

## ARTICLE



# Low-grade non-intestinal-type sinonasal adenocarcinoma: a histologically distinctive but molecularly heterogeneous entity

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Although low-grade non-intestinal-type sinonasal adenocarcinoma (SNAC) is formally a diagnosis of exclusion defined by the absence of salivary or intestinal differentiation, most tumors in this category comprise a distinctive histologic group that are increasingly thought to derive from seromucinous glands. However, the molecular underpinnings of SNAC remain poorly understood, and it is unclear if diverse genetic alterations recently reported in isolated cases should delineate separate subgroups. This study aims to perform comprehensive evaluation of gene fusions and mutations and their histologic correlates in low-grade SNAC to clarify its pathogenesis and classification. We identified 18 non-intestinal-type SNAC that all displayed characteristic tubulopapillary architecture and low-grade cytology, although several cases had other unique histologic features and 3 showed intermixed high-grade areas. Among tumors stained with S100 protein, SOX10, and DOG1, 86% expressed at least one of these seromucinous markers. Of 17 cases with sufficient RNA or DNA available for analysis, likely oncogenic molecular alterations were identified in 76% of cases, most notably including *CTNNB1* p.S33F mutations in 2 cases, concomitant *BRAF* p.V600E and *AKT1* p.E17K mutations in 2 cases, and *ETV6::NTRK3*, *PRKAR1A::MET*, *FN1::NRG1*, and *DNAJB1::PRKACA* fusions in 1 case each. While tumors with most genetic alterations were histologically indistinguishable, cases with *CTNNB1* mutations had intermixed squamoid morules and cases with *BRAF* and *AKT1* mutations showed a myoepithelial cell population and prominent papillary to micropapillary architecture. Overall, these findings confirm previous reports of frequent seromucinous differentiation in low-grade SNAC. However, these tumors display striking molecular diversity with involvement of multiple kinase fusions, leading to frequent activation of signaling cascades including the MAPK pathway. While most genetic alterations are not associated with sufficiently distinctive histologic features to suggest separate classification, biphasic tumors with *BRAF* p.V600E mutations are more unique and may represent a distinctive subgroup.

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## INTRODUCTION

The sinonasal tract can give rise to a diverse range of gland-forming neoplasms. Those tumors that do not fit into defined salivary tumor types or show intestinal differentiation have traditionally been classified as non-intestinal-type sinonasal adenocarcinomas (SNAC)—a category that is further partitioned into low-grade and high-grade groups<sup>1,2</sup>. Although this terminology suggests that non-salivary, non-intestinal SNAC is a diagnosis of exclusion, the majority of tumors in this category actually represent a histologically distinctive group of low-grade adenocarcinomas that demonstrate a recognizable mix of tubular and papillary architecture. While these tumors were historically thought to be of surface origin<sup>3–5</sup>, their consistent tubulopapillary morphology and immunohistochemical expression of putative seromucinous markers S100 protein, SOX10, and/or DOG1 has supported increasing consensus that they actually arise from seromucinous glands<sup>6–9</sup>. As such, the name seromucinous adenocarcinoma has been proposed to describe this distinctive

group of SNAC<sup>7,8</sup>, although this terminology is not currently recommended by the World Health Organization (WHO) Classification of Head and Neck Tumours<sup>1</sup>.

Despite these well-defined histologic characteristics, the molecular underpinnings of low-grade SNAC are still not well understood, with somewhat piecemeal literature assessing for single alterations or documenting findings in individual cases. While *ETV6::NTRK3* and *ETV6::RET* have been identified in several low-grade SNAC, these cases represent <10% of all tumors evaluated<sup>10–12</sup>. A *SYN2::PPARG* fusion has also been reported in a single case of low-grade SNAC, and an *EGFR::ZNF267* fusion was identified in a case with overlapping features of seromucinous hamartoma and low-grade SNAC<sup>13,14</sup>. *CTNNB1* activating mutations and *BRAF* V600E mutations have also been documented in two cases each, with distinctive histologic elements<sup>15,16</sup>. Some of these molecular findings have been proposed to define unique subgroups of SNAC<sup>10–12,17</sup>. However, no study has evaluated both

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gene fusions and mutations in low-grade SNAC to establish their frequency or histologic correlates. This study aims to perform comprehensive histologic and molecular analysis of a large series of non-intestinal-type SNAC to better understand the classification and pathogenesis of these unique neoplasms.

## MATERIALS AND METHODS

### Case selection

We identified 18 cases of non-intestinal-type SNAC from the authors' consultation files and surgical pathology archives that had tissue available for molecular analysis. For inclusion in this study, tumors had to meet the 2017 WHO Classification of Head and Neck Tumours criteria for low-grade non-intestinal-type SNAC, including (1) no diagnostic features of any specific salivary tumor type or intestinal differentiation, (2) tubular and/or papillary architecture, (3) complex growth including back to back or cribriform glands with minimal intervening stroma, and (4) uniform nuclei<sup>1</sup>. As invasive growth is not part of this definition, degree of invasion was not considered as an inclusion criterion in this series. Tumors that harbored some higher grade components were included if recognizable low-grade areas that met the above criteria were also evident. All available hematoxylin and eosin (H&E) sections were reviewed for all cases, and histologic features were tabulated. Because the majority of cases were seen in consultation, detailed clinical and follow-up information was not available.

### Molecular analysis

RNA sequencing (RNA-Seq) was attempted on all 18 cases and DNA next-generation sequencing (NGS) was attempted on 15 cases as described in detail elsewhere<sup>18</sup>. In brief, both RNA and DNA were isolated from 10 µm whole-slide tissue sections using Qiagen AllPrep kits (Qiagen, Germantown, MD). A modified TruSight RNA Pan-Cancer kit (Illumina, San Diego, CA) was used to make a sequencing library containing all exons from 1425 cancer-related genes. Sequencing was performed on the NextSeq 550 (Illumina) with a minimum of 6,000,000 mapped reads. All fusions and variants were reviewed in the Integrated Genomics Viewer (Broad Institute, Cambridge, MA). The Star-Fusion algorithm was used to call fusions, and somatic variants were identified using databases including dbSNP and gnomAD.

### Immunohistochemistry

The results of existing immunohistochemical stains were tabulated for each case. Antibodies used in the majority of cases included AE1/AE3 (clone pck-

26; prediluted; Ventana Medical Systems, Tucson, AZ), CK7 (clone ov-tl; 1:500; Dako, Carpinteria, CA), CK20 (clone Ks20.8; prediluted; Dako), CDX2 (clone EPR2764Y; prediluted; Dako), S100 protein (clone 4C4.9; prediluted; Ventana Medical Systems), SOX10 (clone N-20; prediluted; BioCare Medical, Pacheco, CA), p63 (clone 4a4; prediluted; BioCare Medical), p40 (clone BC28; 1:100; BioCare Medical), DOG1 (clone SP31; prediluted; CellMarque/Sigma-Aldrich, St. Louis, MO), smooth muscle actin (SMA; clone 1A4; prediluted; Ventana Medical Systems), calponin (clone M3556; 1:500; Dako), and beta-catenin (clone 14; 1:1000; BD Biosciences, Franklin Lakes, NJ). In most cases, staining was performed using standard protocols on Ventana BenchMark Ultra autostainers (Ventana Medical Systems) in the presence of appropriate controls, and signals were visualized using the ultraView polymer detection kit (Ventana Medical Systems). In cases where few or no stains were performed at the time of diagnosis, tissue was prioritized for molecular testing over additional immunohistochemistry.

## RESULTS

### Histologic findings

The low-grade SNAC all arose in the nasal cavities of 10 women and 8 men with a median age of 56 years (range 17–86 years). All tumors were composed of crowded, back to back, and confluent tubules, with variable areas of papillary architecture, composed of cuboidal to polygonal cells with a moderate amount of pale eosinophilic to amphophilic cytoplasm. However, a subset of cases had more unique histologic features intermixed. Three tumors demonstrated a diffuse oncocytic appearance, 2 had scattered squamoid morules, and 1 included a prominent population of goblet cells. Although most tumors appeared to be composed of only one cell type, 2 cases had distinct biphasic populations of basal and luminal cells evident on H&E sections. While all cases had areas of low-grade cytology, 1 case had a markedly elevated mitotic rate and 2 had zones of increased cytologic atypia and necrosis, 1 of which demonstrated overt sarcomatoid transformation. One case also arose in association with a respiratory epithelial adenomatoid hamartoma (REAH) but had areas of crowded, back-to-back tubules diagnostic of adenocarcinoma.

### Immunohistochemical findings

Immunohistochemical findings are tabulated in Table 1. All cases tested were positive for AE1/AE3 ( $n = 8$ ) and CK7 ( $n = 13$ ) and

**Table 1.** Immunohistochemical findings.

Case	Age	Sex	AE1/AE3	CK7	CK20	CDX2	S100 protein	SOX10	DOG1	p63/p40	SMA	Calponin	Beta-catenin
1	52	F	NA	+	–	–	NA	NA	NA	–	NA	NA	NA
2	55	M	+	+	–	–	–	+	+	–	NA	NA	NA
3	75	M	+	+	–	NA	+	–	+	–	NA	NA	NA
4	59	F	NA	+	–	NA	NA	NA	NA	NA	NA	NA	NA
5	36	F	+	NA	NA	NA	+	+	NA	–	–	–	NA
6	17	F	NA	NA	NA	NA	+	+	+	NA	NA	NA	Nuclear
7	86	M	Ductal	Ductal	–	–	+	NA	NA	Basal	Basal	NA	NA
8	63	F	Ductal	Ductal	NA	NA	+	+	NA	Basal	NA	Basal	NA
9	38	F	NA	NA	NA	NA	–	NA	NA	Basal	NA	–	NA
10	57	M	NA	+	–	NA	–	–	NA	Basal	–	–	NA
11	64	F	+	+	NA	NA	–	–	–	–	NA	NA	NA
12	51	M	NA	+	–	–	–	+	+	–	–	–	NA
13	57	M	+	+	–	–	–	+	+	–	NA	NA	NA
14	71	F	NA	+	NA	–	+	+	NA	–	NA	NA	NA
15	28	M	+	+	–	–	–	NA	–	NA	NA	NA	NA
16	49	F	NA	NA	NA	NA	+	–	+	NA	NA	NA	NA
17	31	F	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
18	71	M	NA	+	–	NA	NA	NA	NA	NA	NA	NA	NA

NA not attempted. +: positive, -: negative.

**Table 2.** Molecular findings.

Case	RNA-Seq	DNA NGS	Activated Pathways
1	<i>FN1::NRG1</i>	NA	MAPK, PI3K
2	<i>PRKAR1A::MET</i>	NA	MAPK, PI3K
3	<i>ETV6::NTRK3</i>	NA	MAPK, PI3K
4	<i>DNAJB1::PRKACA</i>	–	cAMP
5	–	<i>CTNNB1</i> p.S33F	Wnt
6	–	<i>CTNNB1</i> p.S33F	Wnt
7	Failed	<i>BRAF</i> p.V600E, <i>AKT1</i> p.E17K	MAPK, PI3K
8	–	<i>BRAF</i> p.V600E, <i>AKT1</i> p.E17K	MAPK, PI3K
9	–	–	NA
10	–	–	NA
11	–	<i>CDKN2A</i> homozygous loss	Cell cycle
12	–	<i>TP53</i> p.K132M	Cell cycle
13	–	<i>NF1</i> p.S1262fs and c.888 + 1 G > T	MAPK
14	–	<i>CUL3</i> p.V427fs	Proteolysis
15	–	<i>RAF1</i> amplification	MAPK
16	–	–	NA
17	Failed	–	NA
18	Failed	Failed	NA

NA not attempted. –: negative.

negative for CK20 ( $n = 10$ ) and CDX2 ( $n = 7$ ). There were also 7 cases positive for S100 protein (50%), 7 cases positive for SOX10 (64%), and 6 cases positive for DOG1 (75%). Of 7 cases that had S100 protein, SOX10, and DOG1 staining all performed, 6 (86%) were positive for at least one of these markers; the 1 tumor negative for all 3 stains had an oncocyctic appearance. The 2 cases with clear biphasic cell populations on H&E sections demonstrated positivity for p63 or p40, SMA, and calponin in a basal distribution, with S100 protein and SOX10 expression in both basal and luminal cells. Two additional cases that lacked a clear basal cell layer on H&E also displayed peripheral expression of p40 but were negative for SMA and calponin. The single tumor tested had diffuse nuclear and cytoplasmic beta-catenin positivity.

### Molecular findings

Results of molecular analysis are summarized in Table 2. Among the 18 low-grade SNAC, RNA was successfully amplified for analysis in 15 cases. Four of these cases (27%) demonstrated gene fusions including *ETV6::NTRK3*, *PRKAR1A::MET*, *FN1::NRG1*, and *DNAJB1::PRKACA* in 1 case each. DNA was also successfully amplified in 14 cases, of which putative oncogenic mutations were identified in 9 (64%), encompassing 2 cases with concomitant *BRAF* p.V600E and *AKT1* p.E17K mutations, 2 cases with activating *CTNNB1* p.S33F mutations, 1 case with biallelic *NF1* p.S1262fs and c.888 + 1 G > T mutations, 1 case with *TP53* p.K132M mutation, 1 case with *CUL3* p.V427fs mutation, 1 case with *RAF1* amplification, and 1 case with homozygous *CDKN2A* loss. The remaining cases had only variants of uncertain significance. Overall, of 17 cases with either RNA or DNA available for analysis, 13 cases (76%) harbored a likely oncogenic fusion or driver mutation. Pathways known to be activated by these gene alterations are also noted in Table 2, with 7 of the 13 alterations (54%) leading to MAPK pathway activation.

### Tumors with gene fusions

All 4 SNAC with gene fusions were almost entirely composed of crowded to confluent tubules with a moderate amount of eosinophilic to amphophilic cytoplasm and rare poorly-formed papillary excrescences. This pattern was exemplified by the case with *DNAJB1::PRKACA* fusion, which had no other exceptional features (Fig. 1A). While all four of these tumors largely had monotonous, centrally-placed, round to oval nuclei, the case with *ETV6::NTRK3* fusion harbored areas of markedly increased mitotic activity (Fig. 1B). The case with *FN1::NRG1* fusion arose in association with REAH, which showed prominent invaginations of surface epithelium and basement membrane material amongst less crowded tubules (Fig. 1C) but transitioned to more expansile areas of confluent growth diagnostic of adenocarcinoma (Fig. 1D). The most unusual morphology was seen in the case with *PRKAR1A::MET* fusion, which demonstrated a dominant population of goblet cells with well-defined intracytoplasmic mucin vacuoles and more abundant papillary structures (Fig. 1E) intermixed with focal areas of more conventional serous tubules (Fig. 1F). Notably, despite the goblet cells, this tumor was negative for CK20 and CDX2.

### Tumors with *CTNNB1* mutations

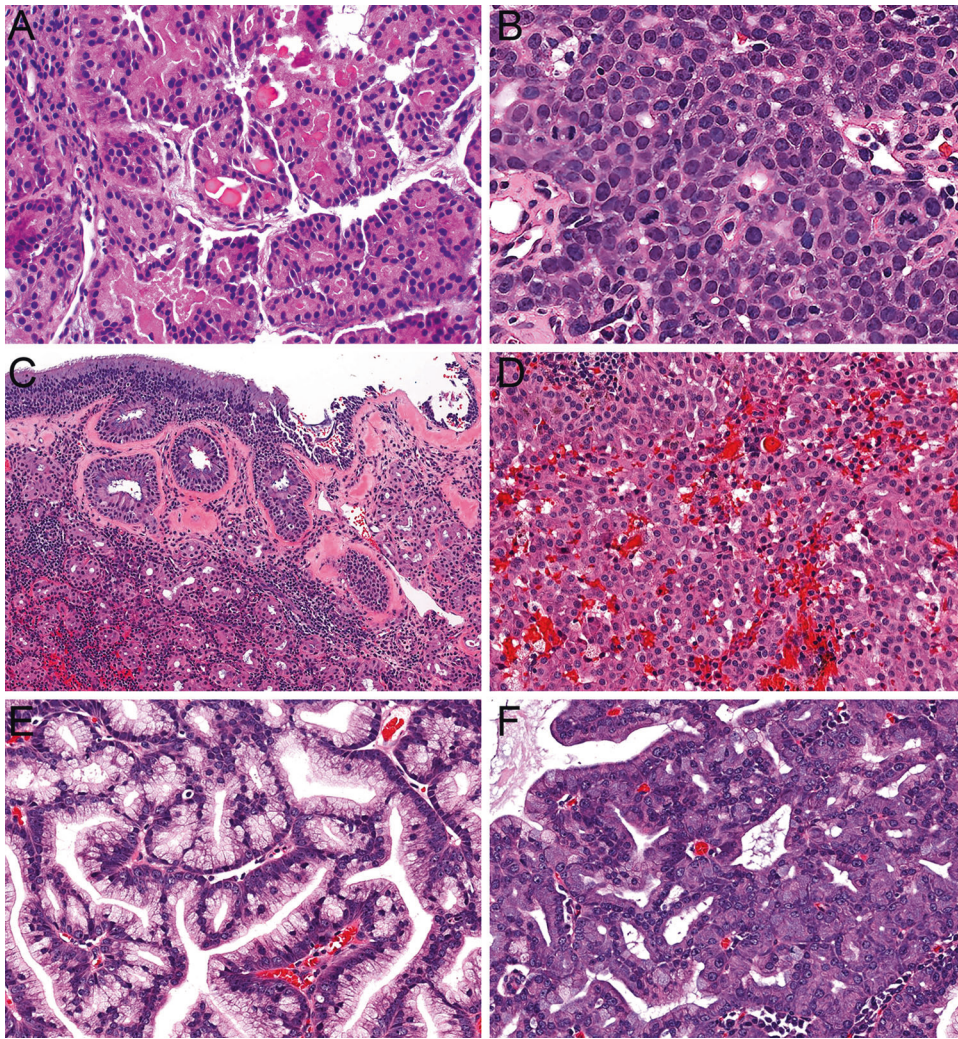
Both tumors that demonstrated activating p.S33F point mutations in the *CTNNB1* gene were uniformly composed of crowded tubules with pale eosinophilic cytoplasm and round nuclei similar to other cases. However, both of these tumors also displayed scattered squamoid morules that ranged from small and focal in one case (Fig. 2A) to large and prominent in 1 case (Fig. 2B). Beta-catenin immunohistochemistry was available for one case and demonstrated diffuse nuclear and cytoplasmic positivity (Fig. 2C). S100 protein and SOX10 were also positive in both tumors.

### Tumors with *BRAF* and *AKT1* mutations

The 2 tumors with concomitant *BRAF* p.V600E and *AKT1* p.E17K mutations both had biphasic cell populations evident on H&E. The peripheral layer was composed of cuboidal cells with scant clear to pale eosinophilic cytoplasm and round nuclei (Fig. 3A). The central layer displayed larger cells with highly variable architecture, including both crowded tubules with areas of oncocyctic change that were indistinguishable from most monophasic tumors (Fig. 3B) as well as complex micropapillary excrescences (Fig. 3C) and papillary fronds (Fig. 3D). By immunohistochemistry, the peripheral cells demonstrated a well-developed myoepithelial phenotype with positivity for p63 or p40 (Fig. 3E), S100 protein, SOX10, SMA, and calponin (Fig. 3F). The luminal cells were positive for AE1/AE3 and CK7 as well as S100 protein and SOX10. Notably, both cell populations were intimately intermixed and showed complex confluent growth throughout the tumor, suggesting that the tumor truly had biphasic elements and was not just colonizing existing myoepithelial cells. Interestingly, despite clear low-grade areas, both of these tumors also had high grade components, including nuclear enlargement, increased pleomorphism, and patchy tumor necrosis, and 1 tumor displayed overt sarcomatoid transformation (Fig. 3G) with areas of osteoid production.

### Tumors with other molecular alterations

The remaining cases harbored various molecular findings in association with two discrete histologic patterns. Seven tumors displayed classic crowded tubules with a moderate amount of amphophilic to pale eosinophilic cytoplasm and round nuclei. This group included tumors with biallelic *NF1* p.S1262fs and c.888 + 1 G > T mutations, *TP53* p.K132M mutation, *CUL3* p.V427fs mutation (Fig. 4A), *RAF1* amplification (Fig. 4B), as well as 3 cases with no identifiable genetic alterations or which failed nucleic acid amplification (Fig. 4C). There were also 3 tumors that had an oncocyctic appearance composed of back-to-back tubules and trabeculae of larger cells with abundant eosinophilic granular



**Fig. 1 SNAC with gene fusions.** The tumor with *DNAJB1::PRKACA* fusion exemplified the crowded tubules of SNAC (A; case 4). The case with *ETV6::NTRK3* fusion was notable for a markedly elevated mitotic rate despite otherwise low-grade cytology (B; case 3). Areas of invaginated surface epithelium lined by dense basement membrane material diagnostic of REAH (C; case 1) were adjacent to more confluent glands in a SNAC with *FN1::NRG1* fusion (D; case 1). The tumor with *PRKAR1A::MET* fusion showed prominent goblet cells with well-defined intracytoplasmic mucin vacuoles (E; case 2) intermixed with more conventional serous tubules (F; case 2).

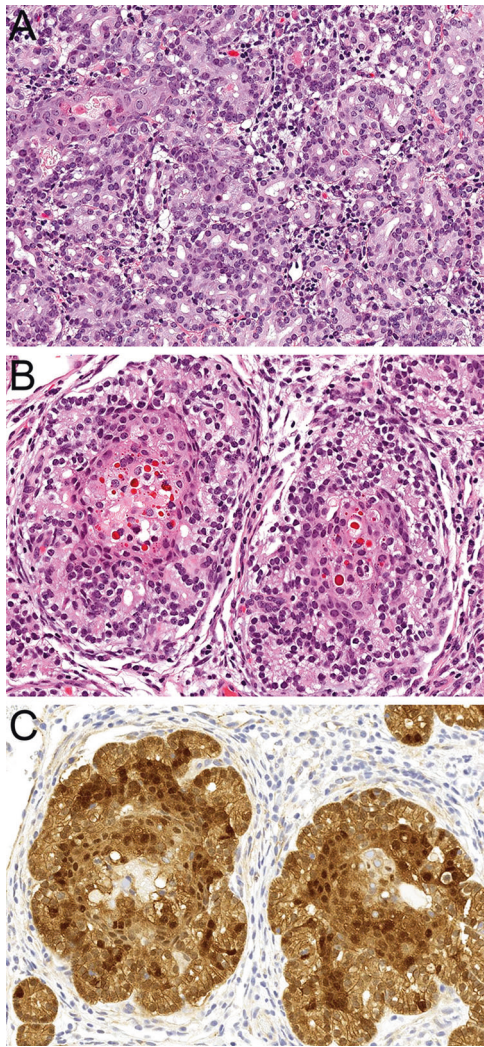
cytoplasm and prominent nucleoli. One of these tumors had homozygous *CDKN2A* loss and was composed of a monophasic population of confluent tubules (Fig. 4D). Interestingly, 2 of these tumors (Fig. 4E), both of which lacked identifiable mutations or fusions, had some degree of biphasic differentiation. Although not well-visualized on H&E sections, they harbored peripheral cells that were positive for p40 (Fig. 4F) and negative for other myoepithelial markers calponin and SMA as well as S100 and SOX10.

## DISCUSSION

Low-grade non-intestinal-type SNAC is formally defined as a diagnosis of exclusion that encompasses any cytologically bland, gland-forming sinonasal tumor that lacks salivary or intestinal differentiation. In practice, however, almost all SNAC that fall into this category comprise a discrete group of tumors with recurrent tubulopapillary architecture that are increasingly regarded to be of seromucinous gland origin. Despite this well-defined histologic profile, the molecular underpinnings of low-grade SNAC are not well-understood. Recent reports of several different gene fusions and mutations in small numbers of cases have raised questions as

to whether further molecular subtypes of SNAC should be defined. This study performed comprehensive RNA and DNA sequencing on a large series of low-grade non-intestinal-type SNAC to better understand their classification and pathogenesis.

First, our findings confirm previous reports of frequent seromucinous differentiation in low-grade SNAC, both at a histologic and immunohistochemical level. Despite some variable elements, all SNAC included in this series displayed histologic seromucinous features in the form of consistent tubulopapillary architecture, which traditionally has been regarded as analogous to the terminal tubules of the seromucinous glands<sup>6</sup>. While a subset of cases had oncocyctic, mucinous, or even myoepithelial differentiation, these elements may actually recapitulate other cell types that are also normal constituents of seromucinous glands. Although comprehensive immunohistochemical analysis was not possible because we prioritized molecular testing in cases with limited tissue, 86% of SNAC that underwent staining for S100 protein, SOX10, and DOG1 expressed at least 1 of these putative seromucinous markers. These results are in keeping with previous reports of a recurrent seromucinous immunophenotype in low-grade SNAC<sup>7</sup>. Furthermore, 3 cases that showed seromucinous differentiation but harbored both low-grade and high-



**Fig. 2 SNAC with *CTNNB1* mutations.** The tumors with *CTNNB1* activating mutations both were composed of crowded tubules with pale eosinophilic cytoplasm and round nuclei, with squamoid morules that ranged from small and focal (**A**; case 5) to large and prominent (**B**; case 6). The one case tested displayed diffuse nuclear and cytoplasmic beta-catenin positivity (**C**; case 6).

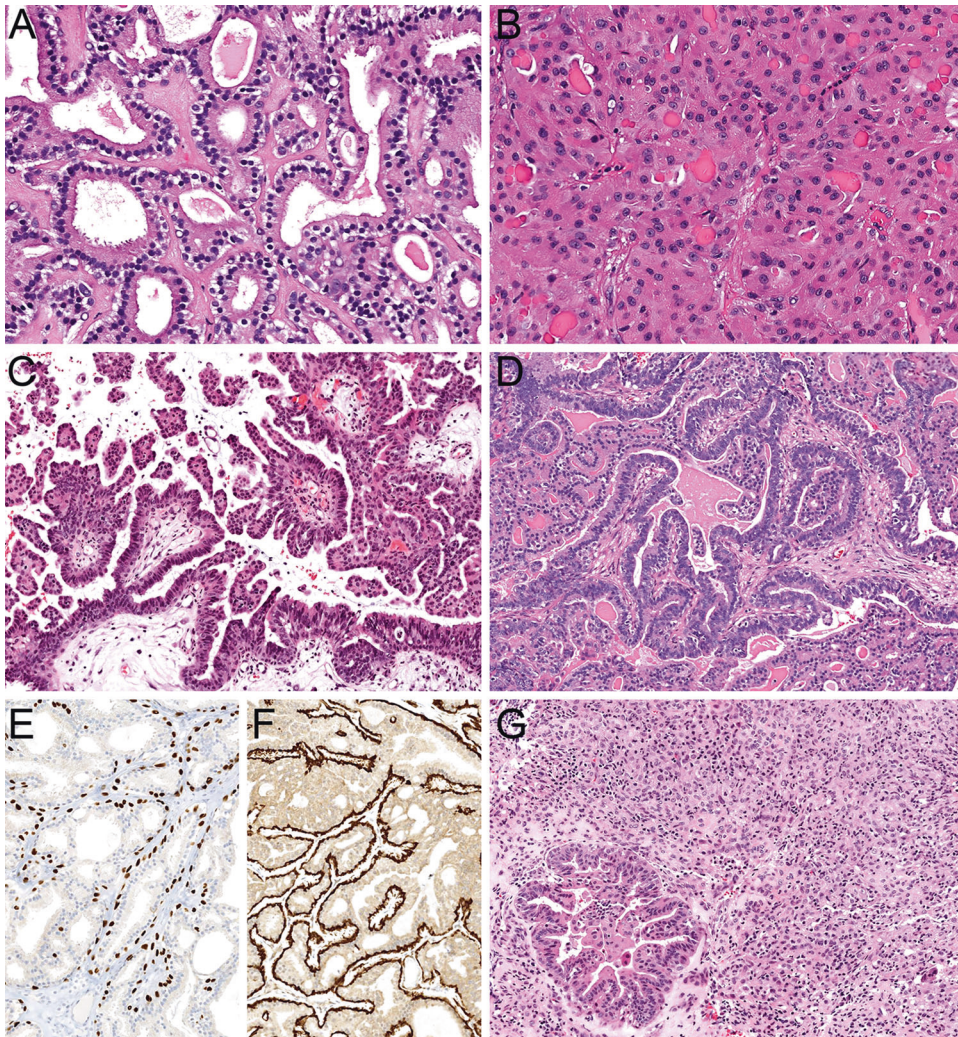
grade areas add to existing evidence that the seromucinous phenotype may transcend conventional grade-based grouping of non-intestinal-type SNAC<sup>7,19</sup>. These histological and immunohistochemical findings support previous proposals for reclassifying low-grade non-intestinal-type SNAC as seromucinous-type adenocarcinoma<sup>7,8</sup>. Paradoxically, despite their historical designation as “non-salivary,” SNAC with seromucinous features may even be conceptualized as specialized salivary tumors that exclusively arise in the sinonasal tract. However, it is essential to note that seromucinous differentiation is not specific for malignancy in the sinonasal tract, as it is also seen in seromucinous hamartomas and REAH. Both of these benign lesions not only can be difficult to distinguish from SNAC in limited samples, but also are occasionally seen adjacent to low-grade SNAC and may represent precursor lesions<sup>13,20</sup>. While one SNAC in this series was indeed associated with a REAH, all SNAC included here clearly displayed the crowded, back-to-back glands that define the presence of adenocarcinoma according to the 2017 WHO Classification of Head and Neck Tumours.

Our findings also highlight marked genetic heterogeneity in these low-grade SNAC, including a prominent role for diverse

kinase fusions. The SNAC in this series harbored 11 different oncogenic drivers, only 2 of which (*CTNNB1* and *BRAF/AKT1* mutations) were found in more than 1 case. Notably, 4 separate kinase fusions were present, accounting for almost a third of tumors. Of course, *ETV6::NTRK3*, as well as *ETV6::RET*, were previously identified in up to 10% of low-grade SNAC<sup>10–12</sup>. However, the other 3 fusions seen in this cohort have never been reported in this tumor type. *DNAJB1::PRKACA* was originally described as the defining molecular alteration in fibrolamellar hepatocellular carcinoma and was subsequently identified in intraductal oncocytic papillary neoplasm of the pancreas<sup>21–24</sup>. *FN1::NRG1* and *PRKAR1A::MET* appear to be entirely novel fusions, although both *NRG1* and *MET* are rarely rearranged with various partners in multiple tumor types, most commonly lung adenocarcinoma<sup>25–28</sup>. In combination with previous reports of *SYN2::PPARG* and *EGFR::ZNF267* fusions in single cases<sup>13,14</sup>, these findings suggest that many different kinase fusions may serve as drivers for a significant subset of SNAC. The majority of SNAC that lacked fusions showed alterations in other common oncogenic drivers including *CTNNB1*, *BRAF*, *AKT1*, *TP53*, *CDKN2A*, *NF1*, and *RAF1*. While *CUL3* is a less well-established driver, it has been reported as a tumor suppressor gene in esophageal squamous cell carcinoma and papillary renal cell carcinoma, where loss of function mutations are thought to contribute to oncogenesis via dysregulation of transcription factor NRF2<sup>29,30</sup>. As such, we regard the loss of function frameshift mutation seen in 1 SNAC as potentially oncogenic as well. Because several of these fusions, *BRAF* p.V600E mutations, and *RAF1* amplification represent drug targets, comprehensive molecular analysis could point to treatment options in rare SNAC that demonstrate aggressive behavior. Interestingly, most of these alterations also activate cellular signaling cascades, including the MAPK pathway in 54% of cases, suggesting some degree of unified pathogenesis despite this genetic variability. Regardless, the degree of heterogeneity at the molecular level indicates that a strategy relying on any single alteration for diagnosis of SNAC is impractical.

Given this diverse genetic landscape, we also question whether any single genetic alteration is sufficient to delineate subtypes of low-grade SNAC within the unified histologic spectrum. Specific gene fusions and mutations are increasingly regarded as pathognomonic for sinonasal and salivary gland entities. And indeed, *ETV6::NTRK3* and *ETV6::RET* fusions have been proposed to define a separate category of “*ETV6*-rearranged low-grade SNAC”<sup>10–12</sup>. However, we find it impossible to objectively distinguish SNAC with *ETV6::NTRK3* fusions, either in this series or published literature, from most other cases at a histologic level. In a background of multiple molecular alterations, it is difficult to confirm any fusion-defined subgroup of SNAC as unique without a correspondingly distinctive histologic profile. SNAC with *CTNNB1* mutations have also been suggested to represent a separate group<sup>16,17</sup>. These tumors are more identifiable histologically with consistent formation of squamoid morules—a feature that was recognized as unique before their molecular underpinnings were documented<sup>7</sup> and is common to other tumor types with beta-catenin alterations<sup>31–35</sup>. But they also display crowded tubules identical to other seromucinous-type SNAC. Moreover, squamoid morules have also been reported in the SNAC with *SYN2::PPARG* fusion<sup>14</sup>, casting doubt on the specificity of this finding for *CTNNB1* mutations. Undoubtedly, detailed clinicopathologic and molecular analysis of a still-larger group of SNAC will be necessary to better establish whether subgroups can and should be carved out of this category. At this point, we favor that low-grade SNAC are more analogous to soft tissue or thyroid tumors, in which multiple genetic alterations produce similar histologic findings<sup>36,37</sup>, than molecularly-defined sinonasal entities.

Nevertheless, these findings do highlight unique cases of SNAC with well-developed myoepithelial elements, prominent papillary and micropapillary architecture, and recurrent *BRAF* p.V600E and

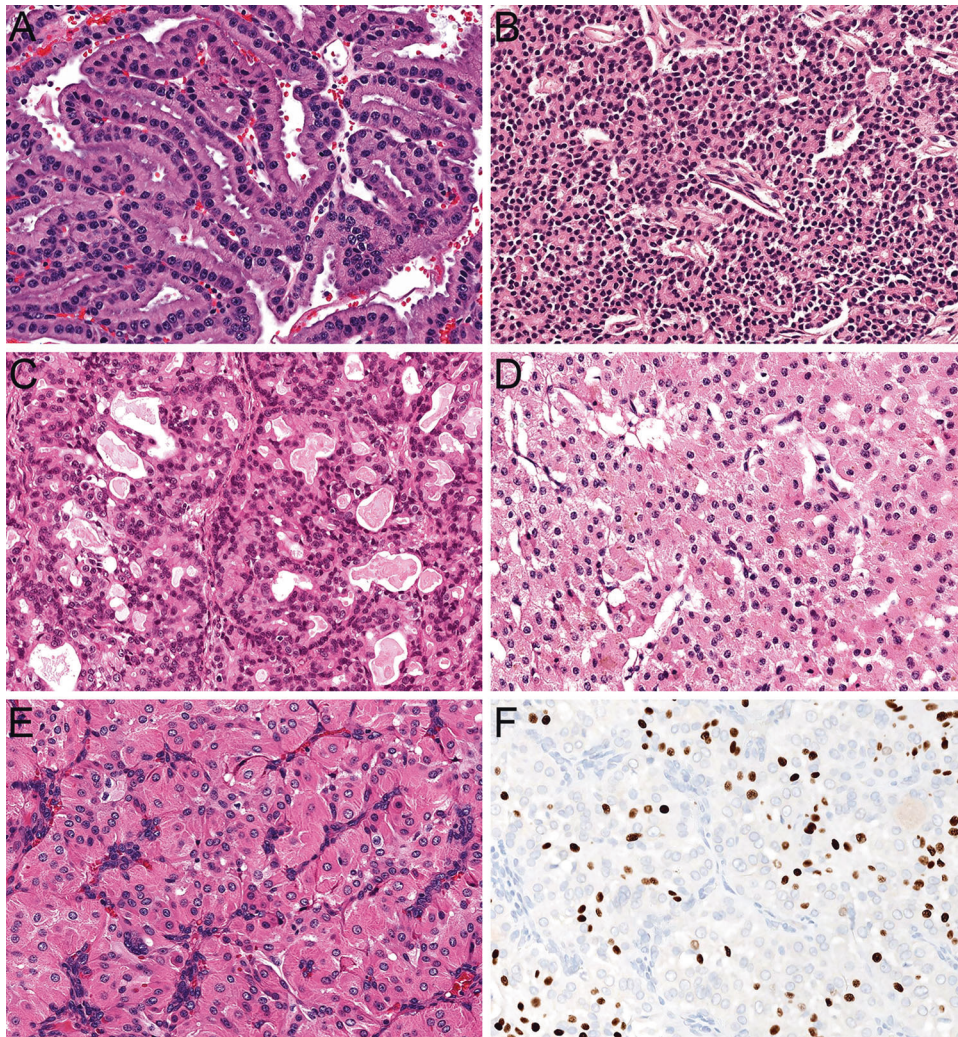


**Fig. 3 SNAC with *BRAF* p.V600E mutations.** Two tumors with concomitant *BRAF* p.V600E and *AKT1* p.E17K mutations demonstrated biphasic populations of peripheral cuboidal cells with clear to pale eosinophilic cytoplasm (**A**; case 8) and central ductal cells that showed variable tubular (**B**; case 7), micropapillary (**C**; case 8), and papillary (**D**; case 7) architecture. The peripheral cells showed a true myoepithelial phenotype with positivity for both p40 (**E**; case 7) and calponin (**F**; case 7). Despite largely low-grade histology, both of these cases also showed high grade areas including overt sarcomatoid transformation in one case (**G**; case 8).

*AKT1* p.E17K mutations that may represent a more distinctive category. In addition to the 2 cases in our series, 4 very similar low-grade SNAC with biphasic cell populations and papillary growth patterns were previously reported<sup>20,38</sup>, of which 2 were also found to have *BRAF* p.V600E mutations<sup>15</sup>, suggesting that these tumors are a recurrent phenomenon. Even in a relatively small cohort, these biphasic, *BRAF*-mutant SNAC have a sufficiently unique histologic and molecular profile to suggest they may represent a distinctive subgroup. The myoepithelial differentiation in these tumors is particularly exceptional among SNAC. Only 1 other SNAC in the literature<sup>7</sup> and 2 cases with oncocyctic features in this series showed peripheral expression of p63 or p40 alone- a finding that likely reflects basal differentiation typical in oncocyctic tumors rather than true myoepithelial elements<sup>39</sup>. The high-grade features in both cases with *BRAF* p.V600E mutation in this cohort, including overt sarcomatoid transformation in one case, are also notable. Of course, these unique characteristics call into question whether these *BRAF*-mutant tumors are really SNAC at all. The biphasic cell populations in these tumors have repeatedly raised the possibility that they should be classified as salivary epithelial-myoeplithelial carcinoma (EMC)<sup>20,38</sup>. However, EMC is vanishingly rare in the sinonasal tract, generally lacks papillary and

micropapillary architecture, and does not harbor recurrent *BRAF* mutations<sup>40–42</sup>, arguing against this diagnosis. These tumors also bear some resemblance to salivary sialadenoma papilliferum and oncocyctic variant of intraductal carcinoma, which share biphasic populations and *BRAF* p.V600E mutations<sup>43–46</sup>. But they lack the surface squamous papillations of sialadenoma papilliferum and demonstrate more architectural diversity than oncocyctic variant of intraductal carcinoma. Without a comfortable fit into existing salivary tumor types, these biphasic, *BRAF*-mutant tumors are probably best regarded as a distinctive group within the SNAC spectrum until more comprehensive analysis can assess whether they represent a unique entity.

In summary, this study reinforces emerging consensus that low-grade non-intestinal-type SNAC is a largely unified histologic group and confirms previous reports of seromucinous differentiation in a majority of cases. Comprehensive molecular analysis also confirms that these low-grade SNAC are a molecularly heterogeneous category, with a particularly prominent role for diverse kinase fusions, although many of these variable alterations may lead to similar downstream signaling amplification including MAPK pathway activation. Tumors with most fusions or mutations did not display sufficiently distinctive histologic features to support



**Fig. 4 SNAC with miscellaneous genetic alterations.** Classic tubulopapillary architecture with crowded, cytologically bland glands were seen in SNAC with *CUL3* mutation (**A**; case 14), *RAF1* amplification (**B**; case 15), and no detectable genetic alterations (**C**; case 16). Oncocytic change was also seen in SNAC with homozygous *CDKN2A* loss and a monophasic cell population (**D**; case 11) as well as cases with no detectable genetic alterations (**E**; case 9) and peripheral cells that expressed p40 (**F**; case 9).

separate classification in this relatively limited cohort. However, SNAC that have recurrent *BRAF* p.V600E mutations, true myoepithelial elements, and prominent papillary and micropapillary architecture may represent a more unique category. Further clinical, histologic, immunohistochemical, and molecular evaluation of larger groups of SNAC will allow for better understanding of its pathogenesis, diagnosis, and potential subgroups in the future.

#### DATA AVAILABILITY

Most data generated or analyzed during this study are included in this published article. Additional data is available from the corresponding author on reasonable request.

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#### AUTHOR CONTRIBUTIONS

L.M.R. and J.A.B. designed the study and prepared the manuscript. L.M.R., J.A.B., L.D.R.T., J.S.G.H., N.R.L., M.W.M., and T.M.S. contributed tumor samples and data analysis. J.G. analyzed sequencing data and interpreted the results. All authors read and approved the final paper.

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#### COMPETING INTERESTS

N.R.L. receives funding from Merck and holds stock in Navigen Pharmaceuticals, neither of which are related to the present study. The other authors have no competing interests to declare.

#### ETHICS APPROVAL AND CONSENT TO PARTICIPATE

This study was approved by the Institutional Review Board at UT Southwestern Medical Center.

#### ADDITIONAL INFORMATION

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