

РОЛЬ EZH2 И ARID1A В ДИАГНОСТИКЕ ПЛОСКИХ УРОТЕЛИАЛЬНЫХ ОПУХОЛЕЙ С АТИПИЕЙ

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Аннотация

Актуальность. Диагностика уротелиальной карциномы in situ имеет большое значение, поскольку обладает прогностической и терапевтической ценностью. Целью исследования было определить роль EZH2 и ARID1A в диагностике карциномы in situ. Материал и методы. Ретроспективное пере-крестное исследование включало 24 образца плоских уротелиальных опухолей, 20 образцов CIS и 10 образцов нормального прилегающего уротелия, взятых при цистоскопической биопсии. Во всех случаях была оценена иммуногистохимическая экспрессия EZH2 и ARID1A. Результаты. Все образцы нормального уротелия показали высокое ядерное окрашивание на ARID1A и отрицательное ядерное окрашивание на EZH2. Высокая экспрессия EZH2 наблюдалась в 80 % образцов CIS по сравнению с 20 % плоских уротелиальных опухолей с атипней (p=0.001), в то время как высокая экспрессия ARID1A наблюдалась в 70,8 % плоских уротелиальных опухолей с атипней по сравнению с 25 % образцов CIS (p=0.001). EZH2 был более точным и специфичным при диагностике карциномы in situ. Заключение. EZH2 и ARID1A являются перспективными диагностическими маркерами уротелиальной карциномы in situ. EZH2 более точен и специфичен, чем ARID1A, в диагностике карциномы in situ по сравнению с другими плоскими уротелиальными опухолями.

Ключевые слова: дисплазия, карцинома in situ, реактивная атипия, EZH2, ARID1A, иммуногистохимия.

THE ROLE OF EZH2 AND ARID1A IN THE DIAGNOSIS OF FLAT UROTHELIAL LESIONS WITH ATYPIA

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Abstract

Background. Diagnosis of urothelial carcinoma in situ is of great importance because it has prognostic and therapeutic value. We aim to determine the utility of EZH2 and ARID1A as a new tool in the diagnosis of carcinoma in situ. Material and Methods. This retrospective cross-sectional study included Twenty-four specimens of flat urothelial lesions, twenty specimens of CIS, and 10 of normal adjacent urothelium that was taken by cystoscopically resection biopsy procedure. Immunohistochemical expression of EZH2 and ARID1A were evaluated in all studied cases. Results. All normal urothelium specimens showed high nuclear staining for ARID1A and negative nuclear staining for EZH2. High EZH2 expression was observed in 80 % of CIS specimens compared to 20 % of flat urothelial lesions with atypia (p=0.001), while high ARID1A expression was observed in 70.8 % of flat urothelial lesions with atypia compared to 25 % of CIS specimens (p=0.001). EZH2 was more accurate and specific in the diagnosis of carcinoma in situ. Conclusion. EZH2 and ARID1A are promising diagnostic markers for urothelial CIS. EZH2 is more accurate and specific than ARID1A in the diagnosis of carcinoma in situ versus other flat urothelial lesions.

Key words: dysplasia, carcinoma in situ, reactive atypia, EZH2, ARID1A, immunohistochemistry.

Introduction

Urothelial carcinoma is a major health problem. The incidence of urothelial carcinoma worldwide is approximately 4.5 % in males and 1.5 in females [1]. In Egypt, it represents the second mostly diagnosed cancers in men [2]. Early diagnosis of UC may decrease the mortality and morbidity rates [3]. According to the latest WHO 2016 committee, flat urothelial lesions with atypia were classified as reactive urothelial atypia, urothelial proliferation of uncertain malignant potential (UPUMP), urothelial dysplasia and urothelial carcinoma in situ [4]. UPUMP contains no true papillary projections but undulations are common with a thickened urothelium but minimal or no cytological atypia. These entities may be seen de novo, and in this setting, the clinical relevance is unknown. More frequently they are seen in patients who have a history of prior carcinoma or seen adjacent to papillary lesions. It is likely that most represent lateral extension (“shoulder lesion”) of a papillary neoplasm [5]. Reactive urothelial atypia may in some instances lead to confusion with dysplasia or cause concern for patients. Nuclemegaly is the most prominent finding in reactive urothelial changes, but the cells often have a single prominent nucleolus and evenly distributed vesicular chromatin. The nuclear borders are smooth. The nuclei are frequently round, and nuclear pleomorphism is lacking [6]. Architecturally, the cells maintain their polarity to the basement membrane although a minimal loss of polarity may be evident. The mitotic rate may be increased with mitoses present predominantly in the basal and intermediate urothelium, but atypical forms are not seen. Intraurothelial acute or chronic inflammation is commonly identified [7]. It is important to recognize that even intraepithelial lymphocytes, by themselves, can result in reactive changes. The cytoplasm may become more basophilic or cosinophilic with loss of cytoplasmic clearing. Clinical history of stones, infection, or frequent instrumentation may be present. Reactive urothelium may be denuded with only a single residual layer of basal cells remaining; the residual cells are not hyperchromatic, not enlarged, and do not possess nuclear membrane irregularity [8]. Urothelial dysplasia (UD) is defined as the loss of polarity with nuclear rounding and crowding and cytologic atypia that is not severe enough to diagnose CIS. CIS and UD are precursor lesions of invasive urothelial carcinoma and their detection, especially CIS, is associated with a significant risk of progression and recurrence [9]. CIS is often multifocal and can occur in the upper urinary tracts and in the prostatic ducts and urethra. CIS exists in two settings; isolated (primary) CIS and secondary CIS associated with papillary urothelial carcinoma. Isolated CIS was rare, accounting for about 10 % of all CIS and 1 % to 3 % of bladder neoplasm [10]. Although nuclear and architectural features are the primary criteria for differentiation between CIS and other flat epithelial lesions with atypia, it may be difficult in some patients. Expression of markers as EZH2 and ARID1A may be helpful [11, 12].

The Enhancer of Zeste Homolog 2 (EZH2) is a core subunit of the polycomb repressor complex 2 (PRC2), which is overexpressed in numerous cancers and mutated in several others. Notably, EZH2 acts as a critical epigenetic repressor through its role in histone methylation, it is also an activator of gene expression, acting through multiple signaling pathways in distinct cancer types. Increasing evidence suggests that EZH2 is an oncogene and is central to initiation, growth and progression of urological cancers [13]. AT-rich interacting rich domain 1 (ARID1A) belongs to a family of proteins containing a highly conserved, approximately 100-amino acid DNA binding domain called ARID (AT-rich interacting domain). Although the ARID domains in general preferentially bind AT-rich DNA sequences, the ARID1A domain of mammalian ARID1A exhibits general DNA binding character without sequence specificity [14]. The ARID1A gene maps to chromosome 1p36.11, a region frequently deleted in cancer. Initial clues that ARID1A was a tumor suppressor came from expression analyses that showed decreased ARID1A expression in 30 % of renal cancer and 10 % of breast cancer [15]. A synthetic lethality relationship between other SWI/SNF components including ARID1A and EZH2 has been
revealed in several tumor entities [16]. We evaluate the validity of EZH2 and ARID1A expression in distinction between carcinoma in situ and other flat urothelial lesions

**Material and Methods**

This retrospective cross-sectional study was performed after approval by the local ethical Committee Zagazig University, Institutional Review Board (IRB) for human studies (reference number is ZU-IRB #:3986-18-9-2017). This study included 54 patients were selected from pathology department archival blocks in collaboration with urology department, Faculty of Medicine, Zagazig University, Egypt, in the period from June 2016 to June 2020. The control group included 10 specimens from adjacent normal mucosa. Twenty-four patients with flat urothelial lesions with atypia (nine specimens of UPUMP, eight specimens of reactive atypia and seven specimens of dysplasia) and 20 specimens of CIS. All specimens were obtained by urethrocytoscop and biopsies. They were classified according to 2016 WHO/ ISUP [4].

**Immunohistochemical staining**

Staining was performed with Dako Autostainer link 48 (Dako) according to manufacturer’s guide.

**Irry antibodies**

1 – ARID1A [Rabbit polyclonal antibody (Cat. GTX129433) isotype: IgG, 1:100-1000 dilution, Gene Tex international corporation].

2 – EZH2 [Rabbit polyclonal antibody, Clone 144CT2.1.1.5, isotype: IgG, 1:50-100 dilution, US Biological life sciences USA] Normal human testicular tissue was used as positive control for EZH2 while normal human breast and kidney tissue served as positive control for ARID1A. Negative controls were obtained by replacing the primary antibodies with non-immune serum.

**The evaluation of ARID1A immunostaining**

Nuclear ARID1A expression was considered positive. Immunoreactivity was assessed by considering the extent and the intensity of nuclear staining in the tumor cells. Grades of 0–3 were assigned according to the percentage of positive tumor cells (0=0 %; 1<25 %; 2=25–50 %; 3>51 %) and the intensity of staining in tumor (0=absence of staining; 1=weak; 2=intermediate; 3=strong intensity). The combined score was calculated by multiplication of the percentage and the intensity grade [17]. Receiver operating characteristic (Roc) curve analysis was used for calculating the cutoff point.

**The evaluation of EZH2 immunostaining**

The EZH2 expression was considered positive in nuclear expression only. The extent of nuclear EZH2 protein staining was graded as follows: 1 (<25 % staining of tumor cells), 2 (25–50 % staining of tumor cells), 3 (50–75 % staining of tumor cells), 4 (>75 % staining of tumor cells). Moreover, the staining intensity was quantified as 0 (indicated no expression), 1 (indicated weak expression), 2 (indicated intermediate expression), 3 (indicated strong expression).

The intensity was multiplied by the extension values to obtain the final immunostaining score (range 0–12) [18]. Receiver operating characteristic (Roc) curve analysis was used for calculating the cutoff point.

**Statistical Analysis**

Statistical analysis was carried out by using SPSS version «20.0» (SPSS Inc.). Fisher's exact and $\chi^2$ tests were applied for comparisons between the nominal variables. Independent student t-test (t) was used to compare two groups of normally distributed data Mann-Whitney U (MW) was used to compare two groups of non-normally distributed data. Sensitivity, specificity, accuracy, and positive predictive values were calculated. The designations true and false were based on the study hypothesis that high EZH2 is expressed in malignancy and true positivity for ARID1A is expected in benign and other flat epithelial lesions.

**Results**

Mean age of the cases was 39.8 ± 1.3, most cases were males (79.2 %). According to Roc curve analysis the cutoff point of EZH2 immunoreactivity in all studied cases was 4 (Fig. 1). They were reclassified into two group; low expression (score<4,) and high expression (score≥4 was). However, the cutoff point
Сравнение EZH2 и ARID1A между CIS и другими плоскими уротелиальными опухолями с атипиею

Comparison of EZH2 and ARID1A between CIS and other flat urothelial lesions with atypia

<table>
<thead>
<tr>
<th></th>
<th>EZH2 Immunoreactivity</th>
<th>p-value</th>
<th>ARID1A Immunoreactivity</th>
<th>p-value</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Low (cut off &lt;4)</td>
<td></td>
<td>Low (cut off &lt;3)</td>
<td></td>
</tr>
<tr>
<td>Normal/Нормальные</td>
<td>10 10 (100.0 %)</td>
<td></td>
<td>0 (0.0 %)</td>
<td></td>
</tr>
<tr>
<td>平贫血橊性悪性/ Flat epithelial lesions with atypia</td>
<td>24 20(80 %)</td>
<td>4(20 %)</td>
<td>1 29.2 %</td>
<td>17 (70.8 %)</td>
</tr>
<tr>
<td>• UPUMP1</td>
<td>9 9 (100 %)</td>
<td></td>
<td>3 (11.1 %)</td>
<td></td>
</tr>
<tr>
<td>• Reактивная атипия/ Reactive atypia</td>
<td>8 7 (87.5 %)</td>
<td>1(12.5 %)</td>
<td>2 25.0 %</td>
<td>6 (75.0 %)</td>
</tr>
<tr>
<td>• Дисплазия/Dysplasia</td>
<td>7 4 (57.1 %)</td>
<td>3 (42.9 %)</td>
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<td>3 (42.9 %)</td>
</tr>
<tr>
<td>CIS2</td>
<td>20 4 (20 %)</td>
<td></td>
<td>3 (75.0 %)</td>
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</table>

Примечание: 1 – UPUMP: уротелиальная пролиферация с неопределенным злокачественным потенциалом; 2 – CIS: Carcinoma in situ; ** – различия статистически значимые (p<0.001).

Note: 1 – UPUMP: urothelial proliferation of uncertain malignant potential, 2 – CIS: Carcinoma in situ. ** – p<0.001 is highly significant.
Expression of EZH2 in ARID1A in CIS versus other flat lesions

<table>
<thead>
<tr>
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<th>ARID1A level</th>
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<tr>
<td></td>
<td>Mean ± SD</td>
<td>Median (Range)</td>
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<td>Flat normal</td>
<td>1.7 ± 1.1</td>
<td>1 (1–4)</td>
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</tr>
<tr>
<td></td>
<td>Reactive atypia</td>
<td>2.0 ± 1.4</td>
<td>1.5 (1–4)</td>
</tr>
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<td>CIS vs dysplasia</td>
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<tr>
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Table 2/Table 2

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Примечание: ^= Тест Манна-Уитни; 1 – CIS: карцинома in situ; 2 – UPUMP: уротелиальная пролиферация с неопределенным злокачественным потенциалом; * – различия статистически значимые (p<0.05); ** – различия статистически значимые (p≤0.001).

Note: ^= Mann-Witenny test; 1 – CIS: carcinoma in situ; 2 – UPUMP: urothelial proliferation of uncertain malignant potential; * – p<0.05 is significant, ** – p≤0.001 is highly significant.

Discussion

Early detection of CIS would help in reduction of bladder cancer incidence, morbidity and mortality. The differentiation of CIS from other flat lesions with atypia is critical because it has both therapeutic and prognostic importance [19, 20] Even after publication of the 4th edition of WHO/ISUP classifications, the distinction between reactive and dysplastic changes has not been resolved, There are still no definite morphological criteria to diagnose CIS, and there is great
inter- and interobserver disagreement [4]. Some forms of CIS may be difficult in diagnosis such as clinging and pagetoid type. So morphology alone is frequently insufficient to differentiate CIS from other flat lesions with atypia [21]. There is always a thorough work to discover a diagnostic marker of CIS to differentiate it from other flat epithelial lesions. Moreover, UPUMP has 40 % risk to turn into low grade urothelial carcinoma after 5 years [4]. Earlier studies suggested the use of CK20, CD44, Ki67, p53, p16 and CK5/6 immunostains as a diagnostic panel for problematic flat lesions with atypia [22], this is a large panel with a high cost and the discriminatory performance may be unsatisfactory in some cases.

EZH2-immunohistochemistry was used in distinction between malignant and reactive mesothelial cells, with 95.5 % sensitivity providing a promising diagnostic marker for malignancy [23]MOC-31, CEA, or B72.3. Also, ARID1A was used as a diagnostic marker for atypical endometrial hyperplasia to
differentiate it from benign hyperplasia [24]. These findings raised our interest for testing both markers in bladder tissue. There are no previous data about the role of EZH2 and ARID1A in distinction between carcinoma in situ and other flat urothelial lesions. In current study, we investigated the use of EZH2 and ARID1A as a diagnostic marker of CIS. We reported that expression of high EZH2 was observed in (80%) of cases of CIS compared to 20% of flat epithelial lesions with atypia and 0% of adjacent normal epithelium with a highly statistically significant difference between them (p-value=0.001), suggesting that EZH2 is a diagnostic marker for CIS. This observation goes in agreement with Warrick et al. (2016) study which found that EZH2 expression was common in bladder cancer with greatest expression seen in CIS in comparison to normal urothelium and recommended EZH2 as a specific marker of CIS and invasive bladder cancer compared to benign urothelium [11]. J.D. Raman et al., (2005) also demonstrated that EZH2 protein levels were higher in UC compared with benign urothelium [25]. Concerning ARID1A our results revealed that ARID1A expression was significantly low in CIS cases (either isolated or concomitant), compared to its high expression in 100% of normal epithelium and 70.8% of flat epithelial lesions with atypia with a highly statistically significance difference between them (p-value=0.0002) suggesting the use of ARID1A as a diagnostic marker for CIS. Similarly, T. Aso et al., (2015) reported that loss of ARID1A tumor suppressor gene may play an important role in carcinogenesis and progression of gastric dysplastic lesions to overt carcinoma [26]. This observation goes in agreement with Q. Cao et al. (2019) who reported that ARID1A expression was significantly downregulated in carcinoma tissues compared with normal tissues (p=0.002) suggesting that ARID1A may act as a tumor suppressor in the development of UC and its downregulation may play a role in progression of flat urothelial lesions with atypia to carcinoma [21].

In comparing the validity of EZH2 and ARID1A in diagnosis carcinoma in situ among the studied group, we found that both high EZH2 and low ARID1A expression can be used as a diagnostic marker to correctly identify cases of CIS (sensitivity 80% and 75% respectively). Moreover, high EZH2 is more specific and accurate than low ARID1A expression in differentiating CIS form normal and other flat epithelial lesions of the bladder (specificity 88.24% and accuracy 85.19%) that will be of great value to surgical pathologists as adjuncts to morphology in this setting especially in small biopsy specimens and urine cytology for accurate diagnosis.

Several therapeutic targets in ARID1A mutated cancers are in development, including EZH2 inhibitors. EZH2 facilitates epigenetic methylation to modulate gene expression. EZH2 inhibition in ARID1A mutated tumors acts in a synthetically lethal manner to suppress cell growth and promote apoptosis [27], revealing a new therapeutic target.

Conclusion

EZH2 and ARID1A expression could be used to differentiate CIS from benign and other flat epithelial lesions. However, EZH2 is more specific and accurate in diagnosis of CIS than ARID1A.

This study has been approved by the Faculty of Medicine, Zagazig University, Institutional Review Board (IRB) for human studies (reference number is ZU-IRB # 3:986-18-9-2017) and the patients have signed an informed written consent.


Diagnostic Value of ARID1A, ARID1B, and ARID2 Expression During Progression of Gastric Cancer. Anticancer Res. 2015 Dec; 35(12): 6819–27.


Variants and new entities of bladder cancer. Raspollini M.R., Cheng L.


AUTHOR CONTRIBUTION

Reham Sameh: study conception and methodology, statistical analysis, data collection and analysis, drafting of the manuscript.
Naglaa Mostafa: data collection and analysis, writing of the manuscript.
Ahmed Embaby: drafting of the manuscript.
Samar Abdel Raouf: data collection and analysis, writing of the manuscript.
Khaled Abdelwahab: data collection and analysis, drafting of the manuscript.

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Conflict of interest
The authors declare that they have no conflict of interest.