTGGAGGCAGGCGGGCCGGCAGACGGCGCGGCCATGG TGAGCAAGGGCGAGGAG	This paper
BR3466 [Fw2, HDR left arm] <i>ΔE2-GFP[mCherry]-Nup85</i>	TCATTGGATACGACCTAGGCTTTCGCCTGGTGTATGGTTCG CAGTTACACTTGAAGTTGAATGGACAGTTGAGGGCTAAC	This paper
BR3467 [Fw1, HDR left arm] <i>ΔE2-GFP[mCherry]-Nup85</i>	GAATGGACAGTTGAGGGCTAACAGCTCCTGTGCCCTTATG TCTTACTAGTGGATTCCAGGTATGGTGAAGCAAGGGCGAGGAG	This paper
BR3468 [Rv1, HDR right arm] <i>ΔE2-GFP[mCherry]-Nup85</i>	CAGTTTCTGCCCTCAGAGAGTCACCAATGTGCCAAGACAGAT GTGAAAAAGAACTGCCTACCCCTGTACAGCTCGTCCATGCCG	This paper
BR3469 [Rv, HDR right arm] <i>ΔE2-GFP[mCherry]-Nup85</i>	GCAACTTGTCTCAAAATAGCAAAAAACAAAGTTCAAAAATA AAAATATTGCCAATAACTCCAGTTCTGCCAGAGAGTC	This paper

**Table S4: Antibodies**

Primary antibodies	Dilution	Source	Identifier
Human autoantibody against-centromeres (CREST serum)	IF: 1/500	ImmunoVision, Springdale, AR	HCT-0100
Rabbit polyclonal anti-GAPDH	WB: 1/2000	Trevigen, Minneapolis, MN	2275-PC100
Mouse anti-gamma-tubulin	WB: 1/50000	Abcam, San Francisco, CA	Ab11316
Rabbit polyclonal antibody anti-Mios	WB: 1/1000	Cell signalling, Danvers, MA	#13557
Affinity-purified rabbit polyclonal anti-Nup43 antibody	WB: 1/1000	From Doye lab (Zuccolo et al., 2007)	# 571/08-111-C1
Rabbit polyclonal anti-mNup85 serum	WB: 1/2000 IF: 1:1000 IP: 1/10	From D. Forbes (Harel et al., 2003)	N/A
Rabbit monoclonal antibody anti-Nup98 C39A3	IF: 1/20	Cell signalling	#2598
Rabbit polyclonal anti-Nup107	WB: 1/1000 IP: 1/10	From Doye lab (Belgareh et al., 2001)	#520-82
Rat monoclonal anti-mouse Nup133 antibody #74; clone 9C2H8)	IF: 1/100	From Doye lab (Berto et al., 2018)	N/A
Rabbit polyclonal antibody anti-Seh1	WB: 1/1000 IP: 1/500	Abcam, San Francisco, CA	ab218531
Rabbit polyclonal antibody anti-Tpr	IF: 1/200	Abcam, San Francisco, CA	ab84516
Rabbit polyclonal antibody anti Ostir	WB: 1/1000	From M.T. Kanemaki (Natsume et al., 2016)	N/A
Rabbit polyclonal antibody anti- Wdr24	WB: 1/1000	Proteintech, Manchester, UK	20778-1-AP

Secondary antibodies	Dilution	Source	Identifier
Cy™3 AffiniPure Donkey anti-rabbit	IF: 1/1000	Jackson ImmunoRes, West Grove, PA	711-165-152
Cy™3 AffiniPure Donkey anti-rat	IF: 1/1000	Jackson ImmunoRes	712-165-153
Cy™3 AffiniPure Goat anti-rabbit	IF: 1/1000	Jackson ImmunoRes	111-165-144
AlexaFluor 568 Donkey anti-rabbit	IF: 1/1000	ThermoFisher Scientific	A10042
Alexa 647 Donkey anti-human	IF: 1/500	Jackson ImmunoRes	709-605-149
Cy™5 AffiniPure Donkey anti-rat	IF: 1/1000	Jackson ImmunoRes	712-175-153
Peroxidase AffiniPure Donkey anti-rabbit	WB: 1/5000	Jackson ImmunoRes.	711-035-152
Peroxidase AffiniPure Goat anti-mouse	WB: 1/5000	Jackson ImmunoRes.	115-035-068
Peroxidase AffiniPure Goat anti-ratCy™5 AffiniPure Donkey Anti-Rat	WB: 1/5000	Jackson ImmunoRes.	112-035-167
Rabbit TrueBlot ®: Anti-Rabbit IgG HRP	WB: 1/1000	Rockland, Limerick, PA	18-8816-33

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