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## Understanding phenotypic variation in rodent models with germline *Apc* mutations

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### Abstract

Adenomatous Polyposis Coli (APC) is best known for its crucial role in colorectal cancer suppression. Rodent models with various *Apc* mutations have enabled experimental validation of different *Apc* functions in tumors and normal tissues. Since the development of the first mouse model with a germline *Apc* mutation in the early 1990s, twenty other *Apc* mouse and rat models have been generated. This article compares and contrasts currently available *Apc* rodent models with particular emphasis on providing potential explanations for their reported variation in three areas: 1) intestinal polyp multiplicity, 2) intestinal polyp distribution, and 3) extra intestinal phenotypes.

### Introduction

Tumor suppressor Adenomatous polyposis coli (APC) is critical for maintaining cellular homeostasis in the intestine (1, 2). APC is a large (2843 amino acids), multi-domain protein that has been implicated in many cellular functions including cellular proliferation, differentiation, cytoskeleton regulation, migration and apoptosis (3). Mechanistically, APC is best known for its ability to antagonize Wnt signaling by targeting the oncoprotein  $\beta$ -catenin for proteasomal degradation (4).

Acquiring a somatic *APC* mutation is an early, if not initiating event in the great majority of colorectal tumors (5). Inheriting a germline *APC* mutation results in the development of hundreds to thousands of colonic polyps, a condition termed familial adenomatous polyposis (FAP). These precancerous polyps are thought to initiate following a somatic mutation in the wild-type *APC* allele (6, 7). To avoid the progression of these polyps into invasive carcinoma, prophylactic colon removal is recommended for FAP (8). There are no reports of humans with germline mutation of both *APC* alleles, consistent with early developmental lethality associated with complete loss of APC function (9–11). Germline and somatic *APC* mutations typically result in premature APC protein truncation and group between codons 1250 and 1464, a region termed the “mutation cluster region”, MCR (12).

A meta-analysis of genotype-phenotype correlation in FAP patients showed that germline mutations in the MCR result in the most severe intestinal polyposis phenotype, with up to 5000 polyps (13). Mutations on either side of the MCR are associated with an intermediate intestinal polyposis phenotype, while mutations that result in a truncation in APC after amino acid (a.a.) 1595 or before a.a. 157 are associated with an attenuated phenotype (AFAP), characterized by development of only a few polyps (13). Complete deletion of *APC* has been reported only rarely and results in an intermediate phenotype (14, 15).

Over two-thirds of FAP patients also have extra-colonic manifestations (13). Chronic hypertrophy of retinal pigment epithelium (CHRPE) is the most frequent phenotype, associated with APC truncation between a.a. 311–1446. Desmoid tumors, on the other hand, are associated with APC truncations 3' to the MCR, after a.a. 1400. Duodenal and gastric tumors have been associated with *APC* mutations in two different regions, downstream of codon 1395 and between codons 564–1465 (13). It is important to note that these genotype-phenotype correlations are not rigid or complete, suggesting roles for other genetic and environmental factors in tumor development (13, 16).

For the past two decades, rodent models have been valuable for analysis of *APC* functions in intestinal homeostasis and tumor suppression (17, 18). *APC* is well-conserved between human and rodent, with 92% similarity at the amino acid level (9, 19). Furthermore, some rodent models with germline *Apc* mutations that result in *Apc* protein truncation develop intestinal polyposis similar to that seen in FAP patients (18). A brief summary of all published rodent models with germline *Apc* mutations appears in Tables 1–3, with a schematic provided in figure 1.

Characterization of the many available *Apc* mouse and rat models has aided in discovery of various pathways important in colon carcinogenesis. *Apc* rodent models were also useful for elucidating the effect of various environmental and genetic factors on intestinal tumorigenesis, and for testing potential chemoprevention and therapeutic agents. The many positive contributions of *Apc* mouse models have been reviewed previously (20, 21). As with most experimental systems, studies of the *Apc* models have also led to unanswered questions, particularly regarding phenotypic variation among the different models. Here we review some of these variations, provide potential explanations, and pose challenges for future investigation.

## I- Variation in intestinal polyp multiplicity

As shown in table 1, the average number of polyps varies greatly between different mouse models with germline *Apc* mutations. In addition, the number of polyps also varies in the same *Apc* mouse model maintained in different laboratories (17). These variations in intestinal polyp number in different models likely stem from the nature of the *Apc* mutations as well as environmental and genetic factors (17, 18). We propose that the number of intestinal tumors that develop in different *Apc* models and in the same model analyzed by different laboratories is influenced by one or more of the following factors:

**1- Different rates and mechanisms of wildtype *Apc* allele loss (e.g. LOH, mutation of wildtype *Apc*, gene silencing)**—In both FAP patients and rodent models with germline *Apc* mutations, loss or inactivation of the wildtype *APC/Apc* allele is required for polyp formation (22, 23). The mechanism by which the second wildtype *Apc* allele is lost appears to depend on the *Apc* mouse model (24). Because this second *Apc* “hit” is essential for polyp initiation (10, 22, 25), the rate at which second “hit” occurs will directly affect the number of intestinal polyps. Increasing the expected rate of these second “hits” through introduction of genomic instability, X-ray exposure, or injection with a mutagen significantly increases the number of polyps in *Apc*<sup>Min/+</sup> and *Apc*<sup>1638N</sup> mice (26–30). It has been suggested that certain *Apc* mutations might lead to chromosomal instability, which could affect the rate of wildtype *Apc* loss (31).

*Apc*<sup>1638N/+</sup> mice develop relatively few intestinal polyps and the second *Apc* “hit” is usually inactivation of the wildtype *Apc* allele, predicted to be a rare event (24). On the other hand, *Apc*<sup>Min/+</sup> mice, where the wildtype *Apc* allele is lost by means of a more frequent LOH event, develop considerably more polyps (24). Loss of the wildtype *Apc* allele in both *Apc*<sup>Min/+</sup> and *Apc*<sup>1322T/+</sup> mice, however, is reported to occur via LOH, yet these two mouse

models have widely different polyp numbers (32). Although the rate and underlying mechanism of wildtype *Apc* allele loss might contribute to intestinal polyp numbers in *Apc* mouse models, it is unlikely that these are sole defining parameters.

**2- Different rates of polyp growth due to differences in Wnt signaling**—Polyps must reach a certain size to be detectable. If two polyps are initiated at the same time, a more rapidly growing polyp should be detectable earlier than a slower growing polyp. The most recognized function of *Apc* is to antagonize the Wnt signaling pathway through inhibition of  $\beta$ -catenin's activity as a transcription co-factor (4). As Wnt signaling can drive cellular proliferation, we might expect that different *Apc* mutations would lead to different levels of Wnt signal activation and different corresponding changes in cellular proliferation. In FAP patients, mutations in the MCR are associated with the most severe intestinal phenotypes while mutations outside the MCR lead to reduced polyp multiplicity (13). Notably, *APC* mutations 5' and 3' to the MCR result in higher and lower activation of Wnt signaling, respectively (33). This observation has led to the proposal that submaximal upregulation of Wnt signaling promotes more polyp growth than higher or lower elevation of Wnt signaling; the “just right” hypothesis (34, 35).

Wnt signaling has been assessed in many *Apc* mouse models. Some models have high polyp multiplicity and show elevated Wnt signaling in these polyps (*Apc*<sup>Min/+</sup>, *Apc* <sup>$\Delta$ 716/+</sup>, *Apc*<sup>1322T/+</sup> and *Apc* <sup>$\Delta$ e1-15/+</sup>) (10, 34, 35). Wnt signaling is also elevated in the few polyps that develop in *Apc*<sup>NeoR/+</sup> and *Apc*<sup>NeoF/+</sup> mice (36, 37). *Apc*<sup>mNLS/mNLS</sup> mice have elevated Wnt signaling in intestinal epithelial cells (38, 39). *Apc*<sup>1572T/1572T</sup> embryonic stem cells also have elevated Wnt signaling (38, 39). Neither *Apc*<sup>mNLS/mNLS</sup> nor *Apc*<sup>1572T/+</sup> mice develop intestinal polyps (38, 39).

The “just right” hypothesis is supported by reports of increased polyp multiplicity in *Apc*<sup>1322T/+</sup> and *Apc* <sup>$\Delta$ e1-15/+</sup> mice relative to *Apc*<sup>Min/+</sup> mice (34, 35). Compared to *Apc*<sup>Min</sup>, *Apc*<sup>1322T</sup> protein retains one 20 a.a. repeat which can bind to  $\beta$ -catenin and decrease Wnt signaling (34, 35). The *Apc* <sup>$\Delta$ e1-15</sup> allele results in complete deletion of *Apc* and polyps in *Apc* <sup>$\Delta$ e1-15/+</sup> mice also display less Wnt signaling than polyps in *Apc*<sup>Min/+</sup> mice (34). However, the “just right” hypothesis does not readily explain why *Apc* <sup>$\Delta$ 716/+</sup> mice show higher activation of Wnt signaling and more polyps than *Apc*<sup>Min/+</sup> mice (40). In addition, several groups have reported that although loss of both *Apc* alleles is required to activate Wnt signaling (as assessed by nuclear translocation of  $\beta$ -catenin), this *Apc* loss is not sufficient for full Wnt signal activation (11, 41, 42). To establish the extent to which Wnt signaling and polyp growth contribute to phenotypic variation, Wnt signaling activities and proliferation rates must be directly compared in different *Apc* mouse models.

**3- Different abilities to evade growth inhibitory effects**—Another explanation of variation in polyp number among different *Apc* mouse models is negative selection of particular *Apc* genotypes. This negative selection could contribute to the “Just right” hypothesis. Support for negative selection contributing to polyp phenotypes is provided by the observation that addition of *Cdx2* or *BubR1* mutations to *Apc* <sup>$\Delta$ 716/+</sup> or *Apc*<sup>Min/+</sup> mice, respectively, results in reduced polyp multiplicity and increased apoptotic indices in the small intestines, despite the increased proliferation index in these cells (43, 44). Similarly, induction of a conditional *Apc* mutation in hematopoietic stem cells results in upregulation of Wnt signaling and increased stem cell proliferation with increased apoptosis and eventual exhaustion of the stem cell population (45). If this phenotype holds true for intestinal tissues, the “just right” hypothesis might explain the increased stem cell number in polyps from *Apc*<sup>1322T/+</sup> mice relative to those from *Apc*<sup>Min/+</sup>, despite lower Wnt signaling in polyps from the former model relative to those from *Apc*<sup>Min/+</sup> mice.

**4- Distinctive effects on differentiation**—It is possible that the effect of *Apc* genotypes on enterocyte differentiation contributes to differences in intestinal polyp number. For instance, compared to *Apc*<sup>Min/+</sup> mice, *Apc*<sup>1322T/+</sup> mice have a higher proportion of Paneth cells and cells that express stem cell markers (*Lgr5*, *Bmi1*, *Msi1* and *CD44*), not only in adenomas but also in apparently normal intestinal epithelial cells (35). Cell fates that result from different *Apc* genotypes might alter tumor initiation or growth. Again, Wnt signaling is one of several factors proposed to affect differentiation.

**5- Contributions of genetic modifiers or environmental factors**—It is well established that genetic and environmental factors affect intestinal polyp multiplicity in *Apc* mouse models. Polyp multiplicity in *Apc*<sup>Min/+</sup> mice varies greatly between laboratories (20–100/mouse) (17, 18). This inconsistency might result from variations in diet, emergence of genetic modifiers, and even from different methods of polyp detection. A genetic modifier is a genetic locus that modifies the effect produced by a non-allelic locus. Modifier genes are present in different mouse strains and can even emerge in what is considered a congenic strain (46). Several modifier loci have been found to affect intestinal polyposis in *Apc*<sup>Min/+</sup> mice and are named *modifier of min (Mom)* (reviewed in (18)). Some modifiers are single genes, others are thought to represent contiguous genes, and some remain less well-defined (47). The modifiers appear to function as recessive, dominant or semi-dominant loci (17). The first identified modifier gene, *Mom-1 (Pla2g2a)*, works in a cell-non-autonomous manner, possibly by reducing inflammatory response in the gut (48–50). The *Mom-2 (Atp5a1)* allele is on the same chromosome as *Apc* (chromosome 18) and appears to inhibit loss of the wild-type *Apc* allele (48, 51). The mechanisms of action of other modifiers such as *Mom-3*, *Mom-7*, *Mom-12* and *Mom-13* are not understood (52–54).

Though identified in *Apc*<sup>Min/+</sup> mice, *Mom* genes likely also affect phenotypes of other *Apc* mouse models. For instance, the C3H/HeJ mouse strain carries at least one *Mom* allele that is absent from the C57BL/6 strain, *Mom-1* (48). Both *Apc*<sup>Min/+</sup> and *Apc*<sup>Δ242/+</sup> mice show reduced polyp multiplicity in the first generation mixed C57BL/6: C3H/HeJ mice compared to in C57BL/6 mice (55). At present, there appears to be no direct examination of the effect of specific *modifiers of Min* on different *Apc* mouse models.

Environmental factors, such as intestinal flora, might also contribute to phenotypic variation (56). While intestinal flora appear to increase the number of polyps in *Apc*<sup>Min/+</sup> mice (57), *Apc*<sup>Δ14/+</sup> mice raised in pathogen-free conditions showed significant increases in intestinal polyp number (58).

Diet is another major environmental factor that clearly impacts the mouse phenotype (59–61). Although typically defined, the concentration of various vitamins, fiber, and total fat varies greatly between laboratory mouse diets. In our own experience, switching the mouse diet had a dramatic effect on polyp multiplicity in our *Apc*<sup>Min/+</sup> mouse colony. We found that the polyp burden per mouse significantly increased from 45.9±4.5 in 10 *Apc*<sup>Min/+</sup> mice on Lab diet 5001 (Purina) to 81±9.3 in 25 age-matched *Apc*<sup>Min/+</sup> mice on Harlan 2018 diet (*p*= 0.0006). Notably, the new diet (Harlan 2018) has a 24% increase in fat and decreased fiber, vitamin D, and folic acid by 42%, 67%, and 44%, respectively. Unfortunately, these inter-laboratory variables such as diet confound direct comparison of the phenotypes of *Apc* mouse models studied in different laboratories.

**6- Differences in cellular migration and adhesion**—APC interaction with cytoskeletal components, including actin filaments and microtubules, is thought to affect cell adhesion and migration (62, 63). Decreased cellular adhesion and migration in cells with *APC* mutations is expected to contribute to tumor formation (64). APC interacts with cytoskeletal proteins through its C-terminal region, which is absent in *Apc* from most mouse

models (figure 1). Adding the C-terminal *Apc* region to *Apc*<sup>1322T</sup> (as in *Apc*<sup>ΔSAMP</sup> mice) did not change the phenotype (65). However, it is possible that cytoskeletal alterations affect later stages of tumor progression such as invasion and metastasis, which do not occur in most *Apc* mouse models (66). Currently, evidence supporting a direct role of the *Apc* C-terminus in intestinal phenotype variation among different *Apc* mouse models is lacking.

**7-Differences in technologies used to generate the mouse model**—*Apc* rodent models have been generated using 3 different technologies; chemical mutagenesis screen, insertion of an antibiotic-resistance gene and Cre-lox induced DNA excision. The *Apc*<sup>Min/+</sup> mouse, PIRC rat and KAD rat were generated by chemical mutagenesis which resulted in a single base-pair change in the *Apc* gene (9, 67, 68). Many other models, such as *Apc*<sup>1309</sup>, *Apc*<sup>1638N</sup> and *Apc*<sup>1638T</sup>, were generated through insertion of an antibiotic-resistance gene into the *Apc* gene, thus introducing a nonsense mutation (69–71). In *Apc*<sup>neoF</sup> and *Apc*<sup>neoR</sup> alleles, the antibiotic-resistance gene disrupts an enhancer sequence in intron 13 (36, 37). Other mouse models with *Apc* truncation including *Apc*<sup>1322T/+</sup> and *Apc*<sup>Δe1-15</sup> were generated using Cre-lox mediated-deletion of specific *Apc* regions. The later technology allowed removal of most exogenous DNA sequences originating from the targeting vector including the antibiotic-resistance gene. The *Apc*<sup>mNLS</sup> model contains mutations “knocked into” the *Apc* gene, with the antibiotic-resistance gene subsequently removed by Cre-Lox-mediated deletion (39).

The *Apc*<sup>1638N/+</sup> and *Apc*<sup>1638T/+</sup> models, which differ only by orientation of the inserted neomycin-resistance gene, provide clear evidence for the contribution of extraneous DNA to phenotypic variation (69). *Apc*<sup>1638N/+</sup> mice express so little truncated *Apc* protein that they might be considered virtually null (69, 72); yet the described phenotype of *Apc*<sup>1638N/+</sup> mice is not similar to that of the *Apc*<sup>Δe1-15</sup> model, which has a complete deletion of the *Apc* gene (34, 72). The neomycin-resistance gene clearly affects the phenotypes of these mice and if inserted in reverse orientation, might affect not only *Apc* expression, but also expression of genes upstream of *Apc*. It is possible that the 6-fold difference in intestinal polyp number between *Apc*<sup>1322T/+</sup> and *Apc*<sup>1309/+</sup> mice, which differ by only 13 amino acids, stems from the different technology used in their generation; Cre-lox-mediated deletion in *Apc*<sup>1322T/+</sup> versus insertion of an antibiotic resistance cassette in *Apc*<sup>1309/+</sup>. However, other genetic and environmental factors may contribute to the variation between these two mouse models as well (32, 70). A final illustration of the challenges in generation of *Apc* mouse models is the *Apc*<sup>Δ474/+</sup> mice, which have a duplication of *Apc* exons 7–10. This feature complicates dissection of the contribution of exon duplication to the phenotype (73).

**8-Differences in expression of the mutant allele**—When analyzing the phenotypes of different *Apc* mouse models, another consideration is the level of expression of the mutant allele. *Apc* is a large multi-domain protein. Truncations of *Apc* in most FAP patients and rodent models leave N-terminal domains intact, figure 1. Although normal expression levels of truncated *Apc* protein have been verified in *Apc*<sup>Δ716</sup>, *Apc*<sup>Min/+</sup>, *Apc*<sup>1322T</sup>, and *Apc*<sup>1638T</sup> mice, this is not universally the case (32, 69, 74). In *Apc*<sup>580D</sup>, *Apc*<sup>Δ14</sup>, *Apc*<sup>Δ474</sup>, and *Apc*<sup>Δ242</sup> models, the truncating mutation occurs before the final exon (15), and thus there is the possibility of a nonsense-mediated RNA decay. Truncated *Apc* was not detected in intestinal polyps from *Apc*<sup>Δ580/+</sup> mice or ES cells from *Apc*<sup>Δ15/+</sup> mice (75, 76), which suggests that these alleles might also be virtually null. A related consideration is the effect of the introduced mutation (and possibly the antibiotic selection cassette) on *Apc* folding. Although most of *Apc* is thought to be natively unfolded (77), the effects of mutations on inherently folded domains of *Apc*, and the consequences of potential folding defects in relation to phenotype, are not understood.

## II- Variation in polyp distribution

Tumors in most *Apc* mouse models occur mainly in the small intestine, while germline mutations of *APC* in humans result in tumors predominantly in the large intestine (21, 78). The PIRC *Apc* rat model has tumors in both small and large intestines (9, 13, 79). A pig model with germline *Apc* mutations was recently reported to develop polyps in the colon (80). In addition to this inter-species variation, mouse models with different germline *Apc* mutations show different distributions of intestinal polyps. Analysis of  $Apc^{Min/+}$  mice with different genetic backgrounds has led to the hypothesis that polyp distribution is somehow linked to the mechanism by which the wildtype *Apc* allele is lost (24). Haigis *et al.* showed that in a B6 background,  $Apc^{Min/+}$  mice develop polyps mainly in the distal half of the small intestine, and loss of the wildtype *Apc* allele occurs by means of LOH. In an AKR background,  $Apc^{Min/+}$  mice develop polyps predominantly at the ileo-cecal junction, and inactivation of the wildtype *Apc* allele is achieved through allelic silencing. In the B6 background,  $Apc^{Min/+}$  mice with additional mutations that inactivate the mismatch repair gene *Mlh* develop polyps all over the small intestine, and loss of the wildtype *Apc* allele is achieved through a point mutation.  $Apc^{1638N/+}$  mice develop polyps in a similar distribution, and appear to retain the wildtype *Apc* allele (24).

Mechanistically, two models have been proposed to explain the connection between polyp distribution and loss of the wildtype *Apc* allele. In the first model, the molecular machinery in different intestinal regions determines the mechanism of the second *Apc* “hit” and hence the distribution of polyps. This model is supported by the finding that mice in which the wildtype *Apc* allele is inactivated by the same mechanism (eg.  $Apc^{Min/+}/Mlh^{-/-}$ , and  $Apc^{1638N/+}$ ) have similar polyp distributions (24). However, the finding that both  $Apc^{1322T/+}$  and  $Apc^{Min/+}$  mice lose the wildtype *Apc* allele through LOH, yet have different polyp distributions, does not support this model. A second model proposes that polyp growth is dictated by the *Apc* status but also by the particular environment of the different intestinal regions, independent of the mechanism of the second *Apc* mutation. Supporting this hypothesis,  $Apc^{\Delta 716/+}$  mice with an additional mutation of *Cdx2* exhibit more colonic and fewer small intestinal polyps. Yet, loss of the wildtype *Apc* allele occurs via LOH regardless of *Cdx2* status (44). Similarly, a colonic shift of polyps has been described in  $Apc^{Min/+}$  mice with an additional *BubR1* mutation, although the mechanism of loss of the wildtype *Apc* allele in these mice was not reported (43). Mutation of both *Cdx2* and *BubR1* increases chromosomal instability and changes the proliferation and apoptotic indices in intestines of  $Apc^{\Delta 714/+}$  and  $Apc^{Min/+}$  mice, respectively (43, 44). Further support for the second model comes from  $Apc^{Min/+}$  mice in a 129/Sv background, where additional mutations that inactivate *Smad3* result in increased colonic tumors; yet in both cases, loss of the wildtype *Apc* allele is achieved through LOH (81). Finally, PPAR $\gamma$  agonists increase colonic but not small intestinal tumors in  $Apc^{Min/+}$  mice (82, 83). PPAR $\gamma$  is expressed in higher quantities in the colon and cecum relative to the small intestine, that might account for this differential effect (83).

An expansion of the “just right” hypothesis has been proposed to explain the variation in polyp distribution among FAP patients,  $Apc^{Min/+}$  and  $Apc^{1322T/+}$  mice. The basal level of Wnt signaling is not the same in different intestinal regions. It was proposed that changes in Wnt signaling that result from specific *Apc* mutations cause optimal Wnt signaling for polyp growth only in certain intestinal regions. On the other hand, in other intestinal regions, these same *Apc* mutations will result in a higher or lower Wnt signaling level than what is optimal for tumor growth (84).

Perhaps some of these mechanisms can be clarified by studying  $Apc^{Min-FCCC}$  mice which were generated by mating C57Bl/6J  $Apc^{Min/+}$  males with  $Apc^{+/+}$  females from an independent colony of C57Bl/6 mice maintained at Fox Chase Cancer Center.

$Apc^{Min-FCCC/+}$  mice develop more colon polyps than do  $Apc^{Min/+}$  mice, but the molecular basis behind this polyp shift has not been determined (85). Further clarification of the underlying mechanisms that control polyp distribution might also be achieved through careful analysis of  $Apc^{\Delta14/+}$  and  $Apc^{580D/+}$  mice, which carry similar mutations (truncating the *Apc* protein at amino acid 580) but appear to have different polyp distributions.  $Apc^{\Delta14/+}$  mice develop more colonic polyps than do  $Apc^{Min/+}$  mice.  $Apc^{580D/+}$  mice develop a similar number of colonic polyps as  $Apc^{Min/+}$  mice, although direct comparison of  $Apc^{580D/+}$  and either  $Apc^{\Delta14/+}$  or  $Apc^{Min/+}$  mice has not been reported (75, 86).

### III- Variation in extra-intestinal phenotypes

Although best known for its role to suppress colorectal tumorigenesis, *APC* mutations have been seen in other tumors including breast and liver carcinomas (4). In addition, both FAP patients and rodent models with germline *Apc* mutations develop extra-intestinal phenotypes (see table 1). As with the intestinal phenotype, the underlying mechanism for variation in extra-intestinal phenotypes between FAP patients and *Apc* rodent models as well as among different *Apc* rodent models is not completely understood. FAP patients have increased susceptibility to hepatic, pancreatic, thyroid and brain tumors. They also develop desmoid tumors, dental anomalies, and congenital hypertrophy of retinal pigment epithelium. It is important to note that the penetrance of these extra-intestinal phenotypes is variable in FAP patients (16, 87). The basis behind this variation is not completely understood, although it seems to correlate with the *APC* germline as well as the acquired somatic mutations. (16, 33).

*Apc* rodent models also develop some of these extra-intestinal manifestations, for example,  $Apc^{1638N/+}$  mice develop desmoid tumors (72) and PIRC rats show mandibular osteoma (9). Other phenotypes described in FAP patients have not been reported for *Apc* rodent models. The short life span of most *Apc* rodent models could prevent the full expression of some of these phenotypes. On the other hand, *Apc* rodent models manifest some other extra-intestinal phenotypes that have not been described in FAP patients (table 1). For example, many mouse models with germline *Apc* mutations develop mammary tumors. Although *APC* mutations and promoter methylation have been found in up to 70% of sporadic human breast cancers, FAP patients do not appear at an increased risk for breast tumors (88–90). In addition, adenoacanthoma is a common type of mammary tumor that develops in *Apc* mouse models but it has not been reported in humans (91). Other extra-intestinal phenotypes described in *Apc* rodent models include; splenomegaly, abnormal hematopoiesis, changes in the serum lipid profile, gonadal changes, cutaneous cysts, and thyroid abnormalities. Differences in physiology, life span and genetic content between human, mouse and rat could be underlying causes.

Among different *Apc* mouse models, some extra-intestinal phenotypes, such as anemia and splenomegaly, seem to correlate with the severity of intestinal polyposis. In contrast, mammary gland tumors in *Apc* mouse models appear to correlate with the severity of polyposis in only a few cases, such as in the  $Apc^{Min/+}$  and  $Apc^{\Delta474/+}$  models. Very few  $Apc^{Min/+}$  mice develop mammary tumors, whereas  $Apc^{\Delta474/+}$  mice develop mammary tumors at a rate that is almost double that seen in  $Apc^{Min/+}$  mice (73, 91). In contrast, there are no reports of mammary tumor development in *Apc* mouse models with the most severe intestinal polyposis ( $Apc^{\Delta714}$ ,  $Apc^{1322T}$ , and  $Apc^{\Delta SAMP}$ ) (32, 40, 65). Perhaps mice with severe polyposis die too early, before mammary tumors have a chance to develop.  $Apc^{1572T/+}$  mice, which develop no intestinal polyps, have a fully-penetrant mammary tumor phenotype in females. K14-cre- $Apc^{CKO/+}$  mice are a conditional model in which the  $Apc^{\Delta580}$  allele is expressed only in ectoderm-derived tissues including the mammary gland (75, 92). Mammary tumors from these mice have mutations in the wildtype *Apc* allele that cluster around codon 1530 consistent with the requirement of an optimal level of Wnt

signaling for mammary tumorigenesis (38). It is likely that some of the genetic and environmental factors previously described also account for the variability in extra-intestinal phenotypes among different *Apc* rodent models.

## Conclusions and future directions

APC research has benefitted greatly from different rodent models with germline *Apc* mutations. However genotype/phenotype correlation of these different models is confounded by many genetic and environmental factors. Use of standardized genetic backgrounds and environmental conditions in different laboratories should enable reliable genotype/phenotype analysis of these animals. This standardization will also shed light on the role of different *Apc* mutations in tumorigenesis. When possible, a direct comparative analysis of different models in the same laboratory will illuminate the contribution of many factors described in this review to phenotypic variation in rodent models with germline *Apc* mutations.

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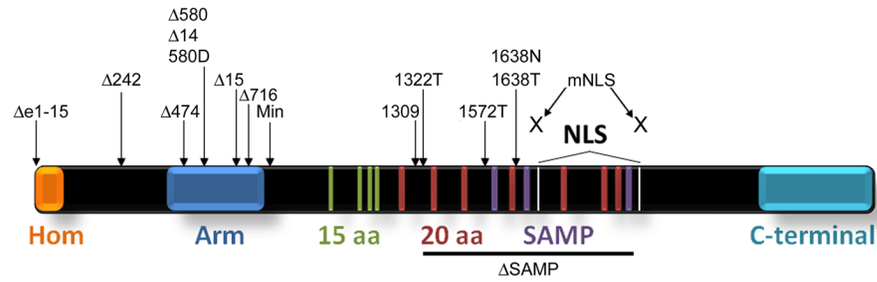
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**Figure 1. Sites of *Apc* mutations in different *Apc* mouse models relative to *Apc* domains**  
 Domains of *Apc* are indicated as follows: Hom = homodimerization, Arm = Armadillo repeats, 15 aa = 15 amino acid repeats, 20 aa = 20 amino acid repeats, Serine-Alanine-Methionine-Proline (SAMP)= axin binding, NLS=nuclear localization signals, and C-terminal includes microtubule, EB1 and PDZ binding domains. The Mutation Cluster Region (MCR) is between codons 1250 and 1464.

**Table 1**Summary of rodent models with germline *Apc* mutations before MCR \*#

Model (Ref)	<i>Apc</i> mutation	Intestinal phenotype	Polyp distribution	Extra-intestinal phenotype
<b>Apc<sup>Δe1-15/+</sup></b> (34)	<ul style="list-style-type: none"> <li>Complete deletion of entire <i>Apc</i> gene</li> </ul>	<ul style="list-style-type: none"> <li>~160/male, ~190/female</li> <li>Benign adenoma</li> <li>Polyps show similar histopathology to those in <i>Apc<sup>Min/+</sup></i> mice</li> </ul>	<ul style="list-style-type: none"> <li>Similar distribution as in <i>Apc<sup>Min/+</sup></i> mice</li> </ul>	<ul style="list-style-type: none"> <li>Anemia</li> </ul>
<b>Apc<sup>Δ242/+</sup></b> (55)	<ul style="list-style-type: none"> <li>β-geo gene trap cassette inserted between exons 7 and 8 leads to stop after codon 242</li> </ul>	<ul style="list-style-type: none"> <li>177 polyps</li> <li>Benign adenoma</li> <li>Polyps show similar histopathology as those in <i>Apc<sup>Min/+</sup></i> mice</li> </ul>	<ul style="list-style-type: none"> <li>Similar distribution as in <i>Apc<sup>Min/+</sup></i> mice</li> </ul>	<ul style="list-style-type: none"> <li>NR</li> </ul>
<b>Apc<sup>Δ474/+</sup></b> (86)	<ul style="list-style-type: none"> <li>Insert of duplicated exons 7–10 leads to frameshift and stop after codon 474</li> </ul>	<ul style="list-style-type: none"> <li>122 polyps</li> <li>Benign adenoma</li> </ul>	<ul style="list-style-type: none"> <li>Mainly small intestine (SI)</li> <li>Some in colon and stomach</li> </ul>	<ul style="list-style-type: none"> <li>Mammary tumors in 18.5% females at 3–5 months (adenoacanthoma)</li> </ul>
<b>Apc<sup>Δ580/+</sup></b> (75)	<ul style="list-style-type: none"> <li>Exon 14 deletion leads to frameshift and stop after codon 580</li> </ul>	<ul style="list-style-type: none"> <li>120 polyps</li> <li>Adenomas</li> </ul>	<ul style="list-style-type: none"> <li>Mainly SI</li> </ul>	<ul style="list-style-type: none"> <li>Anemia</li> </ul>
<b>Apc<sup>Δ14/+</sup></b> (86)	<ul style="list-style-type: none"> <li>Exon 14 deletion leads to frameshift and stop after codon 580</li> </ul>	<ul style="list-style-type: none"> <li>36 polyps</li> <li>Benign adenoma to invasive carcinoma</li> <li>More polyps in germ-free environment</li> <li>Rectal prolapse (61%)</li> </ul>	<ul style="list-style-type: none"> <li>SI</li> <li>More colonic tumors than <i>Apc<sup>Min/+</sup></i> mice</li> </ul>	<ul style="list-style-type: none"> <li>Mammary tumors (9%)</li> <li>Anemia</li> </ul>
<b>Apc<sup>580D/+</sup></b> (93)	<ul style="list-style-type: none"> <li>Exon 14 deletion leads to frameshift and stop after codon 580</li> </ul>	<ul style="list-style-type: none"> <li>Intestinal polyposis</li> </ul>	<ul style="list-style-type: none"> <li>NR</li> </ul>	<ul style="list-style-type: none"> <li>NR</li> </ul>

Model (Ref)	<i>Apc</i> mutation	Intestinal phenotype	Polyp distribution	Extra-intestinal phenotype
<b>Apc<sup>Δ15/+</sup></b> (76)	<ul style="list-style-type: none"> <li>Deletion of the last exon (exon 15) including 3'UTR</li> </ul>	<ul style="list-style-type: none"> <li>185 polyps</li> <li>Adenoma</li> <li>Few adenocarcinoma</li> <li>Normal crypt maturation gradient lost</li> </ul>	<ul style="list-style-type: none"> <li>Mostly SI</li> <li>77% in Ileum</li> </ul>	<ul style="list-style-type: none"> <li>Cutaneous cysts</li> <li>Desmoid tumors</li> <li>Anemia</li> </ul>
<b>Apc<sup>Δ716/+</sup></b> (10, 40)	<ul style="list-style-type: none"> <li>Inserted <i>Neo<sup>R</sup></i> and diphtheria toxin <math>\alpha</math>-subunit genes in exon 15 leads to stop after codon 716</li> </ul>	<ul style="list-style-type: none"> <li>58–256 polyps</li> <li>Benign adenomas</li> </ul>	<ul style="list-style-type: none"> <li>Mainly SI</li> </ul>	<ul style="list-style-type: none"> <li>Anemia</li> </ul>
<b>Apc<sup>Min/+</sup></b> (19, 67, 79)	<ul style="list-style-type: none"> <li>Generated by ENU screen.</li> <li>Nonsense mutation after codon 850</li> </ul>	<ul style="list-style-type: none"> <li>20–100 polyps</li> <li>Benign adenomas</li> <li>Malignant transformation in old mice in some genetic backgrounds</li> </ul>	<ul style="list-style-type: none"> <li>60% in distal 1/3 of the SI</li> <li>Few in colon</li> <li>Very few in stomach</li> </ul>	<ul style="list-style-type: none"> <li>Mammary tumors; 5% old females</li> <li>Anemia</li> <li>Splenomegaly</li> <li>Abnormal hematopoiesis</li> <li>Degeneration of ovarian follicles</li> <li>Underdeveloped seminiferous tubules</li> <li>Abnormal serum lipid profile</li> </ul>
<b>PIRC rat</b> (9, 94)	<ul style="list-style-type: none"> <li>Nonsense mutation after codon 1137</li> </ul>	<ul style="list-style-type: none"> <li>36 polyps and 178 microadenoma (less than 0.5mm), males</li> <li>11 polyps and 35 microadenomas, females</li> <li>Adenoma</li> <li>Adenocarcinoma in older mice</li> </ul>	<ul style="list-style-type: none"> <li>Tumors are in both SI and colon</li> </ul>	<ul style="list-style-type: none"> <li>Benign epidermoid cysts</li> <li>Jaw osteoma in old females</li> </ul>

**Table 2**Summary of rodent models with germline *Apc* mutations within or after MCR <sup>\*/#</sup>

Model (Ref)	<i>Apc</i> mutation	Intestinal phenotype	Polyp distribution	Extra-intestinal phenotype
<b>Apc<sup>1309/+</sup></b> (70, 95, 96)	<ul style="list-style-type: none"> <li><i>Neo<sup>R</sup></i> gene inserted</li> <li>Truncation after codon 1309</li> </ul>	<ul style="list-style-type: none"> <li>33–37 polyps on average</li> <li>Benign adenoma</li> </ul>	<ul style="list-style-type: none"> <li>Mainly SI</li> <li>Few, stomach and colon</li> <li>SI polyps more proximal than <i>Apc<sup>Min/+</sup></i>; only 1/3 distal</li> </ul>	<ul style="list-style-type: none"> <li>Centrilobular cholestasis in liver</li> <li>Microvesicular fatty liver</li> <li>Abnormal serum lipid profile</li> </ul>
<b>Apc<sup>1322T/+</sup></b> (32, 35)	<ul style="list-style-type: none"> <li>Deletion after codon 1322</li> </ul>	<ul style="list-style-type: none"> <li>200 polyps</li> <li>Benign adenomas with severe dysplasia in large polyps</li> <li>Polyps have less Wnt signaling but more stem cells relative to those from <i>Apc<sup>Min/+</sup></i> mice</li> </ul>	<ul style="list-style-type: none"> <li>Most in SI</li> <li>Few in colon &amp; stomach</li> <li>SI polyps more proximal (less than 20% in distal 1/3 of SI)</li> </ul>	<ul style="list-style-type: none"> <li>Anemia</li> <li>Splenomegaly</li> </ul>
<b>Apc<sup>1572T/+</sup></b> (38)	<ul style="list-style-type: none"> <li>PGK-Hygromycin cassette inserted in sense orientation.</li> <li>Stop at codon 1572</li> </ul>	None	<ul style="list-style-type: none"> <li>N/A</li> </ul>	<ul style="list-style-type: none"> <li>Mammary invasive adenocarcinoma in 100% of females and 30% of males</li> </ul>
<b>Apc<sup>1638T/1638T</sup></b> (69, 97)	<ul style="list-style-type: none"> <li>PGK-Hygromycin cassette inserted in sense orientation.</li> <li>Stop at codon 1638</li> </ul>	None	N/A	<ul style="list-style-type: none"> <li>Viable homozygous mutant</li> <li>Post-natal growth retardation</li> <li>Cutaneous cysts in nipples</li> <li>Absent preputial glands</li> <li>Aberrant response of thyroid gland to thyroid stimulating hormone</li> </ul>
<b>Apc<sup>1638N/+</sup></b> (71)	<ul style="list-style-type: none"> <li><i>Neo<sup>R</sup></i> gene inserted in antisense orientation.</li> <li>Stop after codon 1638</li> </ul>	<ul style="list-style-type: none"> <li>&lt;10 polyps</li> <li>Benign adenoma and adenocarcinoma</li> <li>Aberrant crypt foci</li> </ul>	<ul style="list-style-type: none"> <li>SI, colon and stomach</li> <li>Uniformly distributed along SI</li> </ul>	<ul style="list-style-type: none"> <li>Desmoid tumors</li> <li>Cutaneous cysts</li> </ul>

Model (Ref)	<i>Apc</i> mutation	Intestinal phenotype	Polyp distribution	Extra-intestinal phenotype
		<ul style="list-style-type: none"> <li>Liver metastasis in one mouse</li> </ul>		
<b>KAD rat</b> (68)	<ul style="list-style-type: none"> <li>A non-sense mutation in <i>Apc</i> codon 2523</li> </ul>	<ul style="list-style-type: none"> <li>No spontaneous intestinal tumors</li> <li>Homozygous mutant rats have increased incidence and multiplicity of colonic tumors when treated with AOM-DSS relative to treated wildtype rats</li> </ul>	<ul style="list-style-type: none"> <li>Colon (AOM-DSS-induced)</li> </ul>	<ul style="list-style-type: none"> <li>Homozygous mutant animals are viable</li> </ul>

\* *Apc* mouse models reported in this table are on C57Bl/6 background however, with different backcross isogenicity from N2 to > N20

# *Apc* rat models reported in the table are on F344 background

*Apc* models are mouse models unless otherwise noted

NR: not reported

*Neo<sup>R</sup>*: Neomycin resistance gene

Table 3

Summary of mouse models with other germline *Apc* mutations \*

Model (Ref)	<i>Apc</i> mutation	Intestinal phenotype	Polyp distribution	Extra-intestinal phenotype
<b>Apc<sup>mNLS/mNLS</sup></b> (39)	<ul style="list-style-type: none"> <li>Inactivating mutations in the two nuclear localization signals (NLS)</li> </ul>	<ul style="list-style-type: none"> <li>Increased cellular proliferation in intestinal epithelial cells</li> <li>Few spontaneous intestinal polyps</li> <li>Enhanced polyposis in <b>Apc<sup>mNLS/Min</sup></b> mice</li> </ul>	N/A	NR
<b>Apc<sup>ΔSAMP</sup></b> (65)	<ul style="list-style-type: none"> <li>Deletion of codons between 1322 to 2006</li> </ul>	<ul style="list-style-type: none"> <li>Similar to <b>Apc<sup>1322T/+</sup></b></li> </ul>	<ul style="list-style-type: none"> <li>Similar to <b>Apc<sup>1322T/+</sup></b></li> </ul>	<ul style="list-style-type: none"> <li>Similar to <b>Apc<sup>1322T/+</sup></b></li> </ul>
<b>Apc<sup>NeoR</sup></b> and <b>Apc<sup>NeoF</sup></b> (36, 37)	<ul style="list-style-type: none"> <li><i>Neo<sup>R</sup></i> gene in intron 13 in reverse (<b>Apc<sup>NeoR</sup></b>) and forward (<b>Apc<sup>NeoF</sup></b>) direction.</li> <li>Reduced <i>Apc</i> level to 10% and 20%, respectively</li> </ul>	<ul style="list-style-type: none"> <li>0.2 polyps in <b>Apc<sup>NeoR</sup></b></li> <li>1 polyp in <b>Apc<sup>NeoF</sup></b></li> <li>Dysplastic adenomas</li> </ul>	<ul style="list-style-type: none"> <li>SI</li> </ul>	<ul style="list-style-type: none"> <li><b>Apc<sup>NeoR/NeoR</sup></b> embryos show severe developmental abnormalities and die in-utero</li> </ul>
<b>Apc<sup>Δ716/+</sup></b> (98)	<ul style="list-style-type: none"> <li>Mutant <i>Apc</i> allele truncated after codon 716 inserted as transgene in mouse with two wild-type <i>Apc</i> alleles</li> </ul>	<ul style="list-style-type: none"> <li>None</li> </ul>	NR	<ul style="list-style-type: none"> <li>Abdominal hamartoma in one mouse</li> </ul>
<b>Apc<sup>Δ716/Δ716/+</sup></b> (98)	<ul style="list-style-type: none"> <li>Mutant <i>Apc</i> truncated after codon 716 inserted as transgene in <b>Apc<sup>Δ716/+</sup></b></li> </ul>	<ul style="list-style-type: none"> <li>Similar to <b>Apc<sup>Δ716/+</sup></b></li> </ul>	<ul style="list-style-type: none"> <li>Similar to <b>Apc<sup>Δ716/+</sup></b></li> </ul>	<ul style="list-style-type: none"> <li>Similar to <b>Apc<sup>Δ716/+</sup></b></li> </ul>

\* *Apc* mouse models reported in this table are on C57Bl/6 background however, with different backcross isogenicity from N2 to > N20

All models are mouse models

NR: not reported

*Neo<sup>R</sup>*: Neomycin resistance gene