An “expressionistic” look at serrated precancerous colorectal lesions

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Research

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Abstract

Background: Approximately 60% of colorectal cancer (CRC) precursor lesions are the genuinely-dysplastic conventional adenomas (cADNs). The others include hyperplastic polyps (HPs), sessile serrated lesions (SSL), and traditional serrated adenomas (TSAs), subtypes of a class of lesions collectively referred to as “serrated.” Endoscopic and histologic differentiation between cADNs and serrated lesions, and between serrated lesion subtypes can be difficult.

Methods: We used in situ hybridization to verify the expression patterns in CRC precursors of 21 RNA molecules that appear to be promising differentiation markers on the basis of previous RNA sequencing studies.

Results: SSLs could be clearly differentiated from cADNs by the expression patterns of 9 of the 12 RNAs tested for this purpose (VSIG1, ANXA10, ACHE, SEMG1, AQP5, LINC00520, ZIC5/2, FOXD1, NKD1). Expression patterns of all 9 in HPs were similar to those in SSLs. Nine putatively HP-specific RNAs were also investigated, but none could be confirmed as such: most (e.g., HOXD13 and HOXB13), proved instead to be markers of the normal mucosa in the distal colon and rectum, where most HPs arise. TSAs displayed mixed staining patterns reflecting the presence of serrated and dysplastic glands in the same lesion.

Conclusions: Using a robust in situ hybridization protocol, we identified promising tissue-staining markers that, if validated in larger series of lesions, could facilitate more precise histologic classification of CRC precursors and, consequently, more tailored clinical follow-up of their carriers. Our findings should also fuel functional studies on the pathogenic significance of specific gene expression alterations in the initiation and evolution of CRC precursor subtypes.

Background

The World Health Organization's GLOBOCAN database currently shows colorectal cancer (CRC) as the third most commonly diagnosed cancer in males and the second in females, with 1.8 million new cases and almost 861,000 deaths in 2018. But it is also singularly preventable. Its onset is preceded by an interval of approximately 10-15 years, during which benign lesions with different malignant potentials are present in the colon and can be effectively eliminated during screening colonoscopy [1][2][3][4]. Decades of screening colonoscopy data have provided us with a fairly reliable estimate of the cancer risk posed by conventional colorectal adenomas (cADNs), genuinely-dysplastic lesions that account for around 60% of precancerous colon tumors [5][6]. Less is known about the malignancy risk of the more recently defined "serrated" precancerous lesions, so called because of the saw-tooth-like epithelial infolding found in their crypt lumens [7][8]. The past 20 years have witnessed active efforts to characterize these lesions, endoscopically, histologically, and molecularly. Three types of serrated lesions are currently recognized: hyperplastic polyps (HPs), traditional serrated adenomas (TSAs), and sessile serrated lesions (SSL) (the term currently recommended by the World Health Organization for lesions previously referred to as sessile
serrated adenoma/polyps) [9][10][6][11]. HPs and TSAs arise mainly in the distal colon and rectum. HPs account for ~30% of all benign colorectal lesions, and their risk of transformation is considered to be very low. They are generally believed, however, to be precursors to at least some SSLs and TSAs. (The putative pathways underlying these progressions are discussed in a recent review [10].) The cancer risk of TSAs is probably similar to that of cADNs, but they represent only ~1% of precancerous colorectal lesions [10]. As for SSLs, which are usually found proximal to the splenic flexure, they represent about 10% of all precancerous colorectal tumors detected with high-performance colonoscopy [4], and yet they appear to give rise to almost 20% of all CRCs. The long-term risk of developing CRC after endoscopic removal of a large or “advanced” SSL (i.e., one measuring ≥10mm) is as high as that associated with removal of a similarly sized cADN [12][13].

Most CRCs thus arise from cADNs or SSLs. The latter are easier to miss during colonoscopy, partly because of their propensity for the proximal colon. Endoscopic visibility in this area is often reduced due to an insufficient bowel prep, and the examination is frequently marred by technical shortcomings (e.g., omission of cecal intubation, excessively rapid scope withdrawal times). Morphology also plays a role [7][14]. SSLs are nearly always flat or sessile lesions with indistinct borders and colors resembling that of the normal mucosa. They can also be obscured by a mucus cap. Studies conducted using same-day tandem examinations found that ~25% of all precancerous colon lesions are missed on colonoscopy [15]. The miss rate dropped to ~10% for all lesions measuring ≥10mm, but it remained high (~25%) for the lesions that were sessile or flat, the subset that includes SSLs. Most CRCs detected within 3 years of a negative colonoscopy derive from missed or incompletely-excised lesions, and these “interval” or “post-colonoscopy” malignancies are characterized by an over-representation of proximal-colon locations and cancers that develop along the serrated tumorigenic pathway [16][17][18][19].

Three-quarters of the CRCs arising in SSLs (10-15% of all CRCs) have a well-defined phenotype characterized by location in the proximal colon, the BRAFV600E gain-of-function mutation, and methylation of CpG islands cytosines that are generally unmethylated in the DNA of the normal colorectal mucosa [10][20][21]. Most of these CIMP ( CpG island methylator phenotype) cancers are also DNA mismatch repair (MMR)-deficient owing to methylation of the promoter CpG island of the MMR gene \textit{MLH1} (a well-known CIMP target) [22][20]. As a result, \textit{MLH1} expression is silenced, and MMR becomes deficient, as reflected by increased mutation rates and DNA microsatellite instability.

Precancerous lesions can be differentiated endoscopically to some extent, but the process is by no means simple. The location of the lesion within the colon is poorly informative. Conventional adenomas can arise anywhere in the colon. The serrated lesions display some degree of segmental preference (SSLs for the proximal colon, HPs and TSAs for the distal colon and rectum), but these preferences are by no means absolute. Lesion morphology is also of limited value. The Paris classification [23] distinguishes polyoid (i.e., stalked or sessile) precancerous colorectal lesions from those that are nonpolyoid (less elevated than a sessile lesion or flat or depressed relative to the surrounding non-lesional tissue). However, sessile lesions can be cADNs, SSLs, or HPs, whereas TSAs usually resemble cADNs in terms of their elevation above the mucosal plane. The “pit pattern” of a precancerous lesion, as defined during
high-magnification endoscopic examination of the crypt openings on its surface, can reliably predict the histologic diagnosis [24]. However, this approach is not used routinely in most gastroenterology centers, and the results still require histologic confirmation.

Histologic diagnosis itself is also far from being straightforward. Disagreement arises among pathologists regarding several types of resected precancerous colorectal lesions, and the issues underling this inter-examiner variability have been well-reviewed elsewhere [10] [6]. Suffice it to say here that the two most frequently encountered high-risk lesions, cADNs and SSLs, can be readily differentiated, since SSLs rarely exhibit dysplasia. Doubts can arise, however, with “advanced” lesions: SSLs (and TSAs) measuring ≥10 mm can harbor cytologic and architectural features of dysplasia. In this case, differential diagnosis with cADNs might be an issue, especially when the dysplasia resembles that typically found in cADNs [10][6][25][26]. Some lesions also display intratumoral heterogeneity, with certain areas resembling cADN and others more typical of SSL or TSA. TSAs are rare lesions with fairly typical histologic features consisting of eosinophilic cells with elongated nuclei, ectopic crypt foci, and slit-like serration. However, they can be misdiagnosed as cADNs, especially when they are large, polypoid and dysplastic, and located in the distal colon. Large, relatively flat TSAs in the proximal colon can also be mistaken for SSLs.

Problems can also arise in differentiating SSLs and HPs [10][6][27][28][29]. Both display crypt lumens with the characteristic saw-toothed pattern. But in HPs the serration does not extend to the base of the crypts, whereas in SSLs it involves the entire longitudinal axis of the gland, disrupting the simple tubular architecture with asymmetric dilatations that results in bizarre boot- or anchor-shaped crypts. Differential diagnosis is facilitated if the histologic section has been cut parallel to the longitudinal axis of the serrated crypts, but attention is rarely devoted to proper orientation of the specimens resected during endoscopy. Given the markedly different cancer risks associated with SSLs (high) and HPs (very low), an additional tissue staining procedure that would aid pathologists in reliably differentiating between these two types of serrated lesions seems desirable.

Driver gene mutations can also be somewhat informative for typing precancerous colorectal lesions. APC mutations, for example, are typical of cADNs, but these lesions (and in rare cases SSLs) can also harbor KRAS mutations, which are characteristic of TSAs. And while the BRAFV600E mutation is considered typical of SSLs, it can also be found in HPs [10]. As for TSAs, they are generally thought to progress along the KRAS-mutated molecular pathway, but progression also appears to occur along the BRAFV600E or KRAS/BRAF-wild type pathway [30]. Recent molecular studies have revealed that signaling pathways typically involved in colorectal tumorigenesis (e.g., Wnt signaling) have different pathogenic trajectories during the evolution of different CRC precursor types [31][32][33]. For example, somatic APC mutations lead to early constitutive activation of canonical Wnt signaling in cADNs, whereas aberrant Wnt signaling occurs later in SSL and TSA tumorigenesis and is triggered by epigenetic silencing (via CIMP) or genetic mutations affecting Wnt-signaling modulators or antagonists (e.g., SFRPs, AXIN2, RNF43, or RSPOs) [10] [34][35][36]. These differences are also reflected in the transcriptional outputs of the aberrant Wnt signaling, and characterization of the various gene expression profiles would therefore allow more precise classification of precancerous colorectal lesions.
Attempts have recently been made to differentiate CRC precursor lesions based on high-throughput transcriptome profiling data. RNA sequencing studies of serrated lesions and cADNs by our group [21] and that of Delker [37][38] have identified a large number of putative gene expression markers that could be used for this purpose, following their verification and validation with in situ hybridization (ISH) and immunohistochemistry. In the study described below, we used ISH to verify the expression profiles of 21 transcriptome-based RNA molecules that appeared to be promising markers for differentiation between CRC precursors. Such markers could be exploited to create simple tissue-staining tools for refining their routine histologic diagnosis. They could also help pinpoint the molecular pathways active in a given lesion, as a proxy for more complicated genetic and epigenetic analyses. Paradigmatic is the immunostaining for MLH1: negative results indicate that an SSL is dysplastic, almost invariably \( \text{BRAF} \)-mutated, CIMP-positive, and microsatellite instable. And close endoscopic follow-up is required after excision of this type of lesion [10].

**Methods**

**Tissues**

Formalin-fixed, paraffin-embedded colorectal tissues were obtained from the Zurich University Hospital Pathology Archives with local ethics committee approval (No. 2015-00185). Donors provided written consent to tissue testing and data publication. Samples were coded to protect donors’ rights to confidentiality and privacy. None of the donors had a family history of colorectal cancer. SSLs displayed no evidence of dysplasia and exhibited normal staining for the MLH1 protein. The routine histologic diagnosis was confirmed with a second evaluation by a gastrointestinal pathologist. Low magnification, H&E (hematoxylin and eosin) images of each colorectal lesion are shown in supplementary material.

**In-situ hybridization (ISH)**

Three-micrometer-thick tissue sections were mounted on SuperFrost Plus slides (Thermo Scientific, Reinach, Switzerland), stored at 4°C, and analyzed with ISH analysis within one month after sectioning. Deparaffinized sections were processed manually using the \textit{RNAscope 2.5 HD Red} reagents from Advanced Cell Diagnostics (presently Bio-Techne, Abingdon, United Kingdom), according to a protocol based on branched-DNA technology [39][40]. In brief, after blockade of endogenous peroxidase activity, epitope retrieval, and protease digestion steps, the sections were subjected to ISH with 20 pairs of primary oligonucleotide probes for each mRNA of interest. Each probe pair targeted two consecutive 20-to-30-nt regions at a given position within the transcript. (Hybridization of only three of the 20 pairs is sufficient to obtain a signal that can be detected with standard microscopy.) Each hybridized probe pair was then bound by a series of complementary amplification molecules and labeled probes containing a chromogenic enzyme, which markedly enhance signal detection sensitivity. High specificity is also ensured since the amplification cascade begins only after both members of the primary oligonucleotide probe pair have hybridized to the target transcript. The chromogenic reaction generates a single punctate
signal per RNA molecule. Dot size is generally proportional to the number of primary oligonucleotide probe pairs hybridized to the target RNA molecule, and dot aggregates indicate high concentrations of the target. Finally, nuclei were weakly stained with hematoxylin and the slides scanned with an Axio Scan.Z1 (Zeiss, Feldbach, Switzerland). Images shown in this report were obtained from these scannings using the ZEN 2 microscope software (Zeiss, Feldbach, Switzerland). As illustrated in Figure 1 and Table 1, the staining intensity was arbitrarily classified as low, moderate, high, or very high, while the staining distribution was reported both along the longitudinal axis of glands and across the whole section.

Results

We analyzed 12 premalignant colorectal tumor samples (3 cADNs; 3 SSLs; 3 HPs; and 3 TSAs) from the pathology archives of Zurich University Hospital (Table 1). All but one (HP 2) measured ≥10mm and were therefore considered “advanced” lesions.

Sections cut from each tumor block were processed for ISH. Each section was hybridized with primary oligonucleotide probes for one of the 21 RNA targets investigated (Supplementary Table 1) or one of the three RNAs used as staining controls (Supplementary Figure 1). In each section, staining of the normal mucosal crypts at the border of each lesion was assessed as an internal normal-tissue control. The choice of the RNA targets to be verified was based on RNA-sequencing data on precancerous colorectal lesions previously published by our group [21] (Supplementary Figure 2) and others [38] (Supplementary Figure 3).

ISH verification of RNA target expression patterns was undertaken to identify bona fide markers for distinguishing SSLs from cADNs. To this end, we focused our analysis on 12 mRNA targets (Supplementary Table 1). As shown in Table 1, SSLs and cADNs could be readily differentiated from one another using 9 of the 12 candidate markers (8 whose expression was SSL-specific [VSIG1, ANXA10, ACHE, SEMG1, AQP5, LINC00520, ZIC5, FOXD1] and 1 with cADN-specific expression [NKD1]). The ISH expression patterns of these mRNAs in SSLs and cADNs are shown in Figures 2, 3, and 5, and Supplementary Figures 4, 5, 6, 7, 8, 9, 10, 11, and 14. Two of the three remaining candidates (APOBEC1 and MUC5AC) were expressed more intensively in SSLs than in cADNs, but their expression was by no means SSL-specific (Figures 4 and 5, and Supplementary Figures 12 and 13). As for the third, the putatively SSL-specific marker KLK8 (Supplementary Table 1), its staining pattern was uninformative (results not shown), probably due to cross-hybridization with other KLK-family members.

The ISH data summarized in Table 1 also highlight the similarity between the expression patterns of the SSL-specific markers in SSLs and HPs (Supplementary Figures 4-11 and 14). These two serrated lesion types could not be distinguished even with the 9 RNAs that were chosen (on the basis of RNA sequencing data) to differentiate HPs from all other precancerous lesions [38] (e.g., HOXD13, HOXB13, and FAM3A [Supplementary Table 1, Table 1, Figure 6 and Supplementary Figures 15, 16 and 17]). HOXD13 and HOXB13, in fact, seem to be genuine markers not of HPs but of the normal mucosa in the distal colon and rectum, where HPs usually arise. These findings suggest that the specific expression of HOXD13 and
HOXB13 in the distal half of the colon (vs. the proximal colon) is retained in lesions typically occurring in this colorectal segment (Table 1). The upregulated expression of these two genes in HP biopsies processed for RNA sequencing might also have stemmed from normal mucosal contamination of the tumor sample (not uncommon with biopsy of lesions as small as most HPs). These interpretations are also consistent with the expression patterns of EVX2 (a HOXD13 neighbor), PRAC1 (a HOXB13 neighbor), INSL5, OR51E2, CPB1, and ST6GAL2 (Supplementary Table 1, Supplementary Figure 3, ISH data not shown). Along similar—albeit directionally opposite—lines, FAM3B, which was selected for verification because it was specifically unexpressed in HPs, proved instead to be a marker of the normal mucosa of the proximal-colon (HPs are generally located in the distal colon and rectum) (Figure 6, Supplementary Figure 17). Therefore, none of the nine RNAs that were putatively HP-specific could be verified as such.

As shown in Table 1 and all figures, the ISH expression patterns of TSAs were heterogeneous and frequently characterized by a mixture of SSL- and cADN-specific staining patterns.

H&E-stained images of all 12 lesions investigated in this study are shown in Supplementary Figures 18-29.

Discussion

RNA-based gene expression profiles generated by our group have revealed numerous RNA markers that are differentially expressed in SSLs and cADNs [21]. Here, using ISH, we verified the accuracy of 9 of the 12 markers putatively capable of distinguishing between these two major premalignant tumor types. Those that appeared to be SSL-specific, however, were unable to differentiate these lesions from HPs. Evidently, the gene-expression trajectories underlying the early stages of serrated tumorigenesis in these two serrated precursor lesions are common (see Introduction). HP-specific markers were also not found among additional 9 RNAs investigated in this study. The advanced TSAs (>10mm diameter) we investigated showed mixed staining patterns reflecting the coexistence in each lesion of serrated and cADN-like histologic features (Table 1).

One of the three genes that displayed particularly high expression in SSLs and HPs was VSIG1 (V-set and immunoglobulin domain containing 1) (Figure 2, Supplementary Figure 4). Absent in cADNs and the normal colon mucosa, VSIG1 expression was very high along almost the entire length of the serrated crypts in SSLs and HPs, except the very bottom of the crypt and its surface. TSAs, in contrast, display very little VSIG1 expression or none at all (e.g., TSA 2 in Table 1). The expression that is observed is reflected by patchy staining confined to glands with an SSL-like phenotype. The strikingly different VSIG1 expression patterns in serrated crypts (highly expressed) and those of the normal colorectal mucosa (unexpressed) might one day be exploited to improve detection of flat serrated lesions using fluorescein-labeled anti-VSIG1 antibodies during colonoscopy [41].

The VSIG1 protein, a member of the junctional adhesion molecule family, is normally expressed in the gastric mucosa and testis [42][43]. Its ectopic expression in serrated colorectal lesions, which has been documented at both the transcript and protein levels [37][44][21], is thought to reflect aberrant
differentiation toward a gastric-cell phenotype during the development of these tumors. SSLs and HPs also acquire expression of other molecules typically found in the gastric mucosa, e.g., ANXA10 (Annexin 10), a known marker of the normal mucosa of the stomach [45][46][47]. ANXA10 belongs to the calcium-dependent phospholipid-binding annexin protein family, and its function is currently unknown. Like VSIG1, ANXA10 is unexpressed in the normal colon mucosa and in most cADNs. (In rare cases, moderate expression can be observed in a few cells or crypts on the surface of cADNs.) ANXA10 can be considered a bona fide marker of serrated glands in SSLs and HPs (Figure 2 and Supplementary Figure 5), and it is also encountered fairly often in cells on the surface of TSAs and in their SSL-like glands.

The third gene that was highly expressed in SSLs and HPs, ACHE (acetylcholinesterase), is instead typically expressed in conducting tissues, including those of the enteric nervous system, and at neuromuscular junctions [48]. It terminates signal transmission by hydrolyzing the neurotransmitter acetylcholine at cholinergic synapses in the brain and at neuromuscular junctions, and pharmacologic inhibition of this enzymatic activity is used to treat colonic pseudo-obstruction [49]. We found high levels of ACHE mRNA in epithelial cells at the surface of the normal colorectal mucosa, but even higher levels were found in serrated glands, extending about half-way down toward the base of the crypts (Figure 2, Supplementary Figure 6). By contrast, ACHE expression is markedly lower in cADNs, although it may be found in some cells on the surface of adenomatous villi. In TSAs, some serrated glands are strongly positive for ACHE expression, and this feature might be used to better visualize the serrated component of these polyps. ACHE is also expressed in some stromal cells—probably lymphocytes—and in some cells of lymphocytic folliculi. As expected, it is also strongly expressed in the submucosal plexi (Supplementary Figure 6).

Like VSIG1, ANXA10 and ACHE, SEMG1 (Semenogelin 1) also emerged as a good marker of the serrated pathway of tumorigenesis [21][37][38]. It is not expressed in cADNs or in the normal colon mucosa, but moderate levels are found in SSLs and HPs, along the length of serrated crypts and to a somewhat lesser extent at the crypt bases and mouths (Figure 3 and Supplementary Figure 7). SEMG1 expression is also appreciable in regions of TSAs where the serrated glandular differentiation is more obvious. This gene, too, is involved in a curious form of dysregulated cell-fate differentiation that occurs during serrated tumorigenesis. SEMG1 (like SEMG2, which is also strongly expressed in some SSLs [21]), is typically expressed in seminal vesicles, and the SEMG1 protein is a major component of the semen coagulum [50]. Prostate specific antigen-mediated cleavage of SEMG1 yields functional polypeptides that favor semen liquefaction and enhanced sperm motility, and increased sperm levels of SEMG1 are often associated with asthenospermia [51]. It is tempting to hypothesize that an abundant ectopic secretion of semenogelins into the lumens of serrated colon crypts might favor the formation of a tenacious mucus matrix, which would explain the presence of the adhesive mucus cap that often covers SSLs [7].

AQP5 (Aquaporin 5) was also confirmed as a very good marker of serrated tumors: completely absent in the normal mucosa and in cADNs, AQP5 transcript is highly or very highly expressed at the bases of all serrated crypts in SSLs and HPs (Figure 3 and Supplementary Figure 8), and patchy, high-level expression
was also observed in two of the three TSAs we investigated (TSAs 1 and 3 in Table 1). There was no
evidence of AQP5 expression in TSA 2, which was characterized by a more pervasive dysplastic, cADN-
like histology than that seen in TSAs 1 and 3. Ectopic expression of this gene in serrated colorectal
glands is another example of tumor-associated, phenotypic dys-differentiation. AQP5 encodes a water-
channel membrane protein normally expressed in the bronchi, salivary glands, stomach, and testis [48].
AQP5 mutations and polymorphisms are associated with palmoplantar keratoderma [52] and with
outcomes in patients with acute respiratory distress syndrome [53]. Differentiation of alveolar epithelial
cells from type II to type I in the lungs is transcriptionally regulated by the p300/beta-catenin complex
(but not by the CREB-binding protein/beta-catenin complex), with a concomitant increase in the
expression of AQP5 [54]. This finding suggests a functional relationship between APQ5 and Wnt
signaling, but evidently not with the canonical Wnt signaling pathway, which is constitutionally active at
the base of normal colorectal crypts. The fact that AQP5 is unexpressed in the normal colorectal mucosa
and highly expressed at the bottom of serrated crypts (Figure 3 and Supplementary Figure 8) suggests
that a variant form of Wnt signaling, likely resembling that reported for alveolar epithelial cells, is active
during serrated tumorigenesis. Kleeman et al. [31] reported that the Wnt signaling activation observed in
CRCs arising through the serrated pathway is ligand-dependent, i.e., resulting from mutations in genes
encoding RNF43 or RSPOs proteins, which amplify Wnt signal transmembrane transduction. Increased
expression of AQP5 mRNA has also been demonstrated in MMR-deficient CRCs arising via the serrated
pathway [55], suggesting that such variant Wnt signaling might be upregulating the expression of this
gene in the bases of serrated crypts.

In contrast, Wnt signaling activation is ligand-independent in CRCs arising along the conventional
tumorigenic process acting in cADNs, i.e., tumors with mutations in APC or CTNNB1 genes encoding the
intracellular signal transduction proteins Adenomatous polyposis coli or beta-catenin, respectively. The
constitutive activation in cADNs of this canonical Wnt signaling at the base of normal colorectal crypts
upregulates the expression of well-known Wnt target genes, such as CMYC, CD44, NKD1 and AXIN2 [32]
[31]. NKD1 (naked cuticle homolog 1) encodes a protein that negatively regulates canonical Wnt signaling
via mechanisms that are still incompletely understood [56][57]. For these reasons, we tested NKD1 mRNA
expression in this study for its potential to distinguish cADNs (where high levels were expected) from
SSLs and HPs (where expression was very low and confined to a few cells at the bases of serrated
crypts) (Table 1, Figure 5 and Supplementary Figure 14).

Immunohistochemical staining patterns are sometimes difficult to interpret owing to the low specificity of
the available antibodies, and this limitation would have been highly relevant for many of the targets we
investigated in this study. The ISH protocol we used involves hybridization of multiple probes that are
complementary to the RNA targets, thereby providing highly specific results, and its sensitivity is also high
thanks to the use of a series of complementary amplification molecules (see Methods). Unlike
immunohistochemistry, ISH also allows visualization within the tissue of noncoding RNAs, such as
LINC00520, which we found to be upregulated in serrated lesions using RNA sequencing [21]. Our present
findings verify the validity of LINC00520 as a new marker of the serrated pathway: it is moderately to
highly expressed in the upper half of the serrated crypts in SSLs and HPs, virtually absent in cADNs,
except in a few cells at the mouth of the glands, and expressed at low to moderate levels at the surface of normal crypts (Table 1, Figure 3 and Supplementary Figure 9). Therefore, like ACHE, the LINC00520 gene is normally expressed in the superficial epithelium of normal colorectal crypts, and this expression is markedly upregulated in serrated lesions, where it extends deep into the abnormal crypts. This long noncoding RNA regulates endothelial nitric oxide synthase expression [58] and may play role in breast tumorigenesis [59], but its epigenetic regulatory function in the colorectal epithelium is completely unknown.

Two of the serrated-specific targets investigated in this study encode transcription factors, ZIC5 (Zinc finger protein of the cerebellum) and FOXD1 (Forkhead box D1). They are essential for embryonic development of specific tissues [60][61] but absent in most adult tissues, including the normal intestinal mucosa. Developmental transcription factors like these are often found to be ectopically re-expressed in specific tumor cells, and this is the case for ZIC5 and FOXD1 in serrated colorectal tumor cells (Table 1, Figure 4 and Supplementary Figures 10 and 11). Their mRNAs were consistently present in the serrated lesions we investigated: FOXD1 labeling was observed along the entire longitudinal axis of serrated crypts, whereas ZIC5 was generally confined to the lower half. ZIC5 and FOXD1 expression levels in serrated lesion were both low, probably because their mRNAs (like those of most transcription factors) are relatively unstable [62].

Interestingly, ZIC2 and its neighbor, ZIC5, displayed the same staining patterns in serrated lesions (Supplementary Figure 10). Expression of these two genes inhibits the transcriptional activity of beta-catenin/TCF (i.e., the canonical Wnt signaling that occurs in the adult stem cell compartment of the intestinal epithelium), thereby disrupting intestinal epithelial homeostasis [63]. Their re-expression during serrated tumorigenesis once again points to a switch from the canonical Wnt signaling active during conventional adenomatous tumorigenesis to a fundamentally different variant form of this signaling cascade, as previously discussed for AQP5 and NKD1. Indeed, ZIC5/2 expression has been reported in APC-wildtype and MMR-deficient colon cancer cell lines, but levels were almost undetectable in APC-mutant and MMR-proficient lines [63].

RNA-sequencing data [21] revealed SSL-specific upregulation of APOBEC1 (Apolipoprotein B mRNA editing catalytic subunit 1) and, as reported by others [37][44], MUC5AC (Mucin 5AC) expression (in comparison with normal mucosa). Topographical analysis of the ISH tissue-staining patterns confirmed the RNA-sequencing data, but it also highlighted a risk of error associated with conclusions based exclusively on this type of data. As shown in Table 1, Figures 4 and 5, and Supplementary Figures 12 and 13, neither APOBEC1 nor MUC5AC can be considered a bona fide marker of the serrated pathway: both are more strongly expressed in SSLs than they are in cADNs, but cADNs do consistently express both genes, albeit in more restricted areas of the glands or in a patchy pattern. (These staining patterns explain why random selection of endoscopic biopsies for RNA extraction and sequencing can lead to underestimated expression levels of certain genes in some tumor types.)
The RNA-editing enzyme APOBEC1, which deaminates apolipoprotein B mRNA Cytosine666>Uracil in the small intestine [64], is moderately expressed in the surface epithelium of the colorectal mucosa (Supplementary Figure 12). Its high-level expression in SSLs extends down into the serrated crypts but stops short of the crypt base. It remains to be seen whether apolipoprotein B editing and/or APOBEC1-mediated DNA mutagenesis (i.e., C>T transitions stemming from unrepaired cytosine deaminations) are increased in these neoplastic crypts [65][66]. It is interesting to note that C>T transitions at CpG dinucleotides are over-represented in the DNA mutation signature of CRCs displaying MMR-deficiency [67][68], which, as discussed above, is caused by CIMP-mediated silencing of MLH1 expression.

As for MUC5AC, its tumor-associated expression represents another example of dysregulated neoplastic cell-fate differentiation. MUC5AC encodes a typical gel-forming glycoprotein found in normal gastric and respiratory tract epithelial cells [69], and it proved to be an excellent marker of goblet cells in all the tumors we investigated (especially SSLs, which are typically goblet-cell-rich) but not of the goblet cells found in the normal colorectal mucosa (Supplementary Figure 13). The goblet-cell differentiation that occurs in serrated lesions (and in some areas of cADNs and TSAs) thus appears to be epigenetically distinct from that seen in normal colorectal crypts.

Whole-section analysis with ISH or immunohistochemistry facilitates characterization of expression pattern heterogeneity within a tumor, a feature that can be missed with RNA sequencing analysis of random biopsies, as exemplified by our experience with MUC5AC and APOBEC1. When used with reliable antibodies, immunohistochemistry can also identify tumor-specific changes that can escape detection by ISH. A recent example involves AGRN protein expression in the muscularis mucosae of SSLs, which has been shown to distinguish SSLs from HPs despite the fact that the two lesions display similar levels of AGRN mRNA expression levels in the lower half of their serrated crypts [70] (see also Supplementary Figure 2 and 3: AGRN mRNA expression patterns from [21] and [38], respectively).

Nine targets were chosen as putative HP-specific tissue staining markers [38] (RNA-sequencing data shown in Supplementary Figure 3), but none of the nine were verified as such by our ISH findings. Table 1, Figure 6 and Supplementary Figures 15 and 16 show two examples, the HOXD13 and HOXB13 genes. They belong to two different homeobox gene families of transcription factors that play crucial roles in vertebrate embryonic development [71][72]. Their expression in adult tissues is restricted to the distal colon and prostate (both genes), the vagina (HOXD13), and the urinary bladder (HOXB13) [48]. ISH confirmed that HOXD13 is expressed in the normal mucosa of the distal colon, the rectum in particular. It is also expressed at crypt bases in HPs, especially those located in the rectum (Table 1), but low expression is detectable also in the other tumor types investigated. Like HOXD13, HOXB13 expression is restricted to the distal colon and rectum (generally at higher levels than HOXD13), and it is expressed in HPs as well as all other tumor types, with levels in distal-colon tumors that are far higher than those in their proximal-colon counterparts. These two genes are more appropriately considered bona fide markers of the normal epithelium of the distal colon and rectum rather than of HPs. As discussed above, the fact that HPs are much more likely to arise in these segments than SSLs probably explains why these genes would appear to be HP-specific on the basis of RNA-sequencing data.
FAM3B, which encodes a signaling protein normally expressed in the endocrine pancreas and gastrointestinal tract [48][73], was chosen as a putative negative marker of HPs, i.e., one whose nonexpression is specific to these lesions (Supplementary Figure 3). This assumption was also explained by the staining pattern of the normal mucosa: unlike the previously discussed mRNAs, FAM3B is expressed only in the proximal segments of the normal colon, where HPs are rare. It was also variably expressed in the other lesion types from all colorectal segments, but the highest levels were found in tumor glands of proximal-colon lesions (Figure 6, Supplementary Figure 17).

The search for markers that can clearly distinguish SSLs from HPs is obviously going to be difficult. Kanth et al. [38] have identified additional candidate markers that were not included in our investigation. Using RT-PCR, they recently assessed the performance of a 7-marker panel that included five of those candidates, as well as SEMG1 and ZIC5/2. The panel differentiated SSLs from HPs with 89% sensitivity and 88% specificity [74]. However, the gene expression differences between these two types of serrated tumors were less significant when distal- rather than proximal-colon SSLs were considered, suggesting that distinguishing between serrated lesions arising in the same colon segment is still likely to be problematic.

One of the obvious limitations of our study is that it was conducted on an undeniably small number of colorectal lesions. Our aim here was to verify biomarker candidates on the basis of our previously reported RNA-sequencing data. Because the ISH protocol entails synthesis of branched DNA molecules, it is expensive, at least when done manually, as it was in this study, where the priority was to test a relatively large number of promising markers instead of evaluating a few markers in numerous lesions. However, this technique could easily be used with the automatic robotic instruments routinely used for immunohistochemistry in all pathology laboratories. This would reduce costs considerably and greatly facilitate next-step efforts to provide the reliable validation of the most promising markers in larger series of colorectal tumors representing all histologic types, sizes, and colorectal segments of origin.

Descriptive findings like ours can clearly have impact in the clinics, but they can (and should) also serve as springboards for research into the functional significance during serrated colorectal tumorigenesis of the dysregulated gene expression discussed above. The cascade of molecular events that characterizes this process appears to involve a dramatic epigenetic reprogramming, whose early stages are reflected by the recently described proto-CIMP phenotype [21] and intriguing forms of aberrant differentiation at the cellular and tissue levels.

**Abbreviations**

cADNs: Conventional adenomas; CRC: colorectal cancer; CIMP: CpG island methylator phenotype; H&E: Hematoxylin and eosin; HP: Hyperplastic polyp; ISH: in situ hybridization; MMR: DNA mismatch repair; SSL: Sessile serrated lesions; TSA: Traditional serrated adenoma.
Declarations

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Author contributions:

G.M. designed the study, performed in situ hybridization, and wrote the paper.

Declarations:

Ethics approval and consent to participate

Ethics approval was received from the Zürich university hospital ethics committee, Switzerland (Nr. 2015-00185). Donors provided written consent to tissue testing and data publication. Samples were coded to protect donors’ rights to confidentiality and privacy.

Consent for publication

Not applicable.

Availability of data and material

Not applicable.

Competing interests

The author declares that he has no competing interests.

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**Tables**

| Sample | histologic diagnosis | age of diagnosis | sex | lesion segment | lesion's diameter (mm) | JUNT | VASH 1 | ANXA 10 | PDXC 2 | HENX 1 | AQPS | HOK21 | FOSK 1 | MLCX 3 | LINC 2020 | ACRE | APO M8 | BE B1 | HXK 13 | MOKB 13 | KAIX 13 | FAM 3B |
|-------|----------------------|------------------|-----|----------------|-----------------------|------|--------|----------|---------|---------|------|--------|-------|--------|----------|--------|--------|-------|--------|--------|--------|--------|
| eADN 1 | TA with LGD | 76 | M | transverse | 12 x 7 |               |      |        |          |         |         |      |        |       |         |          |        |        |       |         |        |        |        |
| eADN 2 | TVA with LGD and normal HGD | 55 | M | rectum | 55 x 24 |               |      |        |          |         |         |      |        |       |         |          |        |        |       |         |        |        |        |
| eADN 3 | TVA with multifocal HGD | 68 | M | right fissure | 30 x 25 |               |      |        |          |         |         |      |        |       |         |          |        |        |       |         |        |        |        |
| SSL 1 | SSL | 46 | F | ascending | 15 x 10 |               |      |        |          |         |         |      |        |       |         |          |        |        |       |         |        |        |        |
| SSL 2 | SSL | 46 | F | right fissure | 13 x 7 |               |      |        |          |         |         |      |        |       |         |          |        |        |       |         |        |        |        |
| SSL 3 | SSL | 50 | M | ascending | 16 x 6 |               |      |        |          |         |         |      |        |       |         |          |        |        |       |         |        |        |        |
| HP 1 | HP | 78 | M | sigmoid | 16 x 8 |               |      |        |          |         |         |      |        |       |         |          |        |        |       |         |        |        |        |
| HP 2 | HP | 52 | M | transverse | 7 x 7 |               |      |        |          |         |         |      |        |       |         |          |        |        |       |         |        |        |        |
| HP 3 | HP | 60 | F | rectum | 16 x 6 |               |      |        |          |         |         |      |        |       |         |          |        |        |       |         |        |        |        |
| TSA 1 | TSA with dysplasia | 66 | F | cecum | 22 x 11 |               |      |        |          |         |         |      |        |       |         |          |        |        |       |         |        |        |        |
| TSA 2 | TSA with dysplasia | 63 | M | recto-sigmoid | 17 x 16 |               |      |        |          |         |         |      |        |       |         |          |        |        |       |         |        |        |        |
| TSA 3 | TSA (numerous SSA/P-like glands) | 90 | M | sigmoid | 18 x 12 |               |      |        |          |         |         |      |        |       |         |          |        |        |       |         |        |        |        |

Abbreviations: eADN, conventional adenoma; TA, tubular adenoma; TVA, tubulo-villous adenoma; LGD, low-high grade dysplasia; SSL, sessile serrated lesion; TSA, traditional serrated adenoma; F, female; M, male; P, proximal colon; D, distal colon and rectum.

**Full Resolution Figures**
Full resolution figures are available in the supplemental files section.

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