#### 1 TITLE: Germline Missense Variants in CDC20 Result in Aberrant Mitotic Progression and

#### 2 Familial Cancer

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## 29 SUMMARY:

- 30 CDC20 is a co-activator of the anaphase promoting complex/cyclosome (APC/C) and is essential for
- 31 mitotic progression.  $APC/C^{CDC20}$  is inhibited by the spindle assembly checkpoint (SAC), which prevents
- 32 premature separation of sister chromatids and aneuploidy in daughter cells. Although overexpression of
- 33 *CDC20* is common in many cancers, oncogenic mutations have never been identified in humans. Using
- 34 whole exome sequencing, we identified heterozygous missense *CDC20* variants (L151R and N331K)

1 that segregate with cancer in two families. Characterization of these mutants showed they retain APC/C

2 activation activity but show reduced binding to BUBR1, a component of the SAC. Expression of L151R

3 and N331K promoted mitotic slippage in HeLa cells and primary skin fibroblasts derived from carriers.

4 CRISPR/Cas9 was used to generate mice carrying N331K. Homozygous mice carrying N331K were non-

5 viable, however, heterozygotes displayed accelerated oncogenicity in Myc-driven cancers. These findings

6 highlight an unappreciated role for *CDC20* variants as tumor promoting genes in humans.

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8 KEYWORDS: Mitotic spindle, ovarian germ cell tumors, inherited cancer, anaphase promoting
9 complex/cyclosome (APC/C), CDC20

10

#### **11 INTRODUCTION:**

12 CDC20 (cell division cycle 20 homolog) is a co-activator of the anaphase promoting complex/cyclosome 13 (APC/C).<sup>1</sup> The APC/C is a large multi-subunit E3 ubiquitin ligase that is critical for cell cycle regulation.<sup>2</sup> 14 During mitosis, CDC20 binds and activates the APC/C, which ubiquitinates mitotic substrates cyclin B and securin in order for sister chromatids to separate and anaphase to proceed.<sup>3,4</sup> Mitotic progression is 15 regulated by the spindle assembly checkpoint (SAC), also known as the mitotic checkpoint.<sup>3,4</sup> The SAC is 16 17 triggered when cells in mitosis are not ready to proceed to anaphase, such as when kinetochores of 18 chromosomes are not properly attached to the spindle fibers during prometaphase.<sup>3,4</sup> Kinetochores that lack spindle attachment provide a negative signal to the APC/C<sup>CDC20</sup> complex through the mitotic 19 20 checkpoint complex (MCC), formed by the proteins MAD2, BUBR1 and BUB3, along with CDC20.<sup>5,6</sup> 21 The MCC binds APC/C<sup>CDC20</sup> directly and inhibits ubiquitination activity, resulting in a delay in anaphase 22 until all chromosomes are properly attached to the spindles.<sup>7,8</sup> The timing of sister chromatid separation 23 must be perfect in order to maintain proper ploidy.<sup>9</sup> All eukaryotic cells have this fail-safe mechanism to prevent aberrant chromosomal segregation.<sup>10,11</sup> Dysregulation of CDC20 and APC/C activity can lead to 24 problems in chromosome segregation, mitotic progression, and cell viability.<sup>12,13</sup> These aberrant events 25 have many pathological implications, including aneuploidy and tumorigenesis.<sup>12</sup> However, mutations in 26 27 genes coding for mitotic checkpoint proteins such as CDC20 are often embryonic lethal as they are essential to life.<sup>12,14,15</sup> Thus, such mutations are rare in human disease,<sup>12,16</sup> and the clinical significance of 28 29 mutations in CDC20 have been investigated in only a handful of studies in an animal model or human context.<sup>17–23</sup> Although CDC20 is overexpressed in the majority of human cancers, <sup>16,24–30</sup> the involvement 30 31 of CDC20 mutations in cancer has never been established in humans.

32

In the current study, we originally sought to identify the underlying genetic causes of familial malignant
 ovarian germ cell tumors (mOGCTs). We identified 8 families with histories of mOGCT.<sup>31</sup> We were able

1 to obtain biological material from 5 of these families and performed whole exome sequencing (WES) on

2 blood DNA to look for likely pathogenic or pathogenic variants that segregated in one or more families.

3 Using this approach, two distinct missense mutations in the *CDC20* gene were identified in all affected

4 individuals of 2 families. Since an uploidy and chromosome instability are implicated in mitotic

5 checkpoint defects and across different cancers,<sup>11</sup> we sought to investigate the role of CDC20 mutations

6 in cancer patients on cell cycle dysregulation and tumorigenesis. In parallel, we performed WES on germ

7 cell tumors (GCTs) of 85 other individuals with sporadic cases to look for other CDC20 mutations. Here,

8 we report our molecular and *in vivo* findings from our study of two germline *CDC20* mutations in patients

- 9 from two mOGCT families.
- 10
- 11

## 12 **RESULTS:**

#### 13 Identification of CDC20 variants in two families with histories of cancer

14 Saliva or blood samples were collected from five out of eight families previously reported<sup>31</sup> to show

15 mOGCT aggregation and WES analysis was performed on the germline DNA of available affected

16 members (Figures 1A and B, S1, S2, S3 and Table S1A). Two distinct *CDC20* missense variants were

identified in all affected members of two ('fmOGCT4' and 'fmOGCT7') of the five families studied, one

18 variant in each family (Table S1B and C). No other variants in this gene that have met our filtering

19 criteria were found in any of the other three analyzed families (**Table S1D, E and F**). Given that the

20 probability of finding 2 out of 5 families with very rare (allele frequency  $\leq 1 \times 10^{-5}$ ) segregating variants in

21 the same gene is low (p = 0.041) (**Table S2**), we selected the two *CDC20* variants for further analyses.

22 Family fmOGCT4<sup>32</sup> harbors the chr1:43825664T>G (hg19); c.452T>G (NM\_001255.2); p.L151R variant

23 (referred to as L151R onwards), which co-segregates with the disease in all affected women (Figure 1A).

24 Only one male carries the mutation and remains unaffected, most likely due to incomplete penetrance or

25 sex-related factors. Affected women in this family also presented with non-mOGCT cancers. In family

26 fmOGCT7,<sup>33</sup> two affected siblings inherited the chr1:43826548C>G (hg19); c.993C>G (NM\_001255.2);

27 p.N331K variant (referred to as N331K onwards) from their mother who presented with endometriotic

28 cysts and multinodular goiter. Of note, the family presented with a history of goiter, but whether CDC20

is related to that phenotype remains uncertain (Figure 1B). Both variants from both families were

30 heterozygous, expressed at the RNA level, and no loss of the wild-type (WT) allele or second hit in

31 *CDC20* was found in the tumors analyzed, thus suggesting a dominant mode of inheritance (Figure 1A

32 and B). In addition, both variants were predicted to be pathogenic by several *in silico* models (see Figure

33 1C legend).

1 In view of these results, we also screened 85 sporadic GCTs for *CDC20* mutations and found a predicted

2 deleterious mutation in one tumor belonging to a male patient (c.947G>A, p.R316H). Germline DNA was

3 not available in this case. However, there was no evidence that this variant has any functional

4 implications (data not shown).

5

6 WES was also performed on available tumoral material: fresh frozen tissue from fmOGCT7: III-1(mixed 7 GCT) and fmOGCT7: III-2 (Dysgerminoma, DG). As putative driver mutations, we found a variant in 8 AKT2 in the mixed GCT and a variant in KIT in the DG (Table S3). Genome-wide tumoral interrogation 9 was done using the OncoScan microarray assay on FFPE (formalin-fixed paraffin-embedded) samples 10 from these two tumors. Both showed several gains and/or losses mostly affecting whole chromosomes or 11 chromosome arms, resulting in considerable aneuploidy overall and consistent with CDC20 disruption 12 (Figure 2A). The copy number pattern of fmOGCT7 showed shared gains involving chr7, 8, 12 and 20 13 and a partial deletion region on p-arm of chr3 in both sisters (Figure 2B). An euploidy was also detected 14 with the FISH analysis performed on available tumoral slides, which showed a significant gain of 15 chromosomes in the two mOGCTs analyzed (Table S4). Furthermore, blood and skin cell samples were 16 taken from the two sisters (III-1 and III-2) in fmOGCT7 and the karyotypes were analyzed in 50 cells for 17 each. Chromosomal analyses revealed single an euploid cells as well as subclones with a t(3;13) and trisomy X in fibroblast and lymphocyte cultures, respectively (data not shown), from III-1 (Figure 1B). 18 19

20 Immunohistochemistry (IHC) was performed to visualize CDC20 on mOGCTs derived from III-1 and III-21 2 from fmOGCT7, the metastatic melanoma in I-2 from fmOGCT4, and a tissue microarray (TMA) of a 22 series of sporadic mOGCTs. Many of the mitotic figures in the CDC20-mututed tumors, particularly the 23 mOGCTs, were morphologically abnormal with asymmetric divisions and occasional partial or complete 24 ring forms, which is consistent with dysregulated CDC20 function (Figure 2C). Some tumor cells also 25 demonstrated distinctive vesicular nuclei with prominent eosinophilic nucleoli and peri-nucleolar 26 chromatin clearing. More than 50% of the cells in the familial tumors demonstrated nuclear and 27 cytoplasmic CDC20 immunoreactivity and the mOGCTs showed an unusual granular/reticular nuclear 28 staining pattern compared to the more diffuse immunoreactivity seen in most other sporadic tumors 29 (Figure 2C and Table S5). However, there were no significant differences in overall staining extent or 30 intensity between the familial cases and the sporadic mOGCT cases evaluated in the TMA (Table S5). 31 Taken together, these findings suggest that CDC20 mutations may play a role in the tumorigenesis in the 32 aforementioned familial cancers by disrupting the normal function and regulation of CDC20 in mitosis. 33

34

#### 1 CDC20 variants compromise the spindle assembly checkpoint

- 2 Although the L151 residue is close to the MAD2-binding domain (Figure 1C), L151R could not be
- 3 modelled structurally due to its location in a disorganized part of the protein. In contrast, the N331 residue
- 4 is in the highly conserved KEN-box binding site and forms a direct hydrogen bond with the K26 residue
- 5 of the KEN1 motif of BUBR1.<sup>5</sup> Computational structural analysis showed that mutating the N331 residue
- 6 to a lysine (K) removes this hydrogen bond interaction, creating an electrostatic repulsion that could
- 7 potentially weaken the affinity between CDC20 and BUBR1 (Figure 1D).
- 8

9 In order to determine if the identified CDC20 variants affect the SAC, HeLa cells were transfected with

- 10 FLAG-tagged CDC20 constructs expressing WT CDC20 or other variants: L151R and N331K, present in
- 11 the familial cases, and R132A. R132A is a well characterized mutation with defective binding to MAD2
- 12 which serves as a positive control for SAC disruption.<sup>8</sup> Transfected cells were synchronized with
- 13 thymidine (Thy), released, and arrested in mitosis with nocodazole (NOC). The mitotic status of cells was
- 14 monitored by immunoblot detecting the phosphorylated forms of BUBR1 and APC3 as mitotic markers.
- 15 Transfected cells expressing each of the mutant CDC20 constructs displayed increased levels of the
- 16 unphosphorylated forms of BUBR1 and APC3 compared to WT (Figure 3A, compare lane 4 to lanes 5,
- 17 6, and 7). In addition, flow cytometry performed on the Thy-NOC synchronized transfected cells, showed
- 18 that cells expressing the CDC20 mutants displayed a greater proportion of cells with tetraploidy,
- 19 suggesting that the mutant CDC20 expressing cells were more refractive to engagement of the SAC by
- 20 NOC (Figure 3B and C). In order to better characterize perturbations in the cell cycle, live cell, time-
- 21 lapse microscopy, was used to image HeLa cells transfected with bicistronic vectors expressing either WT
- or mutant CDC20 (see Methods) for 40 h after Thy-NOC synchronization. Figure 3D and E shows that
- cells transfected with mutant CDC20 underwent significantly more mitotic slippage and less mitotic cell
- 24 death compared to cells transfected with WT. The elevated levels of slippage observed in cells expressing
- the L151R and N331K mutations suggested that the SAC was attenuated. To test this possibility,
- 26 immunoprecipitations (IPs) were performed on FLAG-tagged CDC20 from transfected HeLa mitotic cell
- 27 extracts followed by immunoblot to detect components of the SAC. We observed that both the N331K
- and L151R mutants had decreased affinity to BUBR1 during mitosis (Figure 3F, lanes 3 and 4; Figure
- 29 S4A, lanes 3 and 4). As expected, the R132A CDC20 mutant had a reduced interaction with MAD2
- 30 (Figure 3F, lane 5). These data suggest that the L151R and N331K mutations disrupt normal mitotic
- 31 progression by compromising interaction with the SAC.
- 32
- 33

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# Familial CDC20 variants retain APC/C co-activator activity, which is required to promote mitotic slippage.

3

4 The L151R and N331K mutants were able to attenuate the SAC in HeLa cells and promote slippage, 5 suggesting that the mutant CDC20 retained activity as an APC/C co-activator. To test this possibility, we 6 reconstituted Cdc20 conditional knockout (KO) mouse embryonic fibroblasts (MEFs) with the bicistronic 7 CDC20 constructs as mentioned above. The Cdc20 KO conditional MEFS are derived from mice in 8 which exon 2 of the Cdc20 gene is flanked by LoxP sequences and knock-out is induced by addition of 4-9 hydroxytamoxifen (4-OHT) to stimulate expression of Cre recombinase.<sup>15</sup> As shown in Figure 4A, approximately 90% of cells become rounded and arrested in mitosis following 48 h of 4-OHT induction. 10 11 MEFs were electroporated with WT or mutant bicistronic CDC20 constructs and analyzed by real-time 12 microscopy as shown schematically in Figure 4B. Figure 4C and D show that reintroduction of WT 13 CDC20 into CDC20 KO cells rescued mitotic arrest in approximately 50% of transfected cells. 14 Reconstitution with the CDC20 point mutants L151R, N331K and R132A were all able to rescue mitotic 15 arrest comparable to WT. In contrast, the mutant R499E, in which the IR tail of CDC20 is mutated, was 16 completely defective for rescue. The IR amino acid motif is present on the C-termini of both APC/C coactivators, CDC20 and CDH1, and is essential for binding and activation of the APC/C.<sup>34</sup> Immunoblotting 17 18 was performed to confirm that the mutants expressed at similar levels (Figure S4B). 19 20 To determine if APC/C binding activity is required for induction of mitotic slippage induced by the 21 familial CDC20 variants, the L151R, N331K or R132A variants were combined with the R499E point 22 mutation (Figure 5A). IPs were performed in mitotic cell extracts to confirm the effects of the mutations. 23 MAD2 binding was decreased in the mutants that include R132A (Figure 5B, lanes 3, 4, and 7), and APC/C interaction was reduced in mutants containing the R499E as previously described<sup>34</sup> (Figure 5B, 24 25 lanes 5-7). All double mutants containing the L151R and N331K mutations had decreased affinity to 26 BUBR1, as well as impaired interaction with BUB3 (Figure 5B, lanes 3-6). All double mutants expressed 27 equally well (Figure S4C). Figure 5C shows that whereas the individual L151R, N331K or R132A 28 mutants were able to significantly increase the level of mitotic slippage, combination of these mutations

- 29 with the R499E IR tail mutation completely reversed this effect. In these experiments, slippage was also
- 30 induced by the R499E mutant and notably, this effect was abrogated by combination with L151R, N331K
- 31 or R132A, confirming that interaction with the SAC is essential for slippage induced by the IR tail mutant
- 32 (Figure 5B). In addition to elevated slippage, time-lapse microscopy in HeLa cells also revealed that
- 33 R132A, L151R and N331K had significantly shorter mitotic transit times in cells that underwent slippage
- in the presence of NOC (Figure 5D). The short mitotic transit times of the mutants were restored to WT

- 1 levels when combined with the R499E mutation. Taken together, these data show that the CDC20
- 2 variants L151R and N331K compromise the mitotic checkpoint via impaired interaction with BUBR1 and
- 3 that interaction and activation of the APC/C via the IR tail is required for this effect (Figure 5E).
- 4

## 5 The CDC20 N331K variant exhibits mitotic defects in primary cells from patients and accelerates 6 tumor formation *in vivo*.

7

Although the functional assays performed above suggest that the L151R and N331K mutations exhibit a compromised SAC, we wished to confirm the potential oncogenicity of these mutants in primary patient cells and *in vivo*. We were able to obtain primary skin fibroblasts from the two siblings in fmOGCT7 who are heterozygous for the N331K mutation (**Figure 1B**) and analyzed them by time-lapse microscopy as shown schematically in **Figure 6A**. Interestingly, fibroblasts from both siblings slipped faster through mitotic arrest in the presence of NOC (average times: patient III-1, 3.5 h; patient III-2, 4.2 h) than two control fibroblast lines (average times: Control 1, 8.9 h; Control 2, 11.1 h) (**Figure 6B and C**).

16 In order to determine if the N331K mutation is able to drive tumor formation *in vivo*, a knock-in mouse 17 model was derived by introducing the N331K mutation into one allele in C57BL/6 mice using CRISPR-Cas9 gene editing (Figure 7A, Figure S5). Mice heterozygous for the N331K mutation ( $Cdc20^{N331K/+}$ ) did 18 not significantly develop any more spontaneous tumors than did WT mice  $(Cdc20^{+/+})$ , nor was there a 19 20 difference in the growth and survival rates between heterozygous and WT animals (Figure S6A and B). However, no live births were observed of homozygous ( $Cdc20^{N331K/N331K}$ ) animals indicating that this 21 22 genotype is non-viable (Figure 7B). Moreover, homozygous embryos died between days E4.5 and E10.5 23 after fertilization and were severely delayed in development compared to WT embryos by the blastocyst 24 stage (Figure S6C and D). In support of the observation that the N331K mutation is non-viable, 25 introduction of this mutation in haploid, S. cerevisiae, also resulted in lethality (Figure S7). Although the 26 N331K variant alone did not result in spontaneous tumorigenesis in heterozygous mice, heterozygous 27 MEFs displayed accelerated mitotic transit compared to WT MEFs under the presence of NOC (averages 28 of 112 min vs. 45 min for WT and mutant, respectively) (Figure 7C, D and E). Since the N331K 29 mutation conferred an aberrant mitotic phenotype to MEFs under a pharmacological stress (NOC), we 30 therefore determined if this mutation could enhance tumorigenicity of a system already in an oncogenic context. We crossed our  $Cdc20^{N331K/+}$  mice with the Eµ-myc cancer model<sup>35,36</sup> to investigate any 31 cooperation of  $Cdc20^{N331K}$  with the Eµ-myc onco-transgene, with which mice develop B-cell lymphomas 32 at 2-3 months of age. Eµ-myc<sup>+</sup> mice carrying one N331K allele ( $Cdc20^{N331K/+}/Eµ-myc^+$ ) developed 33

34 lymphomas faster and experienced accelerated mortality rates compared to Eµ-myc<sup>+</sup> mice that were WT

for Cdc20 ( $Cdc20^{+/+}/E\mu$ -myc<sup>+</sup>) (Figure 7F). Two separate founder mouse lines carrying the N331K allele 1

2 were derived and tested in the E $\mu$ -myc model, and no differences in phenotype were found in the

3 experiments mentioned above (data not shown). Our characterization of the N331K variant in patient skin

4 fibroblasts and in mice demonstrate the potential of the germline N331K Cdc20 mutation to enhance

5 oncogene driven tumors in vivo.

6

#### 7 **DISCUSSION:**

8 Here, we report the identification of two germline CDC20 missense variants (L151R and N331K) that co-

9 segregate with cancer occurrence in two families. To our knowledge, no germline pathogenic variants

10 have previously been identified in CDC20 in human cancer. We have shown that the two variants

11 compromise the mitotic checkpoint in various models, accelerating mitotic transit and causing slippage,

12 likely contributing to the tumorigenic process of the various cancers in the families.

13

14 The impaired interaction between the familial CDC20 mutants and BUBR1 attenuates the SAC, resulting 15 in more mitotic slippage of cells (Figure 8). The reduced affinity of the N331K mutant to BUBR1 is 16 evidenced by the mutation being in the WD40 domain of CDC20, a crucial site of interaction with mitotic substrates that BUBR1 blocks when the SAC is engaged.<sup>37,38</sup> Although the L151R mutation could not be 17 18 modelled computationally, we observed that it also had reduced binding to BUBR1, despite not being 19 located in the WD40 domain. This observation could be due to a drastic change from a medium-sized 20 hydrophobic residue (leucine) to one that is large and positively charged (arginine), which could have an 21 impact on protein folding and BUBR1 binding as well. Moreover, a variant similar to the L151R - an 22 L151F mutation – had been identified in a kidney cancer (http://cancer.sanger.ac.uk/cosmic). As had been 23 previously noted,<sup>39</sup> likely deleterious missense mutations in genes encoding several APC/C proteins have 24 been identified in large-scale sequencing projects and could be selected for, as they result in a "just right" 25 level of genomic instability. At least two CDC20 missense mutations (L151F and P477A, Table S5 in 26 ref<sup>39</sup>) could fall into this group and are classified as probably damaging by Polyphen-2. Both the L151R 27 and N331K mutants had normal binding to MAD2 during the SAC but their abilities to induce mitotic 28 slippage were comparable to that of the R132A mutant, in which CDC20 is mutated in the MAD2 binding 29 domain (KILR motif). This suggests that impaired CDC20 interactions with BUBR1 and MAD2 have 30 comparable phenotypic outcomes relating to mitotic progression and slippage. 31

The lack of live births of N331K homozygous mice ( $Cdc20^{N331K/N331K}$ ) as well as developmental delays in 32

33 homozygous embryos corroborate our observation of no homozygous persons in our mOGCT families.

The significance of CDC20 in cell and organism viability was likewise reported in the study by Li et al.<sup>17</sup>. 34

1 In their study, mice homozygous for the Cdc20 allele in which three amino acid residues critical for Mad2 2 interaction were mutated to alanine (Cdc20<sup>AAA</sup>: K129A-R132A-P137A) died at late gestation and 3 embryos of which also exhibited severe developmental delays. Heterozygotes were viable but developed spontaneous tumors at highly accelerated rates, indicating that the SAC-mediated inhibition of Cdc20 is 4 5 an important tumor-suppressing mechanism. As previously described, the strength of the SAC is crucial 6 in maintaining an orderly mitosis.<sup>12</sup> A weakened SAC leads to premature chromosomal separation and 7 mitotic slippage.<sup>12</sup> Conversely, a hyperactive SAC results in lagging chromosomes (leading to 8 monosomies and trisomies) or cell death.<sup>12</sup> SAC component molecules must also be maintained at specific 9 stoichiometric ratios for proper SAC function. Overexpression of Mad2 in mice leads to aneuploidy and tumorigenesis,<sup>40</sup> and while mice with reduced levels of Cdc20 do not develop cancer, they accumulate 10 significant amounts of aneuploidy.<sup>41</sup> At the extreme end of the spectrum, cells without a SAC undergo 11 death in mitosis.<sup>12</sup> The fact that our heterozygous  $Cdc20^{N331K}$  mice were viable yet heterozygote MEFs 12 displayed an attenuated SAC under NOC could explain why mutant  $Cdc20^{N331K}$  molecules alone cannot 13 14 sustain live homozygous animals – without a WT Cdc20 copy, the SAC is severely weakened, failing to 15 regulate mitosis and maintain normal cell physiology. This suggests that missense mutations in other putative sites of CDC20 are presumably deleterious in development or pathogenic due to the fine balance 16 17 of CDC20 protein function and appropriate stoichiometric levels necessary for a normal SAC. Indeed, the 18 handful of CDC20 likely pathogenic or pathogenic variants identified thus far in clinical association 19 studies are potentially linked to problems with germ cell maturation in human infertility.<sup>19-22</sup> The notion 20 that CDC20 plays a vital role in development could explain the rarity of CDC20 germline alterations in 21 cancer and why those restricted to tumors are likely to be passenger mutations.

22

23 Germline variants in other relevant genes encoding SAC proteins have been associated with an increased

24 cancer risk.<sup>16</sup> Persons biallelic for pathogenic variants in *BUB1B* (encoding the BUBR1 protein) are at

25 high risk of childhood embryonal tumors such as embryonal rhabdomyosarcoma,<sup>42</sup> and also

26 gastrointestinal cancers in adults<sup>43</sup> – both cancers were observed in fmOGCT4. As seen in our CDC20

27 variants, pathogenic variants in *BUB1B* lead to SAC impairment. Defects or disruption in the SAC are

responsible for an increased rate of aneuploidy during tumorigenesis, but whether this represents a cause

29 or effect of oncogenesis is controversial.<sup>11</sup> We found subclones with trisomy X and a t(3;13) in blood and

- 30 skin cells of patients from fmOGCT7, however, such changes can also be seen in individuals without
- 31 CDC20 mutations. The significance of these findings is uncertain, but clearly the N331K CDC20 variant,

32 unlike biallelic *BUB1B* pathogenic variants,<sup>42,43</sup> does not cause mosaic variegated aneuploidy. Recent

33 studies have shown that low levels of an uploidy and transient chromosome instability are sufficient to

34 accelerate cancer progression in *p53*-deficient and *Myc*-driven cancer contexts.<sup>44,45</sup> When we crossed our

 $Cdc20^{N331K}$  mice with the Eµ-myc cancer model, Eµ-myc mice carrying the  $Cdc20^{N331K}$  allele developed 1 2 lymphomas and reached clinical endpoint sooner than animals carrying the WT allele of Cdc20. The 3 ability of N331K to accelerate *Mvc*-driven lymphomagenesis strongly suggests a role of these *CDC20* 4 variants in cancer predisposition. Indeed, there are ongoing efforts to uncover other candidate driver genes in mOGCTs.<sup>46</sup> We found a D820G mutation in the *KIT* gene in the DG arising in III-2 of 5 6 fmOGCT7, which was previously identified in hematopoietic neoplasia and testicular germ cell tumors.<sup>47-</sup> 7 <sup>49</sup> A nearby mutation, D816H, was found to play a transforming role in some seminomas via the KIT signal transduction pathway.<sup>50</sup> Moreover, another study suggested that D820G contributes to drug 8 9 resistance.<sup>51</sup> Therefore, while our data argue that the reported *CDC20* variants are predisposing alleles, we 10 cannot conclude that they are full oncogenic drivers as they likely rely on other activating events to 11 initiate and enhance neoplastic transformation.

12

13 In summary, we describe cancer-promoting germline variants in *CDC20* in two families ascertained

14 because of the occurrence of familial mOGCTs. These variants dysregulate the mitotic checkpoint. The

15 resulting cancers may not be restricted to the ovaries as pathogenic variants in *CDC20* likely also increase

16 the risk of neoplasia at other sites, as illustrated by the families in this study. Further studies are required

17 to determine the range and potential effects of germline *CDC20* variants and to better understand the

18 mechanisms triggering tumorigenesis through aneuploidy.

19

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31

## **32 AUTHOR CONTRIBUTIONS:**

- 33 Conceptualization, O.J.C., E.C., W.D.F., and J.G.T.; Investigation, O.J.C., E.C., M.M.-K., J.N., B.R.,
- 34 S.F., Y.W., C.P., L.J., J.C-Z., L.W., A.B., S.B., C.J.R.S., Z.Z., and W.C.C.; Resources, I.G., S.S., D.S.,

1	M.T., W.D.F.	, and J.G.T.; Writing	– Original Draft,	O.J.C. and E.C.:	Writing –	Review &	Editing. O	J.C.,

- 2 R.S., C.J.R.S., W.D.F. and J.G.T.; Supervision, R.S., C.M.T.G., D.B., J.M., W.D.F., and J.G.T.; Funding
- **3** Acquisition, W.D.F. and J.G.T.
- 4

#### **5 DECLARATION OF INTERESTS:**

- 6 The authors declare no competing interests.
- 7

## 8 MAIN FIGURE LEGENDS:

## 9 Figure 1: Pedigrees of the *CDC20* mutation carriers and locations of the L151R and N331K

## 10 mutations.

11 (A and B) In panel A, fmOGCT4 carries the c.452T>G, p.L151R mutation (\*). In panel B, fmOGCT7

12 carries the c.993C>G, p.N331K mutation (^). In both families, the mutation correlates with the disease,

13 and is expressed and shows no loss of heterozygosity in the tumors analyzed. Probands are identified with

14 an arrow. Affected individuals are filled in grey. Mutated and WT *CDC20* alleles are shown as minus and

15 plus signs, respectively. Age at last follow-up or age at death appears in years below for each individual.

16 Age at diagnosis is shown after disease description below affected individuals. Chromatograms from

17 Sanger sequencing of different tissue types also appear for several individuals. OC: ovarian cancer; CRC:

18 colorectal cancer; GCT: germ cell tumor; R: right; L: left; BC: breast cancer, IDC: invasive ductal

19 carcinoma; DCIS: ductal carcinoma in situ; MNG: multinodular goiter; MGCT: mixed GCT; YST: yolk

20 sac tumor; CC: choriocarcinoma; Hx: family history, or otherwise confirmed by clinical records; gDNA:

21 genomic DNA; cDNA: complementary DNA obtained from RNA; tDNA: DNA from tumoral tissue; Sal:

22 saliva.

23 (C) Schematic representation of the domains and motifs of human CDC20: C-box and IR tail are

24 important for association with APC/C; KEN and CRY sequences are degron recognition sites essential for

25 CDC20 degradation; KILR sequence is part of the MAD2-interacting motif; seven WD40 repeats are

26 important for mediating interaction with other proteins such as BUBR1. Multiple sequence alignment

27 from different organisms of the CDC20 residues surrounding the two familial mutations were performed

using CLUSTAL O (1.2.4). The black background indicates positions with fully conserved residues, dark

- 29 grey indicates conservation of residues of strongly similar properties, and light grey indicates
- 30 conservation of residues of weakly similar properties. The L151R and N331K variants were predicted to
- 31 be pathogenic by several *in silico* models: Combined Annotation Dependent Depletion (CADD) scores
- for L151R and N331K are 26 and 25, respectively; Revels scores for L151R and N331K are 0.47 and
- 33 0.15, respectively. A CADD score  $\geq 20$  indicates a deleterious variant. Revel scores range from 0 to 1,

1 and more damaging variants have higher scores. H.s.: *Homo sapiens*; R.n.: *Rattus norvegicus*; X.l.:

- 2 Xenopus; D.m.: Drosophila melanogaster; S.c.: Saccharomyces cerevisiae.
- 3 (D) Computational structural modelling comparing the theoretical protein structures of WT CDC20 and
- 4 the N331K variant (green), and their interactions with the KEN1-box of BUBR1 (cyan). The interactions
- 5 between residues N331 (in CDC20) and K26 (in BUBR1) are highlighted in the insets. The hydrogen
- 6 bond formed between WT CDC20 and BUBR1 is illustrated by the dashed yellow line. The CDC20
- 7 N331K mutation is predicted to prevent this hydrogen bond interaction with BUBR1.
- 8

## 9 Figure 2: Tumoral tissues from mOGCT patients show chromosomal abnormalities.

10 (A) OncoScan analysis on tumors. Copy number changes (CNV) and allelic imbalance (AI) are shown for

- 11 the two affected sisters: fmOGCT7\_III-1 and fmOGCT7\_III-2. The y-axes of the CNV and AI plots
- 12 indicate log2 ratios and B-allele frequency (BAF) ranging from 0 to 1 (with expected BAF value of 0.5
- 13 for heterozygous loci), respectively. Both samples show CNV on several chromosomes, which are also
- 14 reflected in the AI profiles.
- 15 (B) Copy number profiles for both samples (affected siblings of fmOGCT7). Shared copy number gains
- 16 (blue) can be observed on chromosomes 7, 8, 12, 20, as well as a loss (red) on chromosome 3.
- 17 (C) H&E staining and CDC20 immunohistochemistry on patient tumor sections. (i & ii): Malignant
- 18 mixed germ cell tumor (fmOGCT7, III-1). (i) shows histologically mature elements (squamous lined cyst,
- 19 upper left) and yolk sac tumor (YST) (lower right). The inset shows cytologic detail of the YST.
- 20 Vesicular nuclei, prominent eosinophilic nucleoli, and peri-nucleolar clearing are shown with the arrow.
- 21 (ii) is CDC20 immunohistochemistry showing nuclear and cytoplasmic staining in the YST (left) and in
- 22 an adjacent benign gland (right). The inset shows granular nuclear immunoreactivity in the YST. (iii &
- 23 iv): Dysgerminoma (fmOGCT7, III-2). In (iii), numerous mitotic figures are present including abnormal
- 24 partial ring forms (left arrow) and asymmetric divisions (right arrow). The inset shows cytologic detail
- 25 including cells with prominent nucleoli and peri-nucleolar clearing (arrow). In (iv),
- 26 immunohistochemistry shows nuclear CDC20 expression in most tumor cells together with weaker
- 27 cytoplasmic staining. The inset demonstrates the granular nuclear staining pattern. (v & vi): Metastatic
- 28 melanoma (fmOGCT4, I-2). In (v), many tumor cells show large vesicular nuclei, prominent nucleoli, and
- 29 peri-nucleolar clearing (arrow). The inset shows an abnormal, asymmetric mitotic figure (arrow). (vi)
- 30 shows CDC20 immunohistochemistry of melanoma within subcutaneous fat. Most tumor cells show
- 31 moderate nuclear and cytoplasmic staining.
- 32

## 33 Figure 3: The L151R and N331K CDC20 variants from mOGCT patients compromise the SAC in

34 HeLa cells.

- 1 (A) Expression blots of extracts of cells transfected with FLAG-tagged WT CDC20 or mutant constructs
- 2 and synchronized with thymidine (Thy) and nocodazole (NOC). A visual representation of the
- 3 experimental procedure is included above. The phosphorylated forms of BUBR1 and APC3 are used as
- 4 mitotic markers. NTNS: non-transfected, non-synchronized; NTS: non-transfected, synchronized; EV:
- 5 empty vector.
- 6 (B) Flow cytometry analysis of transfected, synchronized cells (same treatment protocol as in B) stained
- 7 with propidium iodide for DNA content. The 8N peaks indicate non-mitotic, polyploid (>4N) cells.
- 8 (C) Quantification of the proportion of non-mitotic, tetraploid cells from the flow cytometry analysis in
- 9 B, representing 3 independent experiments. An ordinary one-way ANOVA with Dunnett's multiple
- 10 comparisons test was used (n = 3; \*: p < 0.05; \*\*: p < 0.01). Error bars: SD.
- 11 (D) Visual representation of the fates of individual cells transfected with bicistronic constructs expressing
- 12 the different CDC20 mutants and GFP observed using real-time microscopy for 40 h after the addition of
- 13 nocodazole to synchronized cells. A visual timeline of the experimental procedure of this mitotic slippage
- 14 assay is included above. Images were taken every 10 min, and each analyzed GFP-positive cell was noted
- 15 for the frame at which it entered mitosis, and when it underwent apoptosis during mitosis or slipped out of
- 16 mitosis. Cells that died or slipped were not analyzed further. The colour scheme of the cell fates is as
- 17 follows: grey, interphase; red, mitosis resulting in death; blue, mitosis resulting in mitotic slippage. The
- 18 frames were converted to and shown in minutes (x-axis). N = 100 cells per construct (y-axis). These
- 19 results are representative of 3 independent time-lapse experiments.
- 20 (E) Quantification of the proportion of cells that underwent mitotic slippage across the different CDC20
- 21 mutants from the time-lapse experiment in panel **D**. An ordinary one-way ANOVA with Dunnett's
- 22 multiple comparisons test was used (n = 3; \*\*\*: p < 0.001). Error bars: SEM.
- 23 (F) Immunoblots of different mitotic proteins from FLAG co-immunoprecipitations (co-IPs) on mitotic
- 24 extracts from cells transfected with FLAG-tagged CDC20 WT or mutant constructs. The visual
- 25 representation of the experimental procedure is included above. EV: empty vector.
- 26

#### 27 Figure 4: Familial *CDC20* variants retain APC/C co-activator activity.

- 28 (A) Representative images confirming that the majority of Cdc20 conditional knockout (KO) mouse
- 29 embryonic fibroblasts (MEFs) treated with 4-OHT (4-hydroxytamoxifen) undergo mitotic arrest by 48 h.
- 30 (B) Visual timeline of the experimental procedure of the mitotic rescue assay performed in *Cdc20* KO
- 31 MEFs. Bicistronic constructs expressing the different CDC20 mutants with GFP were electroporated into
- 32 the MEFs.
- 33 (C) Visual representations of the fates of individual cells electroporated with bicistronic constructs
- 34 expressing the different mutant CDC20 mutants with GFP observed using real-time microscopy for 40 h

1 after thymidine release in 4-OHT. Images were taken every 10 min, and each analyzed GFP-positive cell

- 2 was noted for the frame at which it entered mitosis, and when it underwent apoptosis during mitosis or
- 3 exited mitosis. Cells that underwent mitotic progression were not followed further within the 40 h. Cells
- 4 that stayed arrested in mitosis by the end of the 40 h were labelled as "mitotic arrest". The colour scheme
- 5 of the different cell fates is as follows: grey, interphase; orange, continued mitotic arrest; red, mitosis
- 6 resulting in death; blue, mitosis resulting in mitotic progression (either cell division or mitotic slippage).
- 7 The frames were converted to and shown in minutes (x-axis). N = 100 cells per construct (y-axis). These
- 8 results are representative of 3 independent time-lapse experiments.
- 9 (D) Comparison of the differences in percent rescue of GFP-positive cells from mitotic arrest for the
- 10 different CDC20 mutants in the real-time microscopy experiment in C. An ordinary one-way ANOVA
- 11 with Dunnett's multiple comparisons test was used (n = 3; \*\*\*: p < 0.001). Error bars: SD.
- 12

## 13 Figure 5: APC/C co-activator activity of familial *CDC20* variants is still required to promote

- 14 mitotic slippage.
- 15 (A) Schematic showing the different CDC20 protein domains and the locations of the mutations in the
- 16 different double CDC20 mutant expression vectors tested. The double mutants are classified whether they
- are MAD2 binding disruptors (with the R132A mutant), APC/C interaction disruptors (with the R499E
- 18 mutant), or both.
- 19 (B) Immunoblots of different mitotic proteins from FLAG co-immunoprecipitations (co-IPs) on mitotic

20 extracts from cells transfected with FLAG-tagged CDC20 WT or double mutant constructs. Transfected

cells were synchronized with thymidine for 22 h, released for 3 h, arrested with nocodazole for 7 h, and

- 22 harvested. EV: empty vector.
- 23 (C) Quantification of the proportion of cells that underwent mitotic slippage across the different CDC20

24 mutants from time-lapse experiments performed as in Figure 3D. Slippage figures for the single CDC20

- 25 mutants (L151R, N331K and N331K) were taken from **Figure 3D** and used here for comparison
- 26 purposes. An ordinary one-way ANOVA with Dunnett's multiple comparisons test was used (\*\*: p <
- **27** 0.01; \*\*\*: p < 0.001; \*\*\*\*: p < 0.0001). Error bars: SEM.
- 28 (D) Comparison of the mitotic transit lengths before slippage between the different CDC20 single (from
- **Figure 3**) and combination mutants. Only cells that underwent mitotic slippage were considered. Cells
- 30 from a representative experiment were analyzed. Mean slippage times: luciferase, 15 h; WT, 16.2 h;
- 31 L151R, 9.7 h; N331K, 8.3 h; R132A, 8.1 h; R499E, 13.3 h; L151R/R499E, 13 h; N331K/R499E, 16.5 h;
- 32 R132A/R499E, 12.7 h. Thick lines indicate the medians of each plot; thin lines indicate the lower and
- 33 upper quartiles. An ordinary one-way ANOVA with Dunnett's multiple comparisons test was used

1 (n <sub>lu</sub>	uciferase = 3	$30; n_{WT} =$	35; $n_{L151R} =$	83; $n_{N331K} =$	$= 86; n_{R132A}$	$= 89; n_{R499E} =$	75; n <sub>L151R/R499E</sub> =	$= 71; n_{N331K/R499E} =$
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- 2 67;  $n_{R132A/R499E} = 32$ ; \*\*\*\*: p < 0.0001).
- 3 (E) Summary of the interactions and mitotic activities of the CDC20 double mutants.
- 4

## 5 Figure 6: Functional analyses of the CDC20 N331K mutant in patient skin fibroblasts demonstrate

- 6 a SAC impairment phenotype.
- 7 (A) Visual timeline of the experimental procedure of the mitotic slippage assay on N331K patient skin8 fibroblasts.
- 9 (B) Visual representation of N331K mitotic skin fibroblasts observed using time-lapse microscopy for 72
- 10 h. Fibroblasts were obtained from the two sisters in fmOGCT7 (who are heterozygotes for the N331K
- 11 variant III-1 and III-2, see Fig. 1B) and WT non-carriers. Images were taken every 15 min following

12 nocodazole treatment. Each analyzed cell was noted for the frame at which it entered mitosis and when it

13 slipped into interphase again. Bar cut-offs into white spaces indicate cell death. The colour scheme is as

14 follows: grey, interphase; blue, mitosis. The frames were converted to and shown in minutes (x-axis). N =

- 15 100 cells per patient (y-axis).
- 16 (C) Comparison of mitotic length for patient and control fibroblasts. Only cells that completed mitosis
- 17 and entered interphase again were considered. NT = not treated with nocodazole. An ordinary one-way
- 18 ANOVA with Tukey's multiple comparisons test was used ( $n_{III-1}=70$ ;  $n_{III-2}=63$ ;  $n_{Control 1}=56$ ;  $n_{Control 2}=57$ ;
- 19  $n_{\text{III-1 NT}}=65$ ;  $n_{\text{Control 2 NT}}=72$ ; \*\*\*\*: p < 0.0001). The solid lines indicate the medians of each plot; the
- 20 dotted lines indicate the lower and upper quartiles.
- 21

#### 22 Figure 7: The *Cdc20* N331K mutant exhibit mitotic defects and accelerate tumor formation *in vivo*.

- 23 (A) Schematic of the incorporation of the N331K mutation in the *Cdc20* locus in *Mus musculus* using the
- 24 CRISPR-Cas9 gene editing technology. An HpaI restriction site was included for genotyping purposes.
- **25** (B) Summary of the genotypes of newborn pups obtained from intercrosses of  $Cdc20^{N331K/+}$  mice.
- 26 (C) Visual timeline of the experimental procedure of the time-lapse microscopy performed on MEFs
- 27 extracted from the N331K mouse.
- 28 (D) Scatter violin plots of the duration of mitosis for +/+ (WT) and N331K/+ (het) MEFs from the
- 29 different isolated mouse embryos. The numbers indicate the MEF IDs. Images were taken every 15 min
- 30 following nocodazole treatment. Each analyzed cell was noted for the frame at which it entered mitosis
- 31 and when it entered interphase again. Only cells that completed mitosis and entered interphase again were
- 32 considered  $(n_1 = 37; n_2 = 26; n_3 = 36; n_6 = 41; n_7 = 22; n_4 = 33; n_5 = 18; n_8 = 36)$ . The thick lines indicate
- the medians of each plot; the thin lines indicate the lower and upper quartiles.

- 1 (E) Quantification of the average duration of mitosis before slippage between all N331K/+ and WT MEFs
- 2 analyzed in panel C. The individual means in panel C were taken and used to compute the average times
- 3 here. An unpaired, two-tail t test was used (+/+: n = 3, avg. time = 112 min; N331K/+: n = 5, avg. time = 112 min; N33
- 4 45 min; \*\*\*\*: p < 0.0001). Error bars: SEM.
- 5 (F) Survival (Kaplan-Meier) plot of animals from a cross between N331K heterozygote mice and mice
- 6 carrying the Eµ-myc onco-transgene. All mice that were observed in moribund states (due to signs of
- 7 cancer/lymphoma swollen lymph nodes, hunched posture, roughed fur, hyperventilation, and reduced
- 8 activity) were euthanized and were considered at endpoint for the calculation of the curves. A log-rank
- 9 (Mantel-Cox) test was performed to compare the survival curves.  $Cdc20^{N331K/+}/E\mu$ -myc<sup>-</sup>: n=22;

10  $Cdc20^{+/+}/E\mu$ -myc<sup>+</sup>: n = 36, median survival = 137 d;  $Cdc20^{N331K/+}/E\mu$ -myc<sup>+</sup>: n = 32, median survival = 83 11 d; p < 0.0001.

12

# Figure 8: A proposed mechanism by which CDC20 mutants L151R and N331K attenuate the SAC, resulting in aberrant mitotic progression.

- 15 Left panel: During the SAC in cells with WT CDC20, proteins MAD2, BUBR1, BUB3 and CDC20<sup>MCC</sup>
- 16 form the mitotic checkpoint complex (MCC) to inhibit the APC/C<sup>CDC20</sup> complex. The CDC20 molecules
- 17 that are part of the APC/C or MCC are indicated by their superscripts.
- 18 **Right panel:** In cells with mutant (L151R or N331K) CDC20, mutant CDC20<sup>MCC</sup> molecules have an
- 19 impaired interaction with BUBR1 (shown by red, dotted double arrows). Thus, defective MCCs
- 20 outcompete the fully functional MCCs (as indicated by >>>), and fewer APC/C<sup>CDC20</sup> complexes are
- 21 inhibited during the SAC. Cells experience a disrupted SAC and undergo mitotic slippage.
- 22
- 23

#### 24 METHODS:

#### 25 Patients

- 26 Germline DNA from blood or saliva was obtained from all available affected members from 5 mOGCT
- 27 families and sequenced by WES. Additional germline DNA from non-affected family members in
- 28 fmOGCT4 and fmOGCT7 was obtained to validate CDC20 mutations and assess co-segregation with the
- 29 disease. Additional biological samples were collected when possible. Tumoral formalin-fixed paraffin-
- 30 embedded (FFPE) and fresh frozen (FF) specimens from the mOGCTs and skin primary fibroblasts were
- also collected from the two sisters in fmOGCT7 (III-1 and III-2). In addition, 85 more FF DNA samples
- 32 were tested from an anonymized cohort of individuals (16 male and 69 female) with sporadic GCTs.
- 33 Informed consent was obtained from participants and all analyses were approved by the appropriate ethics
- 34 committee.

#### 1 Whole-exome sequencing (WES) analysis

2 Genomic DNA from blood and FFPE specimens were respectively subjected to the Agilent 50Mb exon

- 3 capture kit and the Nextera Rapid Exome kit, following the manufacturers' protocols. Each sample was
- 4 sequenced on Illumina HiSeq 2000 (100bp paired-end reads) at the McGill University and Génome
- 5 Québec Innovation Centre (MUGIC). Reads were first trimmed based on base quality (>30) and were
- 6 aligned to the human genome (UCSC hg19, NCBI build 37) using BWA (v. 0.5.9).<sup>52</sup> Then, the Genome
- 7 Analysis Toolkit (GATK)<sup>53</sup> performed local realignment of reads around small insertions and deletions
- 8 (indels) and Picard was used to remove duplicate reads from the final alignment for each sample. GATK
- 9 was used to ensure an average coverage of 80X of the consensus coding sequence (CCDS) in all samples.
- 10 Germline variants including single nucleotide variants (SNVs) and indels were called using SAMtools
- 11 (version 1.8)<sup>54</sup>. Somatic SNVs and indels were called using MuTect<sup>55</sup> and Indelocator
- 12 (https://confluence.broadinstitute.org/display/CGATools/Indelocator), respectively, and were then
- 13 annotated with ANNOVAR. To remove common variants and false positive calls, candidate variants were
- 14 subjected to several filtering steps and eliminated if they fulfilled any of the following criteria: (i)
- 15 genomic position of variant covered by <5X and less than 4 reads supporting the alternative variant, (ii)
- variant has allelic ratio <10% for SNVs or <15% for indels, (iv) variant has allele frequency >0.001 in our
- 17 non-cancer (>2000 exomes sequenced previously in our centre) or ExAC databases, or variant seen as
- 18 homozygote in ExAC database. Finally, only the most likely damaging variants (missense, nonsense,
- 19 splicing [within 6-bp of a splicing junction] and coding indels) that were from highly conserved regions
- 20 across species and predicted to be disease causing by at least 3 out of 5 bioinformatic software (SIFT,
- 21 PolyPhen, MutationTaster, Revel and MCAP) were considered for further analysis. The Integrative
- 22 Genomics Viewer<sup>56</sup> was used for the manual examination and visualization of all potential candidate
- 23 variants. We prioritized heterozygous variants from highly conserved regions across species shared
- among affected family members.
- 25

## 26 Simulation study

- 27 To estimate the probability of finding a pathogenic mutation/variant in the same gene in exome-
- 28 sequenced individuals with cancer in 2 out of 5 families, a simulation study was performed. We used the
- 29 pedigree structures of the families studied and varied two parameters: the number of genes of interest, G,
- 30 where the presence of a pathogenic variant is assessed, and the frequency of such variants in the general
- 31 population. The steps taken are explained in detail in **Tables S2A and S2B**.
- 32
- 33
- 34

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#### 1 Sanger sequencing

2 All the mutations found by WES were validated by Sanger sequencing using standard protocols. CDC20

3 screening on the 85 sporadic GCTs was also performed by Sanger on exons 3 to 6 and 8 to 10, which

4 include important functional motifs. All primers are available upon request.

5

## 6 Molecular modeling

7 Coordinates for human CDC20 in complex with the KEN Box of BUBR1 (PDB code 4GGD) were

- 8 visualized using PyMol.<sup>57</sup> In silico mutation and selection of viable rotamer for the CDC20 K331 residue
- 9 that prevents any steric clashes was also performed using PyMol.
- 10

#### 11 Plasmids

12 CDC20 cDNA was cloned into the multiple cloning site of the expression vector p3XFLAG-myc-CMV-

13 26 (Sigma). Site-directed mutagenesis was used to generate the *CDC20* mutants of interest: L151R was

14 generated through the c.452T>G nucleotide change; N331K through c.993C>A (although fmOGCT7

15 harbours a c.993C>G change, they both give rise to the same amino acid change and equivalent effects on

16 the mitotic checkpoint have been confirmed in both mutants – data not shown); R132A through

17 c.394\_395delinsGC; and R499E through c.1495\_1497delinsGAG. The p3XFLAG-myc-CMV-26

18 constructs were used in CDC20 expression and immunoprecipitation experiments. To follow transfected

19 cells through in the time-lapse microscopy experiments, the GFP-expressing bicistronic reporter construct

20 pIRES2-EGFP (BD Biosciences Clontech) was used. The pIRES2-EGFP vector contains an internal

21 ribosomal entry site (IRES), which allows the expression of both CDC20 and GFP proteins

22 independently. WT and CDC20 mutants were PCR-amplified from the p3XFLAG-myc-CMV-26 vector

and cloned into the multiple cloning site of pIRES2-EGFP. Double CDC20 mutants were created with

24 either additional site directed mutagenesis or sub-cloning from the previously made single-mutant

25 constructs. All double mutants were cloned into p3XFLAG-myc-CMV-26 and pIRES2-EGFP vectors and

tested as described above for the single mutants. All mutations were validated by Sanger sequencing.

27

#### 28 Cell lines, cell culture, transfection, and treatments of eukaryotic cells

29 HeLa cells and primary fibroblasts were regularly tested for mycoplasma by microscopy inspection

30 following DAPI staining. HeLa cells, primary fibroblasts, and mouse embryonic fibroblasts (MEFs) were

31 cultured using DMEM (Wisent Bioproducts: 4.5 g/L glucose, with L-glutamine, sodium pyruvate and

32 phenol red) with 10% fetal bovine serum (Sigma-Aldrich, Canadian origin) and gentamycin at 50 µg/mL

33 (Wisent Bioproducts). HeLa cells were seeded at 50-60% confluency and Lipofectamine® 2000

34 (ThermoFisher Scientific) was used for transfection following manufacturer's instructions, using a 1:2.5

- 1 or 1:2 ratio of DNA: Lipofectamine. Cells were treated with the appropriate drugs 24 h post-transfection.
- 2 To assess mitotic checkpoint function, transfected cells, fibroblasts and MEFs were synchronized with
- 3 0.61 mg/mL thymidine (Sigma) for 22-24 h, released for 3-4 h, and treated with 100 ng/mL nocodazole
- 4 (Sigma-Aldrich) overnight. For the co-immunoprecipitations, mitotic shake-off by dish tapping or strong
- 5 pipetting of transfected HeLa cells was performed after a 20 h treatment with 9 µM RO3306 (Alexis
- 6 Biomedicals) released for 75 min in nocodazole, or after thymidine-nocodazole synchronization as
- 7 described above. Cdc20 conditional knockout (KO) MEFs (Dr. Marcos Malumbres Lab) were
- 8 electroporated with the bicistronic *CDC20* constructs using the Gene Pulser Xcell<sup>™</sup> Electroporation
- 9 System (Bio-Rad) according to the manufacturer's instructions. MEFs were treated with 0.61 mg/mL
- 10 thymidine and 1 µM 4-hydroxytamoxifen (4-OHT, from Selleckchem) for 22-24 h to synchronize and
- 11 induce the KO of *Cdc20*. Synchronized cells were released into 4-OHT alone prior to live imaging.
- 12

## 13 Western Blotting

- 14 Transfected HeLa were harvested and lysed with lysis buffer (50 mM Tris-HCl pH 7.5, 150 mM NaCl,
- 15 0.5% NP-40, protease inhibitors tablet from Roche, 50 mM NaF, 10 mM glycerol 2-phosphate disodium
- 16 salt hydrate, 0.1 mM Na<sub>3</sub>VO<sub>4</sub> and 1 mM DTT). 40 µg of total protein were separated by SDS-PAGE and
- 17 probed with antibodies against BUBR1 (BD BioSciences), CDC27 (BD BioSciences), FLAG (Sigma),
- 18 CDC20 (Santa Cruz), β-actin (Sigma) MAD2 (Covance), and BUB3 (BD BioSciences).
- 19

## 20 Cell cycle analysis (Flow cytometry)

- 21 Transfected HeLa were harvested and fixed with ice-cold ethanol. Cellular DNA was stained using
- 22 propidium iodide and analyzed using the FACSCaliburs Analyser (Beckton-Dickinson). Data was
- 23 analyzed using FlowJo and ModFit softwares.
- 24

## 25 Time-lapse microscopy

26 Cells were synchronized with thymidine for 22-24 h, released for 3-4 h and imaged after the addition of 27 nocodazole. The Zeiss Axiovert 200M microscope was set to take pictures using the brightfield or FITC 28 filter at 10X magnification every 10 or 15 min for transfected HeLa or Cdc20 conditional KO MEFs with 29 bicistronic CDC20 constructs, mouse embryonic fibroblasts, or patient skin primary fibroblasts. For 30 analysis, 100 cells were followed frame by frame for each cell line. Only green cells were monitored for 31 pIRES2-EGFP-transfected cells. Attached cells were considered to be at interphase and rounded up cells 32 were counted as mitotic. Mitotic slippage times were taken from nuclear envelop breakdown to cell death, 33 slippage, or cytokinesis. Mitotic fates (death or slippage) were determined by cell morphology: cell death

was identified by signs of blebbing while slippage was characterized by cells flattening from mitosis
without appreciable cytokinesis.

3

#### 4 Immunoprecipitation

5 Mitotic extracts were quantified and normalized to the same concentration. Approximately 90% of the 6 extract was incubated with anti-FLAG M2 beads (Sigma, EZview RED FLAG M2 Affinity Gel) for 2 h, 7 or with anti-FLAG (Sigma) antibody overnight and then with equilibrated protein G beads (Millipore 8 Sigma, Protein G Agarose, Fast Flow) for 2h. After several washes with lysis buffer, samples were 9 resuspended in 4X loading buffer (200 mM Tris-HCl, 4% SDS, 40% glycerol, 4% β-mercaptoethanol, 10 0.012% bromophenol blue and 50 mM DTT) and resolved by SDS-PAGE. Mouse TrueBlot ULTRA 11 (anti-Mouse Ig (Rockland) secondary antibody) was used to probe for FLAG and CDC20 primary 12 antibodies.

13

### 14 OncoScan

- 15 Tumor samples from fmOGCT7 (III-1 mixed GCT and III-2 DG) were treated with the OncoScanTM
- 16 FFPE Assay Kit then subjected to the OncoScanTM array. The Nexus Express Software for OncoScan
- 17 v.3.1 (BioDiscovery, El Segundo, California) was used to call copy number (CNV) and allelic imbalance
- 18 (AI) across the genome. Briefly, the Fast Adaptive States Segmentation Technique (FASST2) algorithm
- 19 (a Hidden Markov Model based approach) segments copy number regions based on median log-ratio
- 20 derived from all probes in a given region. The segments were detected with a significance threshold of
- 21  $1.0^{-8}$ , a minimum of 10 probes per segment, and a maximum probe spacing of 1000 kbp between adjacent
- 22 probes. Then, the  $\log_2$  ratio was considered for the segments' classifications as follows: gains (> 0.08),
- 23 losses (< -0.1), high copy gains (> 0.7) and big losses (< -1.1). The AI state was determined for each
- 24 segment based on the median B-allele frequency (BAF) of heterozygous makers. Similar AI results were
- 25 obtained with WES data using ExomeAI (data not shown).<sup>58</sup>
- 26

## 27 Immunohistochemistry

- 28 Immunohistochemistry (IHC) was performed at the Segal Cancer Centre Research Pathology Facility
- 29 (Jewish General Hospital). Tissue samples were cut at 4 µm, placed on SuperFrost/Plus slides (Fisher)
- 30 and dried overnight at 37°C, before IHC processing. The slides were then loaded onto the Discovery XT
- 31 Autostainer (Ventana Medical System). All solutions used for automated IHC were from Ventana
- 32 Medical Systems unless otherwise specified. Slides underwent de-paraffinization, heat-induced epitope
- retrieval (CC1 prediluted solution Ref: 950-124, standard protocol). Immunostaining for p53 and CDC20
- 34 was performed online using a heat protocol. Briefly, a pre-diluted mouse monoclonal anti-p53 (Clone

- 1 Bp53-11, Ventana Medical Systems, Inc.) and a rabbit polyclonal anti-CDC20 (Abcam #ab72202) at
- 2 1:300 were manually applied for 32 min at 37°C, followed by the appropriate detection kit (OmniMap
- 3 anti-Mouse-HRP, Ref: 760-4310 and OmniMap anti-Rabbit-HRP, Ref: 760-4311) for 8 min, followed by
- 4 ChromoMap-DAB (Ref: 760-159). The omission of the primary antibody was included as a negative
- 5 control. Slides were counterstained with hematoxylin for 4 min, blued with Bluing Reagent for 4 min,
- 6 removed from the autostainer, washed in warm soapy water, dehydrated through graded alcohols, cleared
- 7 in xylene, and mounted with Eukitt Mounting Medium (EMS, Ref: 15320). Slides were analyzed by
- 8 conventional light microscopy (or scanned).
- 9

#### 10 Fluorescence *in situ* hybridization

11 Fluorescence in situ hybridization (FISH) analyses were performed on paraffin slides with tumor tissue from affected individuals from the mOGCT families as previously described.<sup>59</sup> For the detection of 12 13 chromosomal aneuploidies, commercially available chromosome enumeration probes for chromosomes 4, 14 10 and 20 (Abbott) were applied. In the familial cases, the locus specific LSI RUNX1/RUNX1T1 15 (8q21.3;21q22) Dual Color Dual Fusion probes (Abbott) mixed with a chromosome enumeration probe for 16 chromosome 18 (Abbott) were used. The FISH results on the tissue microarrays (TMAs) were evaluated 17 using Zeiss fluorescence microscopes equipped with the appropriate filter sets. For digital image acquisition 18 and processing, the ISIS digital image analysis system (MetaSystems, Altlussheim, Germany) was applied.

19

## 20 Generation of the *Cdc20*<sup>N331K</sup> mouse model

- 21 The GRCm38.p4 assembly from the NBCI database was used to map the *Cdc20* genomic sequence in
- 22 Mus musculus. A single-stranded guide RNA (sgRNA) sequence (ATGATAACATTGTCAACGTG) and
- 23 a single-stranded deoxy-oligonucleotide (ssODN) donor repair template
- 24 (GCCTTGATGTTGAGTGAATGTTTGCAGGGGGAGCCCATCCACTTTCTCCAGGACCACT
- 25 AGGCCACACGTTAACAATCTTATCATTGCCACCACTTGCCAGATGTCGTCCATCTGG
- 26 GGCCCAGCGAAGC) were used to respectively target the specific *Cdc20* locus and introduce the
- 27 c.993C>G, p.N331K mutation and a silent base substitution for an HpaI site. The sgRNA (Alt-R®
- 28 CRISPR-Cas9 crRNA and tracrRNA), recombinant Cas9 protein (Alt-R® S.p. Cas9 Nuclease V3) and
- the ssODN (Ultramer DNA Oligo) were all purchased from Integrated DNA Technologies.
- 30 Microinjections (50 ng/µl Cas9, 50 ng/µl sgRNA, 30 ng/µl ssODN and 100 mM KCl) were performed on
- 1 side of fertilized C57BL/6 embryos (Charles River) at 1.5 dpc. Injected embryos were subsequently
- 32 implanted in pseudo-pregnant females. All microinjections were performed by personnel at the McGill
- 33 Integrated Core for Animal Modeling (MICAM). Mosaic CRISPR-edited mice were screened with
- targeted deep sequencing (Illumina MiSeq PE250) using the following primer sequences:

#### 1 ACACTGACGACATGGTTCTACAAGTCCCTCGTTTATAGCTGAG and

#### 2 TACGGTAGCAGAGACTTGGTCTACCATGATGTTCGGGTAGC. MiSeq results were analyzed using

- 3 the Integrative Genomics Viewer (IGV) software.<sup>56</sup> Founders were obtained and backcrossed with WT
- 4 C57BL/6 mice. Genotyping was performed using restriction fragment length polymorphism analysis:
- 5 PCR was performed (with primers ACAATTAAACTCAACCCCGCC and
- 6 GAATGGACCAGTCCAACAAGG) and amplicons were digested with the HpaI restriction enzyme.
- 7 Genotypes were confirmed by Sanger sequencing.  $Cdc20^{N331K}$  mosaic founders were generated by
- 8 personnel in the McGill Integrated Core for Animal Modeling (MICAM). Two separate founder lines
- 9 were maintained throughout the study (see Figure S5 for more details). Animals were kept in specific-
- 10 pathogen-free (SPF) rooms. All animal studies were performed in accordance with ethical guidelines and
- 11 protocols approved by our research institute.
- 12

#### 13 Primary mouse embryonic fibroblasts

- 14 Pregnant females from  $Cdc20^{N331K/+}$  heterozygote intercrosses were euthanized 13.5 dpc. Embryos were
- 15 harvested, minced, digested in 0.25% trypsin/EDTA (Wisent Bioproducts) containing 10 mM HEPES
- 16 (Wisent Bioproducts) at 37°C, and filtered through a 100-µm cell strainer. MEFs were maintained at 37°C
- 17 in DMEM (Wisent Bioproducts: 4.5 g/L glucose, with L-glutamine, sodium pyruvate and phenol red)
- 18 with 10% fetal bovine serum (Sigma-Aldrich, Canadian origin). MEFs were subjected to the time-lapse
- 19 microscopy experiments as described above.
- 20

## 21 Survival and tumor studies in mice

- 22  $Cdc20^{N331K/+}$  and  $Cdc20^{+/+}$  mice were monitored once a week for any signs of disease and cancer. Mice
- 23 were euthanized at clinical endpoint (i.e., presence of tumors, inactivity, etc.). The weights of newborns
- 24 were taken weekly for the first 9 weeks after birth. To study the viability of  $Cdc20^{N331K/N331K}$  homozygote
- embryos, zygotes from heterozygote crosses were harvested from females at 0.5 dpc and cultured in
- 26 EmbyroMax® Advanced KSOM media (Sigma). For tumor formation studies, *Cdc20*<sup>N331K/+</sup> mice were
- 27 crossed with the Eµ-myc cancer model (Dr. Jerry Pelletier Lab). Mice were monitored weekly and
- 28 euthanized when in moribund states and/or at clinical endpoint (signs of cancer/lymphoma swollen
- 29 lymph nodes, hunched posture, inactivity).
- 30

## 31 Checkpoint cell survival assay in *S. cerevisiae*

32 The endogenous copy of *S. cerevisiae* CDC20 was replaced using KanMX in transformed yeast carrying a

- duplicate copy of CDC20 in a URA3 plasmid. Then, a CDC20 copy with the N407K mutation (equivalent
- to N331K in humans) or the well-known CDC20 KEN mutant (N405A/N407A/Q473A/R517L)<sup>5</sup> was

1 transformed using the LEU selection marker. The CDC20 WT URA3 plasmid was driven out by 5FOA 2 treatment. The cell survival assay was performed at several dilutions yeast using 50 µg/ml nocodazole for 3 up to 4 h. 4 5 **Statistical analysis** 6 Statistical analysis was performed with the GraphPad Prism (Version 9) software (GraphPad, San Diego, 7 CA). For comparisons of more than 2 groups in the functional assays, an ordinary one-way ANOVA was 8 performed followed by post-hoc pairwise comparisons (Dunnett's or Tukey's multiple comparisons test). 9 A p-value < 0.05 was considered statistically significant. For data presented in scatter-violin plots, the 10 median and upper and lower quartiles were considered. For histograms, data were expressed as mean  $\pm$ 11 standard error of the mean (SEM) or mean ± standard deviation (SD). All other statistical analyses are 12 described in the relevant table or figure legends in the Supplemental Information. 13 14 15 REFERENCES 16 Visintin, R., Prinz, S. & Amon, A. CDC20 and CDH1: A family of substrate-specific activators of 1. 17 APC- dependent proteolysis. Science (80-. ). 278, 460-463 (1997). 18 Manchado, E., Eguren, M. & Malumbres, M. The anaphase-promoting complex/cyclosome 2. 19 (APC/C): cell-cycle-dependent and -independent functions. Biochem. Soc. Trans. 38, 65-71 20 (2010). 21 3. Sivakumar, S. & Gorbsky, G. J. Spatiotemporal regulation of the anaphase-promoting complex in 22 mitosis. Nat. Rev. Mol. cell Biol. 16, 82-94 (2015). 23 4. Peters, J. M. The anaphase promoting complex/cyclosome: A machine designed to destroy. Nature 24 Reviews Molecular Cell Biology vol. 7 644–656 (2006). 25 5. Chao, W. C. H., Kulkarni, K., Zhang, Z., Kong, E. H. & Barford, D. Structure of the mitotic 26 checkpoint complex. Nature 484, 208-213 (2012). 27 6. Herzog, F. et al. Structure of the anaphase-promoting complex/cyclosome interacting with a 28 mitotic checkpoint complex. Science (80-.). 323, 1477-1481 (2009). 29 7. Nilsson, J., Yekezare, M., Minshull, J. & Pines, J. The APC/C maintains the spindle assembly 30 checkpoint by targeting Cdc20 for destruction. Nat. Cell Biol. 10, 1411–1420 (2008). 31 8. Izawa, D. & Pines, J. Mad2 and the APC/C compete for the same site on Cdc20 to ensure proper 32 chromosome segregation. J. Cell Biol. 199, 27-37 (2012). 33 9. Morgan, D. O. Regulation of the APC and the exit from mitosis. Nat. Cell Biol. 1, (1999). 34 10. Musacchio, A. & Salmon, E. D. The spindle-assembly checkpoint in space and time. Nat. Rev.

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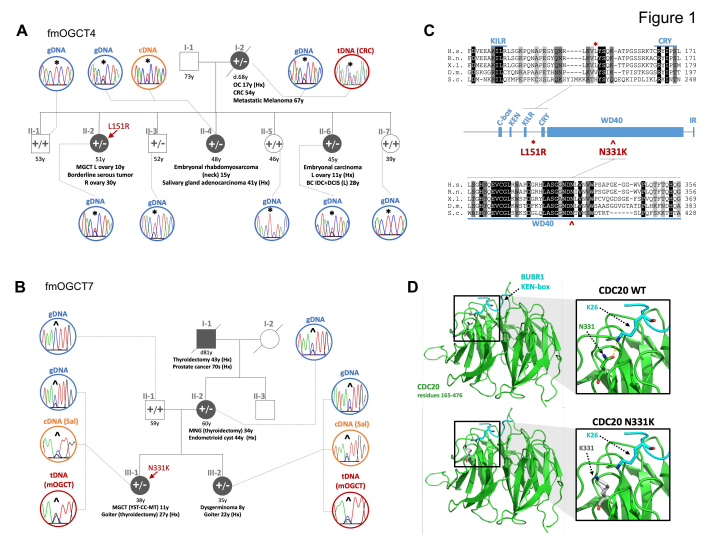
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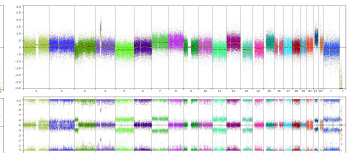
#### Figure 1: Pedigrees of the CDC20 mutation carriers and locations of the L151R and N331K mutations.

(A and B) In panel A, fmOGCT4 carries the c.452T>G, p.L151R mutation (\*). In panel B, fmOGCT7 carries the c.993C>G, p.N331K mutation (^). In both families, the mutation correlates with the disease, and is expressed and shows no loss of heterozygosity in the tumors analyzed. Probands are identified with an arrow. Affected individuals are filled in grey. Mutated and WT *CDC20* alleles are shown as minus and plus signs, respectively. Age at last follow-up or age at death appears in years below for each individual. Age at diagnosis is shown after disease description below affected individuals. Chromatograms from Sanger sequencing of different tissue types also appear for several individuals. OC: ovarian cancer; CRC: colorectal cancer; GCT: germ cell tumor; R: right; L: left; BC: breast cancer, IDC: invasive ductal carcinoma; DCIS: ductal carcinoma *in situ*; MNG: multinodular goiter; MGCT: mixed GCT; YST: yolk sac tumor; CC: choriocarcinoma; Hx: family history, or otherwise confirmed by clinical records; gDNA: genomic DNA; cDNA: complementary DNA obtained from RNA; tDNA: DNA from tumoral tissue; Sal: saliva.

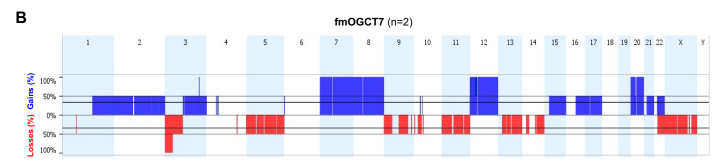
(C) Schematic representation of the domains and motifs of human CDC20: C-box and IR tail are important for association with APC/C; KEN and CRY sequences are degron recognition sites essential for CDC20 degradation; KILR sequence is part of the MAD2-interacting motif; seven WD40 repeats are important for mediating interaction with other proteins such as BUBR1. Multiple sequence alignment from different organisms of the CDC20 residues surrounding the two familial mutations were performed using CLUSTAL O (1.2.4). The black background indicates positions with fully conserved residues, dark grey indicates conservation of residues of strongly similar properties, and light grey indicates conservation of residues of weakly similar properties. The L151R and N331K variants were predicted to be pathogenic by several *in silico* models: Combined Annotation Dependent Depletion (CADD) scores for L151R and N331K are 26 and 25, respectively; Revels scores for L151R and N331K are 0.47 and 0.15, respectively. A CADD score ≥ 20 indicates a deleterious variant. Revel scores range from 0 to 1, and more damaging variants have higher scores. H.s.: *Homo sapiens*; R.n.: *Rattus norvegicus*; X.I.: *Xenopus*; D.m.: *Drosophila melanogaster*, S.c.: *Saccharomyces cerevisiae*.

(D) Computational structural modelling comparing the theoretical protein structures of WT CDC20 and the N331K variant (green), and their interactions with the KEN1-box of BUBR1 (cyan). The interactions between residues N331 (in CDC20) and K26 (in BUBR1) are highlighted in the insets. The hydrogen bond formed between WT CDC20 and BUBR1 is illustrated by the dashed yellow line. The CDC20 N331K mutation is predicted to prevent this hydrogen bond interaction with BUBR1.

fmOGCT7, III-1 (Mixed GCT)



fmOGCT7, III-2 (Dysgerminoma)



MGCT (fmOGCT7, III-1)

Α

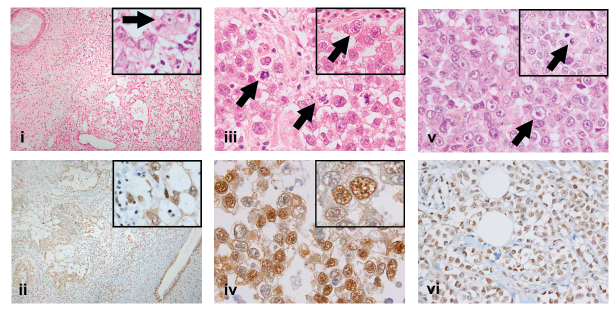
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#### Dysgerminoma (fmOGCT7, III-2)

#### Melanoma (fmOGCT4, I-2)

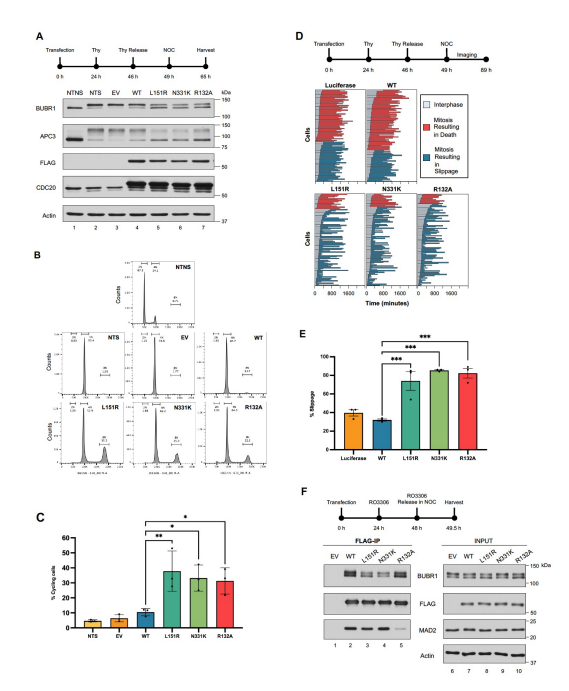


#### Figure 2: Tumoral tissues from mOGCT patients show chromosomal abnormalities.

(A) OncoScan analysis on tumors. Copy number changes (CNV) and allelic imbalance (AI) are shown for the two affected sisters: fmOGCT7\_III-1 and fmOGCT7\_III-2. The y-axes of the CNV and AI plots indicate log2 ratios and B-allele frequency (BAF) ranging from 0 to 1 (with expected BAF value of 0.5 for heterozygous loci), respectively. Both samples show CNV on several chromosomes, which are also reflected in the AI profiles.
 (B) Copy number profiles for both samples (affected siblings of fmOGCT7). Shared copy number gains (blue) can be observed on chromosomes 7,8,12, 20, as well as a loss (red) on chromosome 3.

(C) H&E staining and CDC20 immunohistochemistry on patient tumor sections. (i & ii): Malignant mixed germ cell tumor (fmOGCT7, III-1). (i) shows histologically mature elements (squamous lined cyst, upper left) and yolk sac tumor (YST) (lower right). The inset shows cytologic detail of the YST. Vesicular nuclei, prominent eosinophilic nucleoli, and peri-nucleolar clearing are shown with the arrow. (ii) is CDC20 immunohistochemistry showing nuclear and cytoplasmic staining in the YST (left) and in an adjacent benign gland (right). The inset shows granular nuclear immunoreactivity in the YST. (iii & iv): Dysgerminoma (fmOGCT7, III-2). In (iii), numerous mitotic figures are present including abnormal partial ring forms (left arrow) and asymmetric divisions (right arrow). The inset shows cytologic detail including cells with prominent nucleoli and peri-nucleolar clearing (arrow). In (iv), immunohistochemistry shows nuclear CDC20 expression in most tumor cells together with weaker cytoplasmic staining. The inset demonstrates the granular nuclear staining pattern. (v & vi): Metastatic melanoma (fmOGCT4, I-2). In (v), many tumor cells show large vesicular nuclei, prominent nucleoli, and peri-nucleolar clearing (arrow). The inset shows an abnormal, asymmetric mitotic figure (arrow). (vi) shows CDC20 immunohistochemistry of melanoma within subcutaneous fat. Most tumor cells show moderate nuclear and cytoplasmic staining.





#### Figure 3: The L151R and N331K CDC20 variants from mOGCT patients compromise the SAC in HeLa cells.

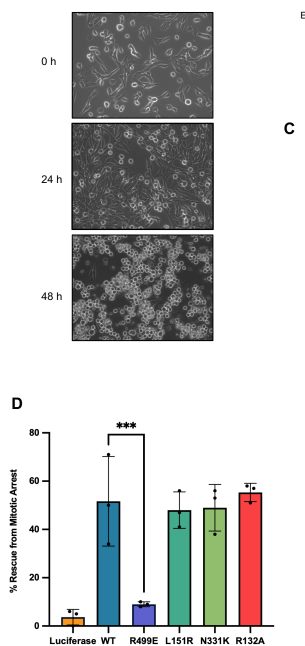
(A) Expression blots of extracts of cells transfected with FLAG-tagged WT CDC20 or mutant constructs and synchronized with thymidine (Thy) and nocodazole (NOC). A visual representation of the experimental procedure is included above. The phosphorylated forms of BUBR1 and APC3 are used as mitotic markers. NTNS: non-transfected, non-synchronized; NTS: non-transfected, synchronized; EV: empty vector.
 (B) Flow cytometry analysis of transfected, synchronized cells (same treatment protocol as in B) stained with propidium iodide for DNA content. The 8N peaks indicate non-mitotic, polyploid (>4N) cells.

(C) Quantification of the proportion of non-mitotic, tetraploid cells from the flow cytometry analysis in **B**, representing 3 independent experiments. An ordinary one-way ANOVA with Dunnett's multiple comparisons test was used (n = 3; \*: p < 0.05; \*\*: p < 0.01). Error bars: SD. (D) Visual representation of the fates of individual cells transfected with bicistronic constructs expressing the different CDC20 mutants and GFP observed using real-time microscopy for 40 h after the addition of NOC to synchronized cells. A visual timeline of the experimental procedure of this mitotic slippage assay is included above. Images were taken every 10 min, and each analyzed GFP-positive cell was noted for the frame at which it entered mitosis, and when it underwent apoptosis during mitosis or slipped out of mitosis. Cells that died or slipped were not analyzed

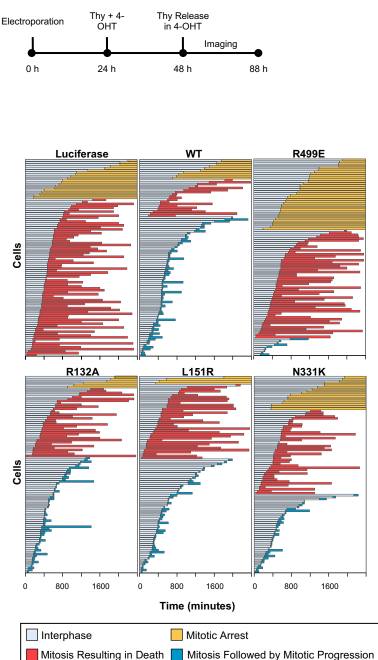
further. The colour scheme of the cell fates is as follows: grey, interphase; red, mitosis resulting in death; blue, mitosis resulting in mitotic slippage. The frames were converted to and shown in minutes (x-axis). N = 100 cells per construct (y-axis). These results are representative of 3 independent time-lapse experiments.

(E) Quantification of the proportion of cells that underwent mitotic slippage across the different CDC20 mutants from the time-lapse experiment in panel D. An ordinary one-way ANOVA with Dunnett's multiple comparisons test was used (n = 3; \*\*\*: p < 0.001). Error bars: SEM.</p>
(F) Immunoblots of different mitotic proteins from FLAG co-immunoprecipitations (co-IPs) on mitotic extracts from cells transfected with FLAG-tagged CDC20 WT or mutant constructs. The visual representation of the experimental procedure is included above. EV: empty vector.

Figure 4



4-OHT



#### Figure 4: Familial CDC20 variants retain APC/C co-activator activity.

(A) Representative images confirming that the majority of Cdc20 conditional knockout (KO) mouse embryonic fibroblasts (MEFs) treated with 4-OHT (4-hydroxytamoxifen) undergo mitotic arrest by 48 h.

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(B) Visual timeline of the experimental procedure of the mitotic rescue assay performed in Cdc20 KO MEFs. Bicistronic constructs expressing the different CDC20 mutants with GFP were electroporated into the MEFs.

(C) Visual representations of the fates of individual cells electroporated with bicistronic constructs expressing the different mutant CDC20 mutants with GFP observed using real-time microscopy for 40 h after thymidine release in 4-OHT. Images were taken every 10 min, and each analyzed GFP-positive cell was noted for the frame at which it entered mitosis, and when it underwent apoptosis during mitosis or exited mitosis. Cells that underwent mitotic progression were not followed further within the 40 h. Cells that stayed arrested in mitosis by the end of the 40 h were labelled as "mitotic arrest". The colour scheme of the different cell fates is as follows: grey, interphase; orange, continued mitotic arrest; red, mitosis resulting in death; blue, mitosis resulting in mitotic progression (either cell division or mitotic slippage). The frames were converted to and shown in minutes (x-axis). N = 100 cells per construct (y-axis). These results are representative of 3 independent time-lapse experiments. (D) Comparison of the differences in percent rescue of GFP-positive cells from mitotic arrest for the different CDC20 mutants in the real-time microscopy experiment in **C**. An ordinary one-way ANOVA with Dunnett's multiple comparisons test was used (n = 3; \*\*\*: p < 0.001). Error bars:

SD.

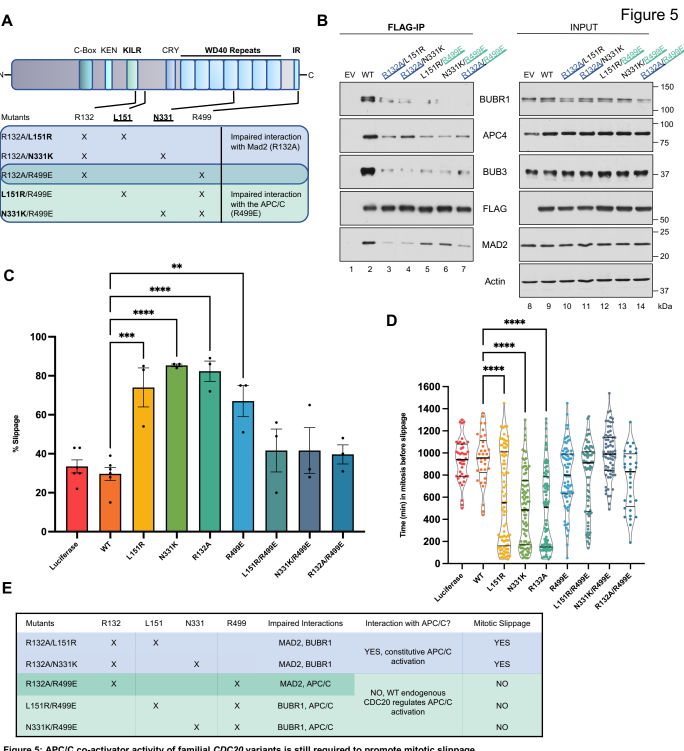


Figure 5: APC/C co-activator activity of familial CDC20 variants is still required to promote mitotic slippage.

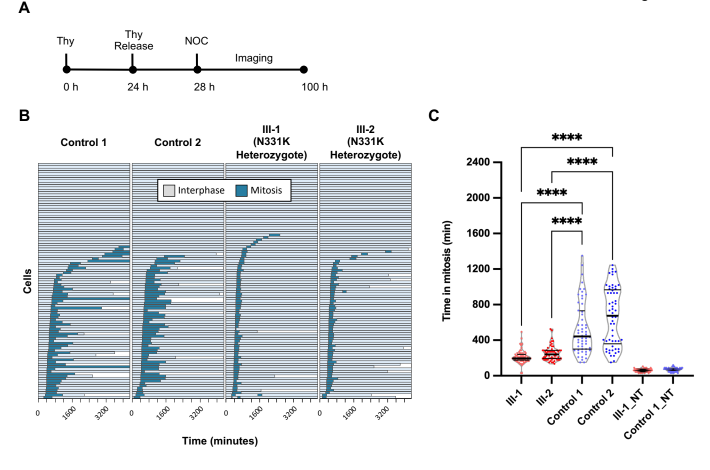
(A) Schematic showing the different CDC20 protein domains and the locations of the mutations in the different double CDC20 mutant expression vectors tested. The double mutants are classified whether they are MAD2 binding disruptors (with the R132A mutant), APC/C interaction disruptors (with the R499E mutant), or both.

(B) Immunoblots of different mitotic proteins from FLAG co-immunoprecipitations (co-IPs) on mitotic extracts from cells transfected with FLAGtagged CDC20 WT or double mutant constructs. Transfected cells were synchronized with thymidine for 22 h, released for 3 h, arrested with nocodazole for 7 h, and harvested. EV: empty vector.

(C) Quantification of the proportion of cells that underwent mitotic slippage across the different CDC20 mutants from time-lapse experiments performed as in Figure 3D. Slippage figures for the single CDC20 mutants (L151R, N331K and N331K) were taken from Figure 3D and used here for comparison purposes. An ordinary one-way ANOVA with Dunnett's multiple comparisons test was used (\*\*: p < 0.01; \*\*\*: p < 0.001; \*\*\*\*: p < 0.0001). Error bars: SEM.

(D) Comparison of the mitotic transit lengths before slippage between the different CDC20 single (from Figure 3) and combination mutants. Only cells that underwent mitotic slippage were considered. Cells from a representative experiment were analyzed. Mean slippage times: luciferase, 15 h; WT, 16.2 h; L151R, 9.7 h; N331K, 8.3 h; R132A, 8.1 h; R499E, 13.3 h; L151R/R499E, 13 h; N331K/R499E, 16.5 h; R132A/R499E, 12.7 h. The thick lines indicate the medians of each plot; the thin lines indicate the lower and upper quartiles. An ordinary one-way ANOVA with Dunnett's multiple comparisons test was used (n<sub>luciferase</sub> = 30; n<sub>WT</sub> = 35; n<sub>L151R</sub> = 83; n<sub>N331K</sub> = 86; n<sub>R132A</sub> = 89; n<sub>R499E</sub> = 75; n<sub>L151RR499E</sub> = 71; n<sub>N331K/R499E</sub> = 67; n<sub>R132A/R499E</sub> = 32; \*\*\*\*: p < 0.0001).

(E) Summary of the interactions and mitotic activities of the CDC20 double mutants.

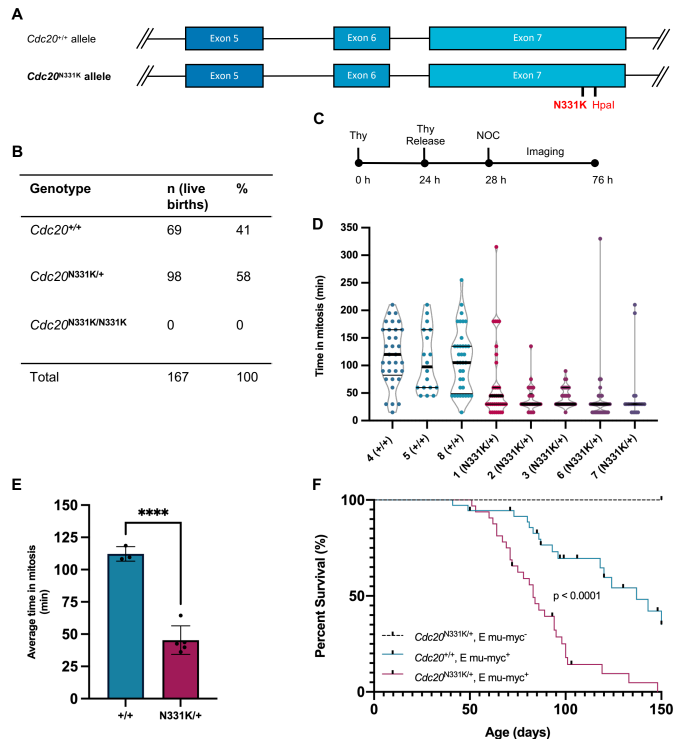


# Figure 6: Functional analyses of the CDC20 N331K mutant in patient skin fibroblasts demonstrate a SAC impairment phenotype.

(A) Visual timeline of the experimental procedure of the mitotic slippage assay on N331K patient skin fibroblasts.

(B) Visual representation of N331K mitotic skin fibroblasts observed using time-lapse microscopy for 72 h. Fibroblasts were obtained from the two sisters in fmOGCT7 (who are heterozygotes for the N331K variant – III-1 and III-2, see Fig. 1B) and WT non-carriers. Images were taken every 15 min following nocodazole treatment. Each analyzed cell was noted for the frame at which it entered mitosis and when it slipped into interphase again. Bar cut-offs into white spaces indicate cell death. The colour scheme is as follows: grey, interphase; blue, mitosis. The frames were converted to and shown in minutes (x-axis). N = 100 cells per patient (y-axis).

(C) Comparison of mitotic length for patient and control fibroblasts. Only cells that completed mitosis and entered interphase again were considered. NT = not treated with nocodazole. An ordinary one-way ANOVA with Tukey's multiple comparisons test was used ( $n_{III-1}=70$ ;  $n_{III-2}=63$ ;  $n_{Control 1}=56$ ;  $n_{Control 2}=57$ ;  $n_{III-1_NT}=65$ ;  $n_{Control 2_NT}=72$ ; \*\*\*\*: p < 0.0001). The solid lines indicate the medians of each plot; the dotted lines indicate the lower and upper quartiles.



#### Figure 7: The Cdc20 N331K mutant exhibit mitotic defects and accelerate tumor formation in vivo.

(A) Schematic of the incorporation of the N331K mutation in the Cdc20 locus in Mus musculus using the CRISPR-Cas9 gene editing technology. An Hpal restriction site was included for genotyping purposes.

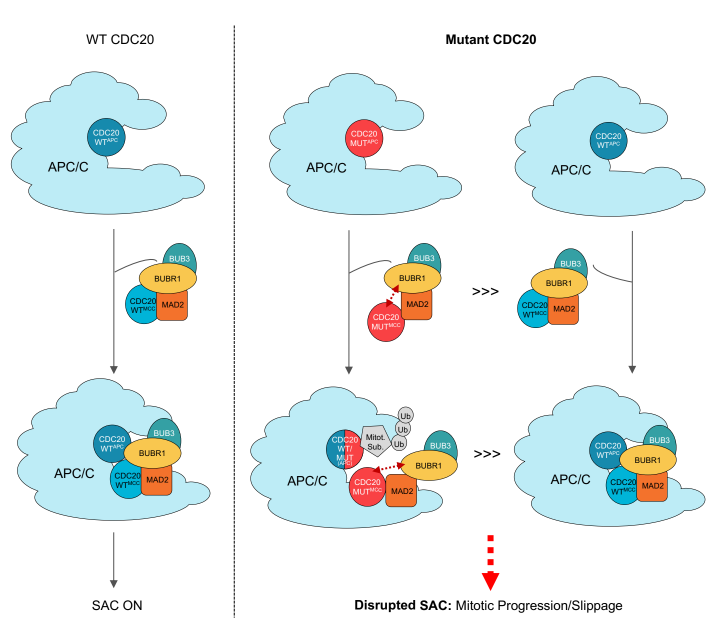
(B) Summary of the genotypes of newborn pups obtained from intercrosses of Cdc20N331K/+ mice.

(C) Visual timeline of the experimental procedure of the time-lapse microscopy performed on MEFs extracted from the N331K mouse. (D) Scatter violin plots of the duration of mitosis for +/+ (WT) and N331K/+ (het) MEFs from the different isolated mouse embryos. The numbers indicate the

MEF IDs. Images were taken every 15 min following nocodazole treatment. Each analyzed cell was noted for the frame at which it entered mitosis and when it entered interphase again. Only cells that completed mitosis and entered interphase again were considered (n1 = 37; n2 = 26; n3 = 36; n6 = 41; n7 = 22; n4 = 33; n<sub>5</sub> = 18; n<sub>8</sub> = 36). The thick lines indicate the medians of each plot; the thin lines indicate the lower and upper quartiles.

(E) Quantification of the average duration of mitosis before slippage between all N331K/+ and WT MEFs analyzed in panel C. The individual means in panel C were taken and used to compute the average times here. An unpaired, two-tail t test was used (+/+: n = 3, avg. time = 112 min; N331K/+: n = 5, avg. time = 45 min; \*\*\*\*: p < 0.0001). Error bars: SEM.

(F) Survival (Kaplan-Meier) plot of animals from a cross between N331K heterozygote mice and mice carrying the Eµ-myc onco-transgene. All mice that were observed in moribund states (due to signs of cancer/lymphoma - swollen lymph nodes, hunched posture, roughed fur, hyperventilation, and reduced activity) were euthanized and were considered at endpoint for the calculation of the curves. A log-rank (Mantel-Cox) test was performed to compare the survival curves. Cdc20<sup>N331K/+</sup>/Eµ-myc<sup>+</sup>: n = 22; Cdc20<sup>+/+</sup>/Eµ-myc<sup>+</sup>: n = 36, median survival = 137 d; Cdc20<sup>N331K/+</sup>/Eµ-myc<sup>+</sup>: n = 32, median survival = 83 d; p < 0.0001.



## Figure 8: A proposed mechanism by which CDC20 mutants L151R and N331K attenuate the SAC, resulting in aberrant mitotic progression.

Left panel: During the SAC in cells with WT CDC20, proteins MAD2, BUBR1, BUB3 and CDC20<sup>MCC</sup> form the mitotic checkpoint complex (MCC) to inhibit the APC/C<sup>CDC20</sup> complex. The CDC20 molecules that are part of the APC/C or MCC are indicated by their superscripts.

**Right panel:** In cells with mutant (L151R or N331K) CDC20, mutant CDC20<sup>MCC</sup> molecules have an impaired interaction with BUBR1 (shown by red, dotted double arrows). Thus, defective MCCs outcompete the fully functional MCCs (as indicated by >>>), and fewer APC/C<sup>CDC20</sup> complexes are inhibited during the SAC. Cells experience a disrupted SAC and undergo mitotic slippage.