Atypical Mesonephric Hyperplasia of the Uterus Harbors Pathogenic Mutation of Kirsten Rat Sarcoma 2 Viral Oncogene Homolog (\textit{KRAS}) and Gain of Chromosome 1q

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Abstract. Background/Aim: Mesonephric carcinoma (MNC) is a rare but notable entity of the female genital tract. While many researchers have acknowledged and studied MNC, much remains unknown on the characteristics of mesonephric remnant (MNR) or hyperplasia (MNH). There has not been any study examining the molecular features of MNR and MNH so far. The aim of this study was to investigate the clinicopathological and molecular characteristics of ten uterine mesonephric lesions, including two MNR without atypia, four MNHs without atypia, and three MNHs with atypia. Materials and Methods: We reviewed the electronic medical records and all available slides of ten cases from multiple institutions. Targeted sequencing and array comparative genomic hybridization were performed. Results: Three atypical MNHs displayed nuclear enlargement, mild-to-moderate nuclear pleomorphism, and nuclear membrane irregularity, and harbored pathogenic Kirsten rat sarcoma 2 viral oncogene homolog sarcoma 2 viral oncogene homolog (\textit{KRAS}) mutation. Two of those that co-existed with MNC harbored the same sequence alterations as each of their adjacent MNC. One of the three atypical MNHs harbored chromosome 1q gain. Conclusion: Atypical MNH is a potential premalignant lesion in which \textit{KRAS} mutation and chromosome 1q gain play an important role in the early stage of mesonephric carcinogenesis.

This mesonephric duct is a precursor of the male genital tract present during human embryogenesis (1). In males, it gives rise to the internal genitalia, including the epididymides, vasa deferentia, seminal vesicles, and efferent ductules of the testes, whereas in females, it regresses with some remnants persisting in the broad ligament and the uterine cervix (2). Mesonephric remnant (MNR) and hyperplasia (MNH) are not uncommon findings in specimens of conizations and hysterectomies, being reported in up to one-third of resected adult uterine cervixes (1). They might also be present within the wall of the vagina and uterine body, as well as the ovarian hilum and mesosalpinx (3).

Mesonephric carcinoma (MNC) is a rare malignant neoplasm thought to arise from the embryonal remnants of mesonephric tubules and ducts, comprising less than 1% of all gynecological tract malignancies (4). MNC arises
typically in the uterine cervix and vagina, although some cases of mesonephric-like carcinoma (MLC) arising in the uterine body and ovary have been reported (5, 6). Despite the rarity of MNC, its aggressive behavior and frequent distant metastasis compared to the more common types of uterine carcinoma warrant greater attention from the clinicians (6-9). MNC is also characterized by molecular aberrations that are significantly different from those found in other types of uterine carcinoma (10). MNC commonly harbors pathogenic mutations in Kirsten rat sarcoma 2 viral oncogene homolog (KRAS) gene and gain of chromosome 1q. In contrast, tumor protein 53 (TP53) mutations are uncommon, and phosphatidylinositol-4,5-bisphosphate 3-kinase catalytic subunit alpha (PIK3CA) and phosphatase and tensin homolog deleted on chromosome 10 (PTEN) mutations have not been identified in MNC.

While many researchers have acknowledged and studied MNC, much remains unknown on the characteristics of uterine mesonephric lesions, including whether MNR or MNH is the precursor of MNC. Moreover, there has not been any study examining the molecular features of MNR and MNH. In this study, we analyzed the clinicopathological characteristics of uterine mesonephric lesions, particularly MNH with atypia, and investigated their association with MNC by examining these lesions at the molecular level as well as their morphologies.

Materials and Methods

Case selection. This study was reviewed and approved by the Institutional Review Board of Severance Hospital (Seoul, Republic of Korea) (4-2017-0968). Using the combination of keywords ‘mesonephric’ and ‘hysterectomy’, we extracted ten cases of MNR or MNH from surgical pathology archives of multiple institutions. All patients underwent hysterectomy for malignant (9/10) and premalignant (1/10) lesions of the female genital tract. Clinical information, including age of patients, reason for surgery (primary indication for surgical treatment), International Federation of Gynecology and Obstetrics (FIGO) stage (11, 12), and surgical procedure, was obtained from the electronic medical record system and pathology reports.

Pathological examination. Three board-certified pathologists (H.Y.W., K.N., and H.-S.K.) specialized in gynecological oncology reviewed all available hematoxylin and eosin-stained slides by light microscopy and determined the following details in the mesonephric lesions: size and extent of the mesonephric lesion, nuclear atypia (enlargement, pleomorphism, membrane irregularity, mitotic count, and atypical mitotic figure), architectural abnormalities (destructive stromal infiltration, confluent growth, back-to-back tubular arrangement, and cribriform pattern), and coagulative tumor cell necrosis. Based on the presence or absence of nuclear atypia, we classified the mesonephric lesions into the following four categories: MNR without atypia, MNR with atypia, MNH without atypia, and MNH with atypia. We performed this four-tiered categorization prior to conducting the ancillary tests, and chose representative slides for each case to perform immunostaining, targeted sequencing, and array comparative genomic hybridization (CGH).

Targeted sequencing. DNA and RNA were isolated from 10-μm thick slices of formalin-fixed, paraffin-embedded (FFPE) tissue using a sterile 26-gauge needle and RecoverAll Multi-Sample RNA/DNA Isolation Workflow (Thermo Fisher Scientific, Waltham, MA, USA). The tumor tissue was obtained by manual microdissection and subjected to extraction of DNA and RNA for library preparation. The normal tissues of each case were obtained from the adjacent non-neoplastic area, DNA and RNA were quantified using the Qubit 2.0 Fluorometer (Thermo Fisher Scientific). DNA and RNA libraries were prepared as previously described (6, 13-22). These DNA libraries were generated from 20 ng of DNA per sample using an Ion AmpliSeq Library Kit 2.0 (Thermo Fisher Scientific) and Oncomine Comprehensive Assay (OCA) v1 panel (Thermo Fisher Scientific). RNA libraries were generated from 15 ng of RNA per sample using the Ion AmpliSeq RNA Library Kit (Thermo Fisher Scientific). Libraries were quantified using the Ion Library Universal Quantification Kit (Thermo Fisher Scientific). The OCA v1 panel (Thermo Fisher Scientific) included 143 genes, of which 73 oncogenes were interrogated for mutational hotspots and 26 tumor-suppressor genes were interrogated for all exons. The panel provided the capability to detect copy number alterations (CNAs) in 49 genes and fusion drivers in 22 genes. The gene list is available at: https://www.thermofisher.com/kr/ko/home/clinical/preclinical-companion-diagnostic-development/oncomine-oncology/oncomine-cancer-research-panel-workflow.html. Consecutively, a 60 pmol/l pool of DNA:RNA libraries at a 4:1 ratio was used to prepare the templated Ion Sphere Particle (Thermo Fisher Scientific). Sequencing was performed using the Ion 540 Kit-Chef (Thermo Fisher Scientific) and Ion S5 system (Thermo Fisher Scientific). Sequencing data of approximately 200 bp reads were generated after 500 flow runs.

Analysis of the sequencing data was performed using the Torrent Suite Software v5.2.2 (Thermo Fisher Scientific). This workflow was created by adding a custom hotspots Browser Extensible Data file to report mutations of interest and a custom CNA baseline (described below) using the manufacturer’s default workflow as described previously (13, 14). The pipeline included signaling processing, base calling, quality score assignment, adapter trimming, read mapping to the human genome assembly GRCh37, quality control of mapping, coverage analysis with down-sampling, and variant calling. The identification of variants was performed using the Torrent Variant Caller plug-in and Ion Reporter Software v5.2 (Thermo Fisher Scientific). Coverage maps were generated using the Coverage Analysis plug-in (Thermo Fisher Scientific). Additionally, ANNOVAR (http://annovar.openbioinformatics.org/) was used for functional annotation of identified single nucleotide polymorphisms (SNPs) to investigate their genomic locations and variation (23). To eliminate error artifacts, sequence data were visually confirmed using the Integrative Genomics Viewer (Broad Institute, Cambridge, MA, USA). This workflow was able to report SNPs and indels in as low as 1% of the variant allele fraction. Based on the results of a feasibility study, the variant allele fraction threshold was established at 5%. Copy number analysis was performed using the copy number module within the aforementioned workflow of the Ion Reporter Software v5.2 (Thermo Fisher Scientific). Copy numbers of 4 or greater were considered concordant if the orthogonal assay also reported a copy number of 4 or greater.
Table I. Clinicopathological characteristics of mesonephric remnants and hyperplasias.

| Category             | Case no. | Age (yrs) | Reason for surgery | FIGO stage | Surgery | Size (mm) | Extent | Nuclear enlargement | Nuclear pleomorphism | Nuclear membrane irregularity | Mitosis (per 10 HPFs) |
|----------------------|----------|-----------|--------------------|------------|---------|-----------|--------|--------------------|----------------------|-------------------------|------------------------|
| MNR without atypia   | 1        | 61        | MNC, cervix        | IB2        | TH+BSO+ PLND+PALND | 4 | Deep    | Absent          | Absent               | Absent                  | 0                      |
|                      | 2        | 59        | EC, grade 2, endometrium | II         | TH+BSO+ PLND+PALND | 5 | Superficial | Absent          | Absent               | Absent                  | 0                      |
|                      | 3        | 56        | MNC, cervix        | IB1        | TH+BSO | 11 | Deep    | Absent          | Absent               | Absent                  | 0                      |
|                      | 4        | 51        | U-EA, cervix       | IB1        | RH+BSO+PLND | 11 | Superficial and deep | Absent          | Absent               | Absent                  | 0                      |
|                      | 5        | 55        | EC, grade 1, endometrium | IA         | TH+BSO+ PLND+PALND | 7 | Superficial | Absent          | Absent               | Absent                  | 0                      |
|                      | 6        | 60        | HGSC, grade 3, bilateral ovaries | IIC     | TH+BSO+PLND+PALND+TOMT TH+LSO | 7 | Superficial | Absent          | Absent               | Absent                  | 0                      |
|                      | 7        | 55        | HSIL (CIN 3), cervix | NA         |               | 8 | Superficial | Absent          | Absent               | Absent                  | 0                      |
| MNR with atypia      | 8        | 55        | PD-EA, cervix      | IB1        | RH+BSO+PLND | 7 | Superficial | Present        | Mild to moderate | Present                  | 0                      |
|                      | 9        | 54        | MNC, cervix        | IIB        | TH+BSO | 12 | Superficial and deep | Present        | Mild to moderate | Present                  | 1                      |
|                      | 10       | 55        | MNC, cervix        | IVB        | TH+RSO | 11 | Superficial | Present        | Mild to moderate | Present                  | 0                      |

MNR: Mesonephric remnant; MNR: mesonephric hyperplasia; MNC: mesonephric carcinoma; EC: endometrioid carcinoma; U-EA: usual-type endocervical adenocarcinoma; HGSC: high-grade serous carcinoma; HSIL: high-grade squamous intraepithelial lesion; CIN: cervical intraepithelial neoplasia; PD-EA: poorly differentiated endocervical adenocarcinoma; FIGO: International Federation of Gynecology and Obstetrics; NA: not applicable; TH: total hysterectomy; BSO: bilateral salpingo-oophorectomy; PLND: pelvic lymph node dissection; PALND: paraaortic lymph node dissection; RH: radical hysterectomy; LSO: left salpingo-oophorectomy; TOMT: total omentectomy; HPFs: high-power fields.

Results

Clinical characteristics. Table 1 summarizes the clinicopathological characteristics of 10 patients with MNR or MHN. The patients’ age ranged between 51 and 61 (mean=56.1) years. Primary indications for the surgical procedures yielding specimens were cervical MNC (4/10), endometrial endometrioid carcinoma (2/10), endocervical adenocarcinoma (2/10), ovarian high-grade serous carcinoma (1/10), and cervical high-grade squamous intraepithelial lesion (1/10). None of the patients had any history of oral contraceptive use, postmenopausal hormone replacement therapy, or preoperative chemotherapy.

All four patients with MNC (cases 1, 3, 9, and 10) were postmenopausal and presented with vaginal bleeding. At the time of initial diagnosis, the FIGO stages were IB1, IB2, IIB (parametrial involvement), and IVB (pulmonary metastasis), respectively.

Pathological characteristics. Ten cases were classified into two MNRs without atypia (cases 1 and 2), five MHNs without atypia (cases 3-7), and three MHNs with atypia (cases 8-10). None of the cases was classified as MNR with atypia.

Array CGH. Both FFPE and reference DNA (NA10851) were labeled using an optimized version of the protocol for ULS labeling of FFPE DNA (Agilent Technologies, Santa Clara, CA, USA). Heat fragmentation at 95°C was required before labeling when the average fragment size was greater than 7.0 kb. Accordingly, 500 ng of FFPE DNA and reference DNA was then chemically labeled by incubating with ULS-Cy5 and Cy3, respectively, for 30 min. Labeling reactions were prepared in thin-walled 0.2 ml PCR tubes and incubated on a thermal cycler with a heated lid. Unreacted dye was then removed using KREApure filters (Agilent Technologies). DNA labeling efficiency was assessed by NanoDrop ND-2000 spectrophotometry (Thermo Fisher Scientific) measuring $A_{260}$ for DNA, $A_{550}$ (for Cy5), and $A_{649}$ (for Cy3) values for the determination DNA and fluorophore concentrations. The degree of labeling (DoL) is represented by the number of fluorophore molecules per 100 nucleotides, expressed as a percentage and was calculated from the post-labeling DNA yield and concentration of fluorophores. According to the manufacturer’s recommendations, DoL values between 0.75% and 2.5% were regarded as optimal for Cy5, whereas values between 1.75% and 3.5% were considered optimal for Cy3-labeled DNA.
MNR without atypia. Two MNRs were located in the deep and superficial cervical stroma, respectively. They consisted of clusters (Figure 1A), lobules, or linear arrays (Figure 1B) of small to medium-sized tubules, lined by bland cuboidal-to-columnar epithelial cells. Most of the mesonephric tubules contained dense (Figure 1C) or pale (Figure 1D) eosinophilic intraluminal secretions. Their nuclei were small and uniform (Figure 1E). Nuclear membrane irregularities were absent or
Figure 2. Histological features of mesonephric hyperplasia (MNH) without atypia. (A) Diffuse tubular proliferation without lobular growth is noted. Proliferation of small mesonephric tubules and varying degrees of intervening stroma are observed. (B) Uniform and bland nuclei of MNH are similar to those of mesonephric remnant. (C) Simple small mesonephric tubules radially distributed around large central ducts display luminal dilatation, irregular shape, and haphazard arrangement, but lack architectural abnormalities. Dilated ducts contain nothing or a small amount of eosinophilic secretion, whereas most of the surrounding tubules possess deeply eosinophilic hyaline-like material. (D) The tubular epithelium does not exhibit nuclear atypia. (E) MNH is observed along the pushing border of mesonephric carcinoma. The surrounding tubules and ducts are compressed and stretched over the border. (F-G) Nuclei of the lining epithelium are bland with minimal membrane irregularity. Intraluminal eosinophilic secretions are suggestive of their mesonephric origin.
minimal, and intranuclear grooves were rarely observed (Figure 1F).

**MNH without atypia.** The histological features of five MNHs were similar to those of MNR, but the tubules and ducts were present in greater abundance and were larger than 6 mm (2, 3, 24, 25). Two of the five cases (cases 4 and 5) displayed diffuse proliferation of mesonephric tubules with intervening stroma and without lobular or clustered growth (Figure 2A). Back-to-back tubular arrangement or cribriform architecture was not observed. There was no nuclear atypia, architectural complexity, or mitotic activity (Figure 2B). Two cases (cases 6 and 7) showed a few central mesonephric ducts surrounded by arranged congeries of simple mesonephric tubules (Figure 2C). Most mesonephric tubules were occasionally irregular-shaped and separated by varying amounts of intervening stroma. Elongated ducts contained nothing or a small amount of eosinophilic secretions, whereas most randomly scattered tubules possessed deeply eosinophilic materials. The nuclei of tubules and ducts were clearly bland, without hyperchromasia, pleomorphism, stratification, or mitosis (Figure 2D). In the remaining one case (case 3), the lesion was located adjacent to the expanding edge of the MNC, resulting in the compression, atrophy, and stretch of the surrounding tubules over the relatively well-defined tumor border (Figure 2E). The nuclei of MNH were small and bland with round (Figures 2F-G) or elongated shapes, whereas those of MNC were large, hyperchromatic, and pleomorphic with irregular nuclear membranes and frequent mitotic figures.

**MNH with atypia.** We identified three cases (cases 8-10) of MNH showing nuclear atypia compared to the non-atypical MNH. These lesions did not exhibit any atypical mitotic figure, architectural abnormality, or coagulative tumor cell necrosis. Thus, we considered that overall histological features in these lesions were not sufficient for the diagnosis of MNC. Detailed descriptions of the histological features are as follows.

In case 8, cystically dilated tubular proliferation was observed. Round or ovoid cystic structures were noted in the superficial cervical stroma, with small mesonephric tubules surrounding the cystic structures (Figure 3A). Some dilated tubules contained pale basophilic or mucoid secretions of lower density than those within the small tubules (Figure 3B). Complex glandular configurations were absent, and the intervening stroma was clearly observed. In contrast to cases 3-7, we noted a patchy distribution of nuclear atypia. Compared to the small and bland nuclei of the adjacent non-atypical tubules (Figure 3C), the nuclei of atypical mesonephric tubules displayed enlargement, mild-to-moderate pleomorphism, and irregular membrane (Figures 3D-E). High-power magnifications (Figure 3F-H) more clearly revealed that the nuclei were 1.5- to 3-times larger and more pleomorphic with irregular membranes than those of their non-atypical counterparts. Mitoses or atypical mitotic figures, as well as peritubular edema, inflammation, and necrosis, were absent.

In case 9, variable-sized cystic structures were randomly scattered at the periphery of the MNC (Figure 4A and B). The carcinoma cells showed tubular, endometrioid-like, cribriform, and glomeruloid architectures merged with round, ovoid, or irregular-shaped, cystically dilated tubules. Although some of these tubules exhibited the typical appearance of mesonephric tubules lined by cuboidal or flat epithelium and containing eosinophilic secretions (Figure 4C), many displayed nuclear atypia and stratification (Figure 4D). The nuclei were 1.5- to 2-times larger and showed mild-to-moderate pleomorphism and increased nuclear membrane irregularities than those of the non-atypical counterparts, displaying a morphology nearly identical to the nuclei of the adjacent MNC (Figure 4E). High-power magnifications (Figure 4F-H) displayed more clearly the enlarged nuclei with pleomorphism and irregular membranes in areas of atypical MNH and MNC compared with the non-atypical tubules. A single mitotic figure was detected in 10 high-power fields (Figure 4G). However, these atypical cells did not show any architectural complexity.

In case 10, we noted a closely packed tubular arrangement present around the MNC. At low-power magnifications, the carcinoma area appeared deeply basophilic due to hypercellularity and nuclear hyperchromasia, whereas the MNH area appeared pale or lightly eosinophilic (Figure 5A and B). Despite numerous mesonephric tubules being crowded and haphazardly scattered, the intervening stroma was well preserved (Figure 5C). Some areas showed very small or poorly formed glandular spaces, while most of the tubular lumina were patent with eosinophilic secretions. As in case 9, the nuclei of atypical MNH exhibited enlargement, mild-to-moderate pleomorphism, membrane irregularities, and occasional intranuclear grooves (Figure 5D), and the degree of atypia observed in these nuclei was nearly identical to that displayed by nuclei of the adjacent carcinoma (Figure 5E). We could not detect any mitotic activity, atypical mitotic figure, or coagulative tumor cell necrosis, as well as architectural abnormality, in the areas of MNH. In contrast, the MNC showed obvious back-to-back tubular arrangement, loss of intervening stroma, and cribriform architecture. High-power magnifications of atypical MNH (Figure 5F) revealed nuclear atypia nearly identical to that of MNC (Figure 5G and H).

**Molecular characteristics.** Both targeted sequencing and array CGH followed the histological classification of mesonephric lesions in four categories. Table II summarizes the results of targeted sequencing and array CGH. We found that three MNHs with atypia harbored pathogenic missense mutations of the KRAS codon 12, specifically, c.35G>A.
(p.G12D; case 8), c.34G>T (p.G12C; case 9), and c.35G>T (p.G12V; case 10). In two MNCs that co-existed with MNH with atypia (cases 9 and 10), the same type of KRAS mutation was identified (c.34G>T and c.35G>T, respectively). In contrast, there was no pathogenic mutation in one MNR (case 1) found along with MNC. The remaining cases, consisting of one MNR without atypia and five MNHs without atypia, had wild-type KRAS.
Figure 4. Histological features of mesonephric hyperplasia (MNH) with atypia (case 9). (A) Cystically dilated ductal structures (left and middle one-thirds) are observed at the periphery of mesonephric carcinoma (right one-third). Randomly scattered spaces of various shapes and sizes are observed. (B) Round, ovoid, or irregular-shaped cystically dilated tubules do not show architectural complexity. A difference in epithelial thickness between non-atypical and atypical tubules is noted. (C) Non-atypical MNH shows low cuboidal-to-flattened epithelium with small and bland nuclei. (D) The lining epithelium of atypical MNH is thicker than that of the non-atypical counterpart due to nuclear enlargement and stratification. No stromal desmoplasia or inflammation is observed. (E) Most nuclei observed in mesonephric carcinoma display mild atypia very similar to that of the atypical MNH. A single mitotic figure is detected (left upper corner). (F-H) High-power magnifications of images C-E. The nuclei of (G) atypical MNH demonstrate enlargement and irregular membrane. A single mitotic figure is detected (right lower corner).
Copy number plots obtained by array CGH are shown in Figure 6. All of the two MNRs, five MNHs without atypia, and two of the three MNHs with atypia did not show any CNA. In contrast, one MNH with atypia (case 9) showed chromosome 1q gain, and four MNCs showed aberrations in multiple chromosomes (Table III). The most common alterations observed in MNCs were the gains of chromosome 1q (4/4) and 10 (4/4), followed by gains of chromosome 2 (3/4), 12 (3/4), and 20 (3/4). We also observed the gains of chromosome 7 (1/4), 16 (2/4), and 17 (1/4) in MNCs.

Figure 5. Histological features of mesonephric hyperplasia (MNH) with atypia (case 10). (A) At low-power magnification, atypical MNH (left upper corner) appears pale compared to the hyperchromatic mesonephric carcinoma (MNC; middle and left one-thirds). (B) In another area, atypical MNH (middle and left two-thirds) appears lighter compared to the deeply basophilic MNC (right one-third). (C) Tubular lumina and intraluminal secretions are patent in atypical MNH. (D) Although numerous mesonephric tubules are crowded, the intervening cervical stroma is well preserved. (E) In contrast, MNC displays significant hypercellularity and back-to-back tubular arrangement. (F–H) High-power magnifications of images D and E. (F and G) The tubular epithelial cells show nuclear atypia including enlargement, significant membrane irregularity, and intranuclear grooves. (H) The degree of atypia observed in the nuclei of adjacent MNC is nearly identical to that of the atypical MNH.
Discussion

Ten cases of MNRs and MNHs were explored in search for their clinical and pathological significance. Based on the histological features reported in previous studies (2, 26), we classified the ten cases according to the presence or absence of nuclear atypia. Two cases were MNR without atypia; five were MNH without atypia; and three were MNH with atypia. Among these, we focused on the MNHs with atypia, i.e., atypical MNH. Despite the fact that these lesions have been acknowledged by only two studies (2, 26), it was evidently shown that these lesions were insufficient to call a carcinoma while being clearly different from MNHs and MNRs without atypia. Atypical MNH exhibited nuclear enlargement, mild-to-moderate pleomorphism, and irregular membrane in comparison with the non-atypical counterpart. While so, the atypical MNH cases did not possess the architectural abnormalities of MNC (destructive stromal infiltration, confluent growth, back-to-back tubular arrangement, and cribriform pattern), malignant nuclear features (severe nuclear pleomorphism, increased mitotic activity, and atypical mitotic figure), and coagulative tumor cell necrosis.

In one of the two reports that studied MNH with atypia, Ayroud et al. (26) observed florid tubular and ductular structures in the endocervical mucosa, differing from but intermingled with the endocervical glands. They noted that their intraluminal secretions with eosinophilia had similar characteristics to those of the adjacent MNR. Upon careful histological examination and exclusion of mesonephric and endocervical adenocarcinomas, the authors concluded that the florid tubular structures were hyperplasia with atypia rather than carcinoma, hence the term “atypical” or “florid” MNH. In another study, Ferry and Scully (2) openly stated that the division between MNC and MNH was arbitrary. Nevertheless, they pointed out that in the absence of clear diagnostic features of malignancy, the next step to detect the presence of MNC would be back-to-back glandular aggregation and disorderly invasion. The following step would be to identify cases with MNH appearing “predominantly typical, with only focal glandular crowding...
and nuclear atypia,” and call them MNH with atypia rather than carcinoma. The histological features we identified in our cases were concordant with those reported in these studies.

Thus, bringing together the histological analyses from previous studies and our study, we suggest to call these lesions atypical MNH defined by: the proliferation of mesonephric tubules or ducts, or both where 1) epithelial cells have nuclear atypia (e.g., mild-to-moderate pleomorphism, larger nuclei, and more nuclear membrane irregularities) than those of the adjacent non-atypical MNH, and where 2) epithelial cells do not form architectural abnormalities, such as destructive stromal infiltration, confluent growth, back-to-back glandular arrangement, and cribriform pattern.

All three atypical MNHs were studied at the molecular level for the first time to reveal some significant results. All of them harbored KRAS mutations, of which the type observed in these lesions were the same as those detected in the co-existing MNCs. In addition, one of the atypical MNH exhibited chromosome 1q gain, which was also found in the adjacent MNC amidst its other numerous mutations. The mutation of KRAS and the gain of chromosome 1q have been well acknowledged in MNC. Several targeted sequencing analyses from the past have shown that KRAS is the most common molecular abnormality in MNCs and MLCs. All 10 cases of MLC recently reported by Kolin et al. (27), Liang et al. (28), and Horn et al. (29) showed either p.G12V or p.G12D alterations. In larger cohorts, such as those reported

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Table II. Molecular characteristics of mesonephric remnants and hyperplasias.

| Category                  | Case no. | Gene    | Mutation type | Sequence change | Predicted effect | Copy number alteration          |
|---------------------------|----------|---------|---------------|-----------------|-----------------|---------------------------------|
| MNR without atypia        | 1        | KRAS    | Missense      | c.34G>T         | p.G12C          | Gain of chromosome 1q           |
|                           | 2        |         |               |                 |                 |                                 |
| MNH without atypia        | 3        |         |               |                 |                 |                                 |
|                           | 4        |         |               |                 |                 |                                 |
|                           | 5        |         |               |                 |                 |                                 |
|                           | 6        |         |               |                 |                 |                                 |
|                           | 7        |         |               |                 |                 |                                 |
| MNH with atypia           | 8        | KRAS    | Missense      | c.35G>A         | p.G12D          |                                 |
|                           | 9        | KRAS    | Missense      | c.34G>T         | p.G12C          |                                 |
|                           | 10       | KRAS    | Missense      | c.35G>T         | p.G12V          |                                 |

MNR: Mesonephric remnant; MNH: mesonephric hyperplasia.

Table III. Molecular characteristics of mesonephric carcinoma.

| Category      | Case no. | Gene    | Mutation type | Sequence change | Predicted effect | Copy number alteration          |
|---------------|----------|---------|---------------|-----------------|-----------------|---------------------------------|
| MNC           | 1        | KRAS    | Missense      | c.34G>T         | p.G12C          | 1q, chr 2, chr 5, 7q, chr 10, chr 12, chr 20; Loss: 7p |
|               | 3        | PIK3CA  | Missense      | c.278G>A        | p.R93Q          |                                 |
|               | 9        | KRAS    | Missense      | c.34G>T         | p.G12C          | 1q, 2q, chr 10, 12p, chr 20; Loss: 1p, 3p, 6q, chr 9, chr 15, 18q, chr 19, chr 22 |
|               | 10       | KRAS    | Missense      | c.35G>T         | p.G12V          |                                 |

MNC: Mesonephric carcinoma.
by Mirkovic et al. (10), Euscher et al. (8), and our group (6), the majority of MNCs (9/13) and MLCs (25/30) harbored KRA S mutations. The second most common alteration in MNC was the gain of 1q. Nine of the 13 MNCs and 16 of the 17 MLC reported by Mirkovic et al. (10), Kolin et al. (27), and our group (6) harbored 1q gain. Consistent with these data, all four cases of MNC in this study harbored both KRA S mutations and 1q gain. Our observations regarding the KRA S mutation and chromosome 1q gain in atypical MNH indicate that it could be a clonal lesion, and these two genetic alterations might be early events during the mesonephric carcinogenesis.

Mutations of KRA S result in constitutive activation of mitogen-activated protein kinase, which subsequently activates various downstream targets, leading to expression of genes involved in cellular proliferation, differentiation, and survival (10, 30, 31). Activating KRA S mutations have been detected in many human malignancies, particularly in pancreatic, colorectal, and pulmonary carcinomas, but also in endometrial and endocervical adenocarcinomas (32-36). At the early stage of carcinogenesis, the precancerous cells require KRA S for survival. KRA S activity is also necessary to continue the neoplastic transformation and the regulation of cellular differentiation, indicating that KRA S mutations are important not only for tumor formation but also during early stages of tumor progression (37). Our observation of pathogenic mutations in codon 12 of the KRA S gene in all atypical MNHs and in none of MNHs without atypia suggest that KRA S mutation is one of the early events during the mesonephric carcinogenesis.

1q gain is considered one of the early changes occurring in mammary carcinogenesis since they have been detected as the sole chromosomal abnormality in well-differentiated breast carcinoma with a few alterations (38, 39). These genetic changes and the resulting chromosome imbalances have been thought to play a pathogenic role in breast carcinoma development. Similarly, 1q gain is one of the most common cytogenetic abnormalities in patients with multiple myeloma, occurring in approximately 40% of patients (40, 41). 1q gain occurs with disease progression in multiple myeloma, often by jumping translocations. The copy number can increase over time, with higher rates of 1q gain detection after progression from precursor conditions including monoclonal gammopathy of undetermined significance and smoldering multiple myeloma (41, 42). In addition, 1q gain is associated with increased risk of relapse and adverse outcome in patients with favorable-histology Wilms’ tumor (43). CNA in malignant mesonephric lesions were primarily at the chromosome or arm level, without distinct copy number changes. In our previous study, 1q gain was the most common CNA detected in 91.7% of MLCs (6). Based on the previous data that the majority of MNCs and MLCs harbored 1q gain and on our observations that 1q gain was the only chromosomal abnormality in atypical MNHs, we suggest that this alteration is one of the early stage events of mesonephric carcinogenesis. In 2017, a report by Mirkovic et al. (25) suggested that MNH might not be neoplastic. In their case series, none of the 10 MNH harbored mutations in KRA S. However, because they do not describe any atypical MNH, we believe that their argument does not oppose ours. In agreement with their data, all five MNHs without atypia in our series did not harbor any KRA S mutation or CNA.

In conclusion, we demonstrated three cases of MNH showing mild-to-moderate nuclear pleomorphism, nuclear enlargement, and irregular nuclear membranes compared to the adjacent non-atypical MNH. These atypical MNHs displayed mild to moderate nuclear atypia, the degree of which was similar to that of the adjacent MNC, but without any architectural abnormality. All three atypical MNHs were found to harbor KRA S mutations, and one of them also exhibited chromosome 1q gain. Both the pathogenic mutations in KRA S and gain of 1q were observed in all MNC examined. In contrast, none of the MNRs or MNHs without atypia was shown to have pathogenic mutations or chromosomal aberrations. These findings raise the possibility that atypical MNH is potentially a premalignant lesion of MNC, and that KRA S mutation and 1q gain are early events during mesonephric carcinogenesis.

Conflicts of Interest
None of the Authors has any conflict of interest to declare.

Authors’ Contributions
All Authors made substantial contributions to the conceptualization and design of the study; the acquisition, analysis, curation, and interpretation of the data; drafting of the manuscript; critical revision and editing of the manuscript for important intellectual content; and the approval of the final version to be published.

Acknowledgments
This research was supported by the Basic Science Research Program through the National Research Foundation of Korea (NRF) funded by the Ministry of Education (2016R1D1A1B03935584) and by the NRF funded by the Korean government (Ministry of Science and ICT) (2018R1C1B5043725).

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Received June 22, 2020
Revised July 20, 2020
Accepted July 21, 2020