

Tales from the crypt

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Commentary

Intestinal colonic crypts are derived from a stem cell population located at the base of each crypt. A new analysis of mitochondrial function and of the rates of mitochondrial DNA (mtDNA) mutation in individual crypts shows that mtDNA mutations arise in stem cells — and at a surprisingly high frequency. Because crypts turn over extremely rapidly (about once per week), somatic mtDNA mutations can “take over the system” and even become homoplasmic, in a manner similar to what has been shown to occur in tumors.

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Tales from the crypt

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Intestinal colonic crypts are derived from a stem cell population located at the base of each crypt. A new analysis of mitochondrial function and of the rates of mitochondrial DNA (mtDNA) mutation in individual crypts shows that mtDNA mutations arise in stem cells — and at a surprisingly high frequency (see the related article beginning on page 1351). Because crypts turn over extremely rapidly (about once per week), somatic mtDNA mutations can “take over the system” and even become homoplasmic, in a manner similar to what has been shown to occur in tumors.

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Stem cells are the progenitors of specific cell lineages that become the body's organs and tissues during embryonic development. After birth, however, stem cells continue to play an equally important role in tissue maintenance, as they are called upon to repopulate cells that

turn over constantly. Hematopoietic stem cells were among the earliest identified exemplars of this role, but stem cells exist even in long-lived tissues — for example, muscle “satellite” cells — and, with the discovery in the last few years of stem cell lineages in brain and heart, our whole view of the idea of a “terminally differentiated” tissue has undergone a complete overhaul.

Mitochondrial dysfunction in stem cells

Mitochondria are semiautonomous organelles that are present in essentially all cells of the body. They contain their own DNA (mtDNA) and are the seat of a number of important

housekeeping functions. Foremost among these is the production of energy via the respiratory chain and oxidative phosphorylation, an intricate system composed of five complexes and two electron carriers (Figure 1a). The mtDNA (Figure 1b), a tiny 16.6 kb maternally inherited circular genome present in multiple copies in each organelle (there are about 10,000 mtDNAs in a typical cell), encodes 2 rRNAs, 22 tRNAs, and only 13 polypeptides, all of which are subunits of the respiratory complexes. In the last 15 years, mutations in mtDNA, all of which impair oxidative energy metabolism, have been found to cause a wide spectrum of disorders (1). In these patients, the mutations are typically heteroplasmic; that is, mutated mtDNAs coexist with wild-type mtDNAs in varying proportions, resulting in a mosaic pattern of respiratorily competent and incompetent cells. Respiratorily deficient cells must typically contain at least 80% mutated mtDNA to initiate dysfunction.

Heteroplasmic populations of mtDNA mutations can also arise randomly in somatic cells and can accumulate at low levels in individual cells during the course of normal aging (2). Even more intriguingly, somatic mtDNA mutations arise and are amplified in solid tumors, such as colon cancers (3), although a causative

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Nonstandard abbreviations used: mitochondrial DNA (mtDNA); cytochrome c oxidase (COX); nuclear DNA (nDNA); succinate dehydrogenase (SDH).

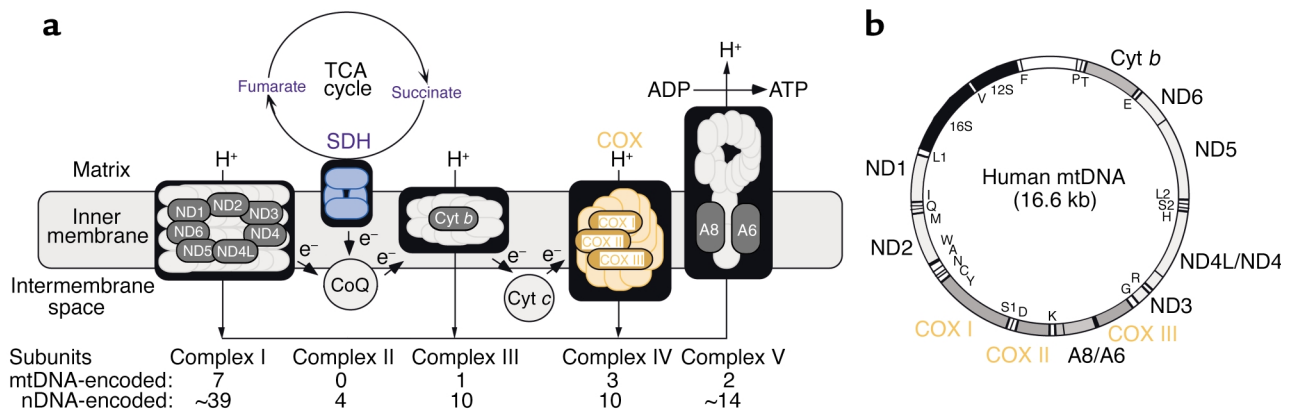


Figure 1
 (a) The respiratory chain. Nuclear DNA-encoded subunits are light gray; mtDNA-encoded subunits (see panel b) are dark gray. (b) Map of the human mitochondrial genome. Polypeptide-coding gene products (outside the circle) specify 7 subunits of NADH dehydrogenase-CoQ oxidoreductase (ND), 1 subunit of CoQ-cytochrome *b* oxidoreductase (Cyt *b*), 3 subunits of COX, and 2 subunits of ATP synthase (A). Protein synthesis gene products (inside the circle) specify 12S and 16S rRNAs, and 22 tRNAs (one-letter code). Figure modified from Schon and Manfredi (10).

relationship between mtDNA mutations and tumorigenesis has not yet been established.

Mitochondria in *every* cell, even those that do not divide, turn over, because they replicate their DNA and divide independently of the cell cycle. In a sense, then, as befitting an organelle that is derived evolutionarily from bacteria, mitochondria are each cell's own "stem" population, continually dividing and replacing themselves within their "hosts."

If mtDNA is always replicating within cells, it stands to reason that somatic mtDNA mutations could arise in stem cells as well, but the confirmation of this supposition, both qualitatively (does it happen?) and quantitatively (at what rate?), has been lacking. This question is not an academic one, as it goes to the heart of issues relating to the accumulation of mutated mtDNAs in disease (especially in the brain, a particularly susceptible tissue), in aging, and in tumorigenesis. It is also a hard question to answer for the simple reason that it has been extremely difficult to identify a stem cell population amenable to be studied easily and in sufficient quantity – until now. In this issue of the *JCI*, Taylor et al. (4) provide convincing evidence that at least one population of stem cells – those giving rise to intestinal colonic crypts – do indeed harbor somatic mtDNA mutations, which, even more

surprisingly, arise at a relatively high frequency. The decision to examine colonic crypts may have been inspired by earlier work on the segregation of mtDNA haplotypes in "transmitochondrial mice" by Shoubridge's group (5). The advantage of studying a colonic crypt is that it is a macroscopically observable clonal population of cells derived from one or two single cells – the stem cells – located at the base of each crypt (Figure 2). Thus, any mtDNA mutation found in the crypt in toto must perforce have been amplified from that very mutation in the stem cell itself, as the crypt is the surrogate of the stem cell.

The assumptions underlying the approach of Taylor et al. (4) were based on a simple syllogism: if (i) a stem cell harbors an mtDNA mutation that disrupts respiratory chain function and if (ii) that mutation expands (in the stem cell) to a level exceeding the threshold for dysfunction (typically >80% mutated mtDNA), then (iii) one ought to be able to observe both the dysfunction (by histochemical and/or biochemical means) and the mtDNA mutation (by genetic means) in the daughter population comprising the entire crypt. In order to assess function, Taylor et al. (4) applied a powerful method of mitochondrial analysis on serial transverse sections from individual human crypts: two-color histochemistry (6) to detect simultaneously the

enzymatic activities of cytochrome *c* oxidase (COX; complex IV of the respiratory chain), which contains both mtDNA- and nuclear DNA (nDNA)-encoded subunits; and succinate dehydrogenase (SDH; complex II of the respiratory chain), which contains only nDNA-encoded subunits. Importantly, when stained for both COX (brown) and SDH (blue) simultaneously, crypts with normal COX activity stained brown (the blue was hidden by the brown stain), whereas those with impaired COX activity stained blue (the brown was absent, revealing the blue). The authors then performed PCR coupled with mtDNA sequencing on the same transverse sections to search for somatic mutations that might correlate with the histochemistry. The two-color approach also allowed them to make a three-dimensional reconstruction of the COX activity in the entire crypt (Figure 2).

What Taylor et al. (4) found was, literally most illuminating. They observed three types of crypts: a majority of all-brown crypts, plus a minority consisting of both all-blue crypts and "mosaic" crypts containing ribbons of brown and blue cells. The first two patterns are consistent with the idea that the crypt has been repopulated from a single stem cell, either COX-normal (i.e., brown) or COX-deficient (blue). The most likely explanation for the third pattern (brown/blue mosaicism)

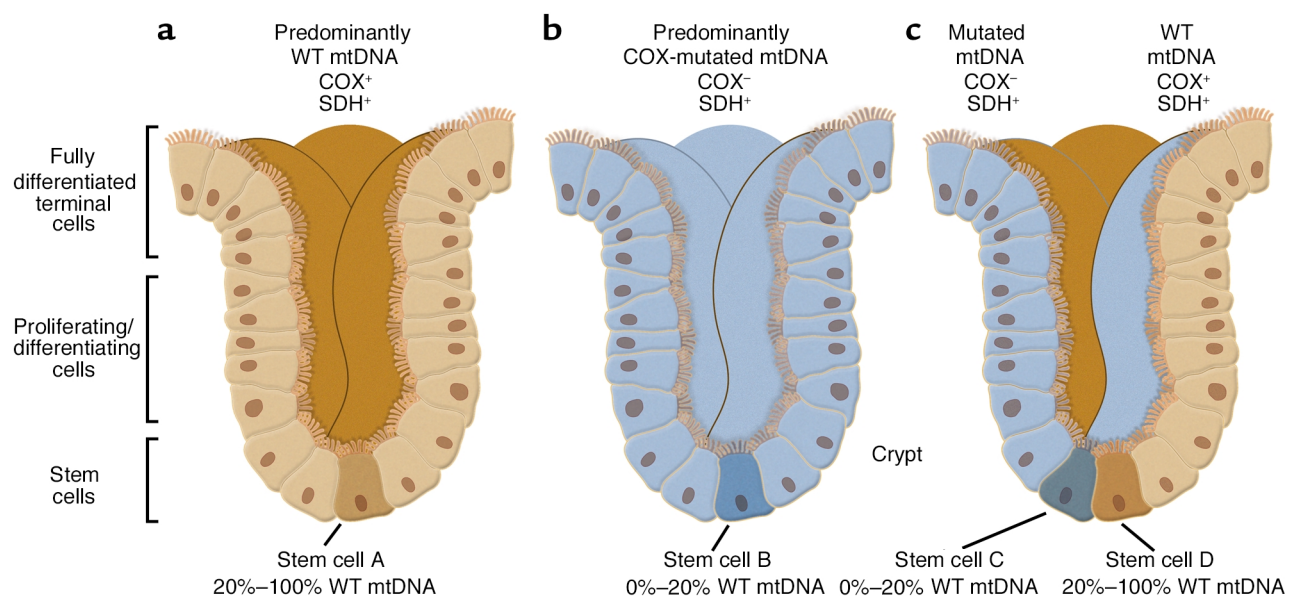


Figure 2

Two-color histochemistry for SDH and COX reveals three types of crypts, with normal (a), deficient (b), and “mosaic” (c) patterns of mitochondrial function, reflecting the phenotype in the stem cells that gave rise to each type of crypt.

is that at least two stem cells — one COX-normal and the other COX-deficient — were involved in crypt formation. Thus, all three patterns are consistent with the notion that a colonic crypt is indeed a clonal population derived from one or more stem cells.

Upon sequencing the sectioned crypts, they found many crypts with a normal mtDNA genotype (no surprise here), but in many crypts they found numerous mtDNA mutations, both in COX-negative crypts (many, but not all, mutations were in COX or protein synthesis genes) and in some of the ostensibly normal COX-positive crypts (mainly neutral mutations or mutations in non-COX genes). Interestingly, there were some COX-negative crypts in which no mtDNA mutations were found at all — presumably there were nuclear mutations in these stem cells that affected COX function. Overall, the amount of mutated mtDNA was extremely high (on average, one mtDNA point mutation per crypt), and, as has been known for a dozen years now (2), the mutational “load” increased with age.

Using mitochondrial function as a stem cell marker

So what is the significance of this work for “mitochondriacs” and for

stem cell biologists? First, we now know that mtDNA mutations can indeed arise in stem cells. Second, by mathematical modeling, Taylor et al. (4) were able to estimate the rate of somatic mtDNA mutation: it is approximately 5×10^{-5} per genome per day, a rate far greater than that of nuclear DNA. Third, the accumulation of mtDNA mutations, often to homoplasmy, in crypt stem cells is highly reminiscent of the dramatic shifts in mtDNA genotypes in solid tumors (3), which, of course, are clonal expansions of a tumor “stem cell.” However, unlike stem cells, tumors are aneuploid. Moreover, they also amplify segments of their chromosomes as “double minutes,” up to 5 Mb in size, that likely contain nuclear-embedded mtDNA pseudogenes. If these are amplified by PCR, pseudogene-derived polymorphisms may be attributed erroneously to mutations in authentic mtDNA.

Finally, the use of mitochondrial markers may allow researchers to track the progeny of multiple stem cells simultaneously. In fact, one can envision the use of COX-negative cells, especially in tissues from aged individuals, as a way to scan or “map” regions that are populated by stem cells. For example, the unusual non-

random distribution of COX-negative neurons in the hippocampus of brains of the elderly is likely due to random mtDNA somatic mutations arising in ependyma-derived *single* neuronal stem cells that repopulate *groups* of COX-negative neurons in the hippocampus (7). Similarly, the accumulation of COX-negative neurons in aging parvocellular, but not magnocellular, neurons of the lateral geniculate nucleus (8) may reflect the fact that a stem cell population exists for the former but not the latter.

Clearly, the use of colonic crypts as a stem cell “model system” allows investigators to address new types of questions in stem cell biology (9). From a mitochondrial standpoint, crypts may be of particular value, especially with regard to studying the mitochondrial population “bottleneck” that occurs during early oogenesis, and in understanding the dynamics of shifts from heteroplasmy to homoplasmy in mitochondrial disease.

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The TRAIL to arthritis

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Antigen-specific lymphocytes are involved in synovial proliferation within inflamed joints. Activated lymphocytes and synoviocytes from patients with rheumatoid arthritis express receptors that can bind TNF-related apoptosis-inducing ligand (TRAIL). A new study demonstrates that DCs pulsed with collagen and transduced with an adenovirus-based vector able to express TRAIL limit the incidence of arthritis in a model of collagen-induced arthritis and joint inflammation (see the related article beginning on page 1332). These results suggest that gene-modified cell therapy represents a therapeutic option for systemic rheumatic diseases.

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TNF-related apoptosis-inducing ligand (TRAIL) is a type II transmembrane protein that belongs to the TNF superfamily. It binds to death receptors (DRs) 4 and 5, two decoy receptors, and a soluble receptor called osteoprotegerin. The TRAIL signaling pathway was identified recently, and it has generated a great deal of interest since TRAIL induces apoptosis preferentially in tumor but not in normal cells, thus providing exciting opportunities for development of novel therapeutic strategies in cancer. TRAIL, like FasL, induces apoptosis by cross-linking and oligomerizing its receptors and forming a death-inducing signaling com-

plex through recruitment of an adapter molecule and the initiator caspase-8 and subsequent mitochondria-dependent or -independent activation of the downstream effector caspase-3. Resistance of tumor cells to TRAIL has been associated either with low expression of its receptors or with defects in the downstream signaling (1).

Rheumatoid arthritis is a chronic inflammatory disorder that affects up to 1% of the population. The exact origin and pathogenesis of the disease are still unknown, and numerous disease-modifying drugs and biologics have been tested. There is a significant need for increased efficacy and safety of these agents (2).

TRAIL controls negative selection of T cells in the thymus

Recent reports have claimed a central role for TRAIL in thymocyte selection. TRAIL^{-/-} mice have larger thymi, and immature CD4⁺CD8⁺ cells expressing high levels of heat-stable antigen are resistant to anti-CD3 antibody-mediated cell death. Similarly, TRAIL^{-/-} mice fail to reduce ovalbumin-specific cells following exposure to ovalbu-

min. Both experiments clearly show that TRAIL is essential for negative selection of T cells in the thymus (3). In vitro, TRAIL blockade enhances the accumulation of concanavalin-stimulated spleen T cells into the S-G2/M cell cycle phase, supporting that TRAIL is important in the control of the lymphocyte cell cycle (4).

TRAIL suppresses the development of arthritis

TRAIL^{-/-} mice are sensitive to the development of collagen-induced arthritis, probably because they fail to delete relevant T cell specificities and because they fail to properly silence activated T cells (3). As predicted, blockade of TRAIL with soluble DR5 administered systemically exacerbates arthritis, whereas direct transfer of a nonreplicative adenovirus expressing TRAIL into the joints of arthritic mice reduces arthritis (4). Injection of a TRAIL-expressing adenovirus into IL-1 β -induced arthritic joints also significantly limits synovial proliferation (5).

An anti-TRAIL receptor antibody has been shown to be quite effective in treating bone-erosive disease in a model that involves transfer of fibrosarcoma cells into mice (6). However, antibodies to DRs, including those against CD95, may be associated with hepatotoxicity (7, 8), precluding their use in the treatment of tumors and autoimmune diseases.

Collagen-pulsed TRAIL-expressing DCs suppress arthritis

In this issue of the *JCI*, Liu et al. (9) report suppression of collagen-induced arthritis using DCs pulsed with collagen and transfected with an adenovirus-based vector expressing the TRAIL gene under the control of the doxycycline-inducible (DOX-inducible) tetracycline response element. The system offered two novel features: DCs were primed to recognize collagen-spe-

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Nonstandard abbreviations used: TNF-related apoptosis-inducing ligand (TRAIL); death receptor (DR); doxycycline (DOX).